# The Enzyme List Class 3 — Hydrolases

# Nomenclature Committee of the International Union of Biochemistry and Molecular Biology (NC-IUBMB)

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# EC 3.1 Acting on ester bonds

This subclass contains the esterase enzymes. The esterases are subdivided into: carboxylic-ester hydrolases (EC 3.1.1), thioester hydrolases (EC 3.1.2), phosphoric-monoester hydrolases, the phosphatases (EC 3.1.3), phosphoric-diester hydrolases (EC 3.1.4), triphosphoric-monoester hydrolases (EC 3.1.5), sulfuric-ester hydrolases, the sulfatases (EC 3.1.6), diphosphoric monoesterases (EC 3.1.7) and phosphoric-triester hydrolases (EC 3.1.8). The nucleases, previously included under EC 3.1.4, are now placed in a number of new sub-subclasses: the exonucleases (EC 3.1.11-16) and the endonucleases (EC 3.1.21-31).;p; EC 3.1.23 and EC 3.1.24

In a previous edition, site-specific endodeoxyribonucleases were set out individually in subclasses EC 3.1.23 and EC 3.1.24 (since deleted), with 113 separate entries. These are now included in three entries EC 3.1.21.3, EC 3.1.21.4 and EC 3.1.21.5. A complete listing of all of these enzymes has been produced by R.J. Roberts and is available at http://rebase.neb.com/rebase/rebase.html.

# EC 3.1.1 Carboxylic-ester hydrolases

EC 3.1.1.1

Accepted name: carboxylesterase

**Reaction:** a carboxylic ester  $+ H_2O =$  an alcohol + a carboxylate

Other name(s): ali-esterase; B-esterase; monobutyrase; cocaine esterase; procaine esterase; methylbutyrase; vitamin

A esterase; butyryl esterase; carboxyesterase; carboxylate esterase; carboxylic esterase; methylbutyrate esterase; triacetin esterase; carboxyl ester hydrolase; butyrate esterase; methylbutyrase; α-carboxylesterase; propionyl esterase; nonspecific carboxylesterase; esterase D; esterase B; esterase

A; serine esterase; carboxylic acid esterase; cocaine esterase

**Systematic name:** carboxylic-ester hydrolase

**Comments:** Wide specificity. The enzymes from microsomes also catalyse the reactions of EC 3.1.1.2

(arylesterase), EC 3.1.1.5 (lysophospholipase), EC 3.1.1.6 (acetylesterase), EC 3.1.1.23 (acylglycerol lipase), EC 3.1.1.28 (acylcarnitine hydrolase), EC 3.1.2.2 (palmitoyl-CoA hydrolase), EC 3.5.1.4

(amidase) and EC 3.5.1.13 (aryl-acylamidase). Also hydrolyses vitamin A esters.

**References:** [97, 141, 207, 325, 1177, 1790, 1889, 2479]

[EC 3.1.1.1 created 1961]

EC 3.1.1.2

Accepted name: arylesterase

**Reaction:** a phenyl acetate  $+ H_2O = a$  phenol + acetate **Other name(s):** A-esterase; paraoxonase; aromatic esterase

**Systematic name:** aryl-ester hydrolase

**Comments:** Acts on many phenolic esters. The reactions of EC 3.1.8.1 aryldialkylphosphatase, were previ-

ously attributed to this enzyme. It is likely that the three forms of human paraoxonase are lactonases rather than aromatic esterases [1434, 645]. The natural substrates of the paraoxonases are lactones [1434, 645], with  $(\pm)$ -5-hydroxy-6*E*,8*Z*,11*Z*,4*Z*-eicostetraenoic-acid 1,5-lactone being the best sub-

strate [645].

**References:** [30, 102, 267, 1445, 1768, 1, 1434, 645]

[EC 3.1.1.2 created 1961, modified 1989]

EC 3.1.1.3

Accepted name: triacylglycerol lipase

**Reaction:** triacylglycerol +  $H_2O$  = diacylglycerol + a carboxylate

Other name(s): lipase (ambiguous); butyrinase; tributyrinase; Tween hydrolase; steapsin; triacetinase; tributyrin es-

terase; Tweenase; amno *N*-AP; Takedo 1969-4-9; Meito MY 30; Tweenesterase; GA 56; capalase L; triglyceride hydrolase; triolein hydrolase; tween-hydrolyzing esterase; amano CE; cacordase; triglyceridase; triacylglycerol ester hydrolase; amano P; amano AP; PPL; glycerol-ester hydrolase; GEH; meito Sangyo OF lipase; hepatic lipase; lipazin; post-heparin plasma protamine-resistant lipase; salt-resistant post-heparin lipase; heparin releasable hepatic lipase; amano CES; amano B; tributyrase;

triglyceride lipase; liver lipase; hepatic monoacylglycerol acyltransferase

**Systematic name:** triacylglycerol acylhydrolase

**Comments:** The pancreatic enzyme acts only on an ester-water interface; the outer ester links are preferentially

hydrolysed.

**References:** [1511, 1760, 2528, 2669, 2670]

[EC 3.1.1.3 created 1961]

EC 3.1.1.4

**Accepted name:** phospholipase A<sub>2</sub>

**Reaction:** phosphatidylcholine +  $H_2O = 1$ -acylglycerophosphocholine + a carboxylate

**Other name(s):** lecithinase A; phosphatidase; phosphatidolipase; phospholipase A

**Systematic name:** phosphatidylcholine 2-acylhydrolase

**Comments:** Also acts on phosphatidylethanolamine, choline plasmalogen and phosphatides, removing the fatty

acid attached to the 2-position. Requires Ca<sup>2+</sup>.

**References:** [628, 801, 1042, 1957, 2497, 3032]

[EC 3.1.1.4 created 1961, modified 1976, modified 1983]

EC 3.1.1.5

**Accepted name:** lysophospholipase

**Reaction:** 2-lysophosphatidylcholine +  $H_2O$  = glycerophosphocholine + a carboxylate

**Other name(s):** lecithinase B; lysolecithinase; phospholipase B; lysophosphatidase; lecitholipase; phosphatidase B;

lysophosphatidylcholine hydrolase; lysophospholipase A1; lysophopholipase L2; lysophospholipase

transacylase; neuropathy target esterase; NTE; NTE-LysoPLA; NTE-lysophospholipase

**Systematic name:** 2-lysophosphatidylcholine acylhydrolase

**References:** [5, 483, 551, 732, 2622, 3033, 3035, 3048, 2345, 1756, 3198]

[EC 3.1.1.5 created 1961, modified 1976, modified 1983]

EC 3.1.1.6

Accepted name: acetylesterase

**Reaction:** an acetic ester +  $H_2O$  = an alcohol + acetate

**Other name(s):** C-esterase (in animal tissues); acetic ester hydrolase; chloroesterase; p-nitrophenyl acetate esterase;

Citrus acetylesterase

**Systematic name:** acetic-ester acetylhydrolase

**References:** [30, 196, 1308]

[EC 3.1.1.6 created 1961]

EC 3.1.1.7

**Accepted name:** acetylcholinesterase

**Reaction:** acetylcholine +  $H_2O$  = choline + acetate

Other name(s): true cholinesterase; choline esterase I; cholinesterase; acetylthiocholinesterase; acetylcholine hydro-

lase; acetyl.β-methylcholinesterase; AcCholE

**Systematic name:** acetylcholine acetylhydrolase

**Comments:** Acts on a variety of acetic esters; also catalyses transacetylations.

**References:** [98, 197, 453, 1653, 2021, 3344]

#### [EC 3.1.1.7 created 1961]

EC 3.1.1.8

Accepted name: cholinesterase

> **Reaction:** an acylcholine +  $H_2O$  = choline + a carboxylate

pseudocholinesterase; butyrylcholine esterase; non-specific cholinesterase; choline esterase II (un-Other name(s):

specific); benzoylcholinesterase; choline esterase; butyrylcholinesterase; propionylcholinesterase;

**BtChoEase** 

**Systematic name:** acylcholine acylhydrolase

> **Comments:** Acts on a variety of choline esters and a few other compounds.

**References:** [98, 102, 1493, 2021, 2548, 2780]

[EC 3.1.1.8 created 1961]

[3.1.1.9 Deleted entry. benzoylcholinesterase; a side reaction of EC 3.1.1.8 cholinesterase]

[EC 3.1.1.9 created 1961, deleted 1972]

EC 3.1.1.10

Accepted name: tropinesterase

> Reaction: atropine +  $H_2O$  = tropine + tropate

Other name(s): tropine esterase; atropinase; atropine esterase

**Systematic name:** atropine acylhydrolase

> **Comments:** Also acts on cocaine and other tropine esters.

**References:** [930, 1956]

[EC 3.1.1.10 created 1961, deleted 1972, reinstated 1976]

EC 3.1.1.11

Accepted name: pectinesterase

> **Reaction:** pectin + n H<sub>2</sub>O = n methanol + pectate

pectin demethoxylase; pectin methoxylase; pectin methylesterase; pectase; pectin methyl esterase; Other name(s):

pectinoesterase

**Systematic name:** pectin pectylhydrolase References:

[593, 1704, 1918]

[EC 3.1.1.11 created 1961]

[3.1.1.12 Deleted entry. vitamin A esterase, now believed to be identical with EC 3.1.1.1 carboxylesterase]

[EC 3.1.1.12 created 1961, deleted 1972]

EC 3.1.1.13

Accepted name: sterol esterase

> **Reaction:** a steryl ester +  $H_2O$  = a sterol + a fatty acid

cholesterol esterase; cholesteryl ester synthase; triterpenol esterase; cholesteryl esterase; cholesteryl Other name(s):

ester hydrolase; sterol ester hydrolase; cholesterol ester hydrolase; cholesterase; acylcholesterol lipase

steryl-ester acylhydrolase **Systematic name:** 

A group of enzymes of broad specificity, acting on esters of sterols and long-chain fatty acids, that **Comments:** 

may also bring about the esterification of sterols. Activated by bile salts.

References: [1216, 2185, 3028, 3127]

[EC 3.1.1.13 created 1961, modified 1990]

#### EC 3.1.1.14

Accepted name: chlorophyllase

**Reaction:** chlorophyll +  $H_2O$  = phytol + chlorophyllide

Other name(s): CLH; Chlase

Systematic name: chlorophyll chlorophyllidohydrolase

**Comments:** Chlorophyllase has been found in higher plants, diatoms, and in the green algae *Chlorella* [2977].

This enzyme forms part of the chlorophyll degradation pathway and is thought to take part in degreening processes such as fruit ripening, leaf senescence and flowering, as well as in the turnover and homeostasis of chlorophyll [2187]. This enzyme acts preferentially on chlorophyll a but will also accept chlorophyll b and pheophytins as substrates [1180]. Ethylene and methyl jasmonate, which are known to accelerate senescence in many species, can enhance the activity of the hormone-inducible

form of this enzyme [1180].

**References:** [1162, 1473, 2977, 2187, 1180]

[EC 3.1.1.14 created 1961, modified 2007]

## EC 3.1.1.15

Accepted name: L-arabinonolactonase

**Reaction:** L-arabinono-1,4-lactone +  $H_2O$  = L-arabinonate **Systematic name:** L-arabinono-1,4-lactone lactonohydrolase

**References:** [3153]

[EC 3.1.1.15 created 1961]

[3.1.1.16 Deleted entry. 4-carboxymethyl-4-hydroxyisocrotonolactonase. This reaction was due to a mixture of EC 5.3.3.4 (muconolactone  $\Delta$ -isomerase) and EC 3.1.1.24 (3-oxoadipate enol-lactonase)]

[EC 3.1.1.16 created 1961, deleted 1972]

# EC 3.1.1.17

Accepted name: gluconolactonase

**Reaction:** D-glucono-1,5-lactone +  $H_2O$  = D-gluconate

**Other name(s):** lactonase; aldonolactonase; glucono-δ-lactonase; gulonolactonase

**Systematic name:** D-glucono-1,5-lactone lactonohydrolase

Comments: Acts on a wide range of hexose-1,5-lactones. The hydrolysis of L-gulono-1,5-lactone was previously

listed separately.

**References:** [298, 322, 2812]

[EC 3.1.1.17 created 1961 (EC 3.1.1.18 created 1961, incorporated 1982)]

[3.1.1.18 Deleted entry. aldonolactonase. Now included with EC 3.1.1.17 gluconolactonase]

[EC 3.1.1.18 created 1961, deleted 1982]

## EC 3.1.1.19

Accepted name: uronolactonase

**Reaction:** D-glucurono-6,2-lactone +  $H_2O$  = D-glucuronate

**Other name(s):** glucuronolactonase

**Systematic name:** D-glucurono-6,2-lactone lactonohydrolase

**References:** [3196]

[EC 3.1.1.19 created 1961]

EC 3.1.1.20

Accepted name: tannase

**Reaction:** digallate  $+ H_2O = 2$  gallate **Other name(s):** tannase S; tannin acetylhydrolase

**Systematic name:** tannin acylhydrolase

**Comments:** Also hydrolyses ester links in other tannins.

**References:** [672]

[EC 3.1.1.20 created 1961]

[3.1.1.21 Deleted entry. retinyl-palmitate esterase. Now known to be catalysed by EC 3.1.1.1, carboxylesterase and EC 3.1.1.3, triacylglycerol lipase.]

[EC 3.1.1.21 created 1972, deleted 2011]

EC 3.1.1.22

Accepted name: hydroxybutyrate-dimer hydrolase

**Reaction:** (*R*)-3-((*R*)-3-hydroxybutanoyloxy)butanoate +  $H_2O = 2$  (*R*)-3-hydroxybutanoate

**Other name(s):** D-(-)-3-hydroxybutyrate-dimer hydrolase

**Systematic name:** (R)-3-((R)-3-hydroxybutanoyloxy)butanoate hydroxybutanoylhydrolase

**References:** [573]

[EC 3.1.1.22 created 1972]

EC 3.1.1.23

Accepted name: acylglycerol lipase

**Reaction:** Hydrolyses glycerol monoesters of long-chain fatty acids

Other name(s): monoacylglycerol lipase; monoacylglycerolipase; monoglyceride lipase; monoglyceride hydrolase;

fatty acyl monoester lipase; monoacylglycerol hydrolase; monoglyceridyllipase; monoglyceridase

**Systematic name:** glycerol-ester acylhydrolase

**References:** [1887, 2311]

[EC 3.1.1.23 created 1972]

EC 3.1.1.24

Accepted name: 3-oxoadipate enol-lactonase

**Reaction:** 3-oxoadipate enol-lactone +  $H_2O = 3$ -oxoadipate

 $\label{eq:othername} \textbf{Other name}(s) : \quad \text{carboxymethylbutenolide lactonase}; \\ \beta\text{-ketoadipic enol-lactone hydrolase}; \\ 3\text{-ketoadipate enol-lactone}$ 

lactonase; 3-oxoadipic enol-lactone hydrolase; β-ketoadipate enol-lactone hydrolase

**Systematic name:** 4-carboxymethylbut-3-en-4-olide enol-lactonohydrolase

**Comments:** The enzyme acts on the product of EC 4.1.1.44 4-carboxymuconolactone decarboxylase.

**References:** [2202, 2203]

[EC 3.1.1.24 created 1961 as EC 3.1.1.16, part transferred 1972 to EC 3.1.1.24]

EC 3.1.1.25

**Accepted name:** 1,4-lactonase

**Reaction:** a 1,4-lactone +  $H_2O$  = a 4-hydroxyacid

Other name(s):  $\gamma$ -lactonase

**Systematic name:** 1,4-lactone hydroxyacylhydrolase

**Comments:** The enzyme is specific for 1,4-lactones with 4-8 carbon atoms. It does not hydrolyse simple aliphatic

esters, acetylcholine, sugar lactones or substituted aliphatic lactones, e.g. 3-hydroxy-4-butyrolactone;

requires Ca<sup>2+</sup>.

**References:** [775, 776]

# [EC 3.1.1.25 created 1972, modified 1981]

EC 3.1.1.26

Accepted name: galactolipase

**Reaction:** 1,2-diacyl-3-β-D-galactosyl-sn-glycerol + **2** H<sub>2</sub>O = 3-β-D-galactosyl-sn-glycerol + **2** carboxylates

Other name(s): galactolipid lipase; polygalactolipase; galactolipid acylhydrolase

**Systematic name:** 1,2-diacyl-3-β-D-galactosyl-*sn*-glycerol acylhydrolase

Comments: Also acts on 2,3-di-O-acyl-1-O-(6-O- $\alpha$ -D-galactosyl- $\beta$ -D-galactosyl)-D-glycerol, and phosphatidyl-

choline and other phospholipids.

**References:** [1108, 1150]

[EC 3.1.1.26 created 1972]

EC 3.1.1.27

Accepted name: 4-pyridoxolactonase

**Reaction:** 4-pyridoxolactone +  $H_2O = 4$ -pyridoxate **Systematic name:** 4-pyridoxolactone lactonohydrolase

**References:** [327]

[EC 3.1.1.27 created 1972]

EC 3.1.1.28

Accepted name: acylcarnitine hydrolase

**Reaction:** O-acylcarnitine +  $H_2O$  = a fatty acid + L-carnitine

**Other name(s):** high activity acylcarnitine hydrolase; HACH; carnitine ester hydrolase; palmitoylcarnitine hydrolase;

palmitoyl-L-carnitine hydrolase; long-chain acyl-L-carnitine hydrolase; palmitoyl carnitine hydrolase

**Systematic name:** *O*-acylcarnitine acylhydrolase

Comments: Acts on higher fatty acid ( $C_6$  to  $C_{18}$ ) esters of L-carnitine; highest activity is with O-decanoyl-L-

carnitine.

**References:** [1773, 1888]

[EC 3.1.1.28 created 1972]

EC 3.1.1.29

**Accepted name:** aminoacyl-tRNA hydrolase

**Reaction:** N-substituted aminoacyl-tRNA +  $H_2O = N$ -substituted amino acid + tRNA

**Other name(s):** aminoacyl-transfer ribonucleate hydrolase; *N*-substituted aminoacyl transfer RNA hydrolase;

peptidyl-tRNA hydrolase

Systematic name: aminoacyl-tRNA aminoacylhydrolase

**References:** [1342]

[EC 3.1.1.29 created 1972]

EC 3.1.1.30

Accepted name: D-arabinonolactonase

**Reaction:** D-arabinono-1,4-lactone +  $H_2O$  = D-arabinonate **Systematic name:** D-arabinono-1,4-lactone lactonohydrolase

**References:** [2227]

[EC 3.1.1.30 created 1972]

EC 3.1.1.31

**Accepted name:** 6-phosphogluconolactonase

**Reaction:** 6-phospho-D-glucono-1,5-lactone +  $H_2O$  = 6-phospho-D-gluconate

**Other name(s):** phosphogluconolactonase; 6-PGL

**Systematic name:** 6-phospho-D-glucono-1,5-lactone lactonohydrolase

**References:** [1405, 1906]

[EC 3.1.1.31 created 1972]

#### EC 3.1.1.32

**Accepted name:** phospholipase A<sub>1</sub>

**Reaction:** phosphatidylcholine +  $H_2O = 2$ -acylglycerophosphocholine + a carboxylate

**Systematic name:** phosphatidylcholine 1-acylhydrolase

**Comments:** This enzyme has a much broader specificity than EC 3.1.1.4 phospholipase  $A_2$ . Requires  $Ca^{2+}$ .

**References:** [883, 2549, 3032, 3034]

[EC 3.1.1.32 created 1972, modified 1976]

# EC 3.1.1.33

**Accepted name:** 6-acetylglucose deacetylase

**Reaction:** 6-acetyl-D-glucose +  $H_2O = D$ -glucose + acetate

Other name(s): 6-*O*-acetylglucose deacetylase
Systematic name: 6-acetyl-D-glucose acetylhydrolase

**References:** [663]

[EC 3.1.1.33 created 1972]

## EC 3.1.1.34

Accepted name: lipoprotein lipase

**Reaction:** triacylglycerol +  $H_2O$  = diacylglycerol + a carboxylate

Other name(s): clearing factor lipase; diglyceride lipase; diacylglycerol lipase; postheparin esterase; diglyceride li-

pase; postheparin lipase; diacylglycerol hydrolase; lipemia-clearing factor

Systematic name: triacylglycero-protein acylhydrolase

Comments: Hydrolyses triacylglycerols in chylomicrons and low-density lipoproteins. Also hydrolyses diacyl-

glycerol.

**References:** [681, 765, 977, 1981, 2089]

[EC 3.1.1.34 created 1972, modified 1976]

#### EC 3.1.1.35

Accepted name: dihydrocoumarin hydrolase

**Reaction:** dihydrocoumarin  $+ H_2O =$  melilotate **Systematic name:** dihydrocoumarin lactonohydrolase

**Comments:** Also hydrolyses some other benzenoid 1,4-lactones.

**References:** [1517]

[EC 3.1.1.35 created 1972]

## EC 3.1.1.36

Accepted name: limonin-D-ring-lactonase

**Reaction:** limonoate D-ring-lactone +  $H_2O$  = limonoate

Other name(s): limonin-D-ring-lactone hydrolase; limonin lactone hydrolase

**Systematic name:** limonoate-D-ring-lactone lactonohydrolase

**Comments:** Limonoate is a triterpenoid.

**References:** [1776]

[EC 3.1.1.36 created 1972]

EC 3.1.1.37

Accepted name: steroid-lactonase

**Reaction:** testololactone +  $H_2O$  = testolate **Systematic name:** testololactone lactonohydrolase

**References:** [1164]

[EC 3.1.1.37 created 1972]

EC 3.1.1.38

Accepted name: triacetate-lactonase

**Reaction:** triacetate lactone +  $H_2O$  = triacetate

Other name(s): triacetic lactone hydrolase; triacetic acid lactone hydrolase; TAL hydrolase; triacetate lactone hydro-

lase

**Systematic name:** triacetolactone lactonohydrolase

**References:** [1401]

[EC 3.1.1.38 created 1972]

EC 3.1.1.39

Accepted name: actinomycin lactonase

**Reaction:** actinomycin +  $H_2O$  = actinomycinic monolactone

Systematic name: actinomycin lactonohydrolase

**References:** [1186]

[EC 3.1.1.39 created 1972]

EC 3.1.1.40

**Accepted name:** orsellinate-depside hydrolase

**Reaction:** or sellinate depside  $+ H_2O = 2$  or sellinate

**Other name(s):** lecanorate hydrolase

**Systematic name:** orsellinate-depside hydrolase

**Comments:** The enzyme will only hydrolyse those substrates based on the 2,4-dihydroxy-6-methylbenzoate struc-

ture that also have a free hydroxy group ortho to the depside linkage.

**References:** [2588]

[EC 3.1.1.40 created 1976]

EC 3.1.1.41

Accepted name: cephalosporin-C deacetylase

**Reaction:** cephalosporin  $C + H_2O =$  deacetylcephalosporin C + acetate

**Other name(s):** cephalosporin C acetyl-hydrolase; cephalosporin C acetylase; cephalosporin acetylesterase;

cephalosporin C acetylesterase; cephalosporin C acetyl-esterase; cephalosporin C deacetylase

**Systematic name:** cephalosporin-C acetylhydrolase

**Comments:** Hydrolyses the acetyl ester bond on the 10-position of the antibiotic cephalosporin C.

**References:** [848]

[EC 3.1.1.41 created 1976]

#### EC 3.1.1.42

Accepted name: chlorogenate hydrolase

**Reaction:** chlorogenate +  $H_2O$  = caffeate + quinate **Other name(s):** chlorogenase; chlorogenic acid esterase

**Systematic name:** chlorogenate hydrolase

**Comments:** Also acts, more slowly, on isochlorogenate. No other substrates are known.

**References:** [2577, 2578]

[EC 3.1.1.42 created 1981]

#### EC 3.1.1.43

**Accepted name:** α-amino-acid esterase

**Reaction:** an  $\alpha$ -amino acid ester + H<sub>2</sub>O = an  $\alpha$ -amino acid + an alcohol

**Other name(s):**  $\alpha$ -amino acid ester hydrolase

**Systematic name:** α-amino-acid-ester aminoacylhydrolase

**Comments:** Also catalyses  $\alpha$ -aminoacyl transfer to a number of amine nucleophiles.

**References:** [1399, 1400, 2848]

[EC 3.1.1.43 created 1983]

#### EC 3.1.1.44

**Accepted name:** 4-methyloxaloacetate esterase

**Reaction:** oxaloacetate 4-methyl ester +  $H_2O$  = oxaloacetate + methanol

**Systematic name:** oxaloacetate-4-methyl-ester oxaloacetohydrolase

**References:** [639]

[EC 3.1.1.44 created 1983]

# EC 3.1.1.45

**Accepted name:** carboxymethylenebutenolidase

**Reaction:** 4-carboxymethylenebut-2-en-4-olide +  $H_2O$  = 4-oxohex-2-enedioate

Other name(s): maleylacetate enol-lactonase; dienelactone hydrolase; carboxymethylene butenolide hydrolase

**Systematic name:** 4-carboxymethylenebut-2-en-4-olide lactonohydrolase

**References:** [2572]

[EC 3.1.1.45 created 1983]

# EC 3.1.1.46

Accepted name: deoxylimonate A-ring-lactonase

**Reaction:** deoxylimonate  $+ H_2O =$  deoxylimononic acid D-ring-lactone

Systematic name: deoxylimonate A-ring-lactonohydrolase

**Comments:** The enzyme opens the A-ring-lactone of the triterpenoid deoxylimonic acid, leaving the D-ring-

lactone intact.

**References:** [1066]

[EC 3.1.1.46 created 1983]

# EC 3.1.1.47

**Accepted name:** 1-alkyl-2-acetylglycerophosphocholine esterase

**Reaction:** 1-alkyl-2-acetyl-sn-glycero-3-phosphocholine +  $H_2O$  = 1-alkyl-sn-glycero-3-phosphocholine + ac-

etate

**Other name(s):** 1-alkyl-2-acetyl-*sn*-glycero-3-phosphocholine acetylhydrolase; alkylacetyl-GPC:acetylhydrolase

**Systematic name:** 1-alkyl-2-acetyl-*sn*-glycero-3-phosphocholine acetohydrolase

References: [240]

[EC 3.1.1.47 created 1984]

EC 3.1.1.48

**Accepted name:** fusarinine-C ornithinesterase

**Reaction:**  $N^5$ -acyl-L-ornithine ester + H<sub>2</sub>O =  $N^5$ -acyl-L-ornithine + an alcohol

**Other name(s):** ornithine esterase; 5-*N*-acyl-L-ornithine-ester hydrolase

**Systematic name:**  $N^5$ -acyl-L-ornithine-ester hydrolase

**Comments:** Hydrolyses the three ornithine ester bonds in fusarinine C. Also acts on  $N^5$ -dinitrophenyl-L-ornithine

methyl ester.

**References:** [697]

[EC 3.1.1.48 created 1984]

EC 3.1.1.49

**Accepted name:** sinapine esterase

**Reaction:** sinapoylcholine +  $H_2O$  = sinapate + choline

**Other name(s):** aromatic choline esterase

**Systematic name:** sinapoylcholine sinapohydrolase

**References:** [2121]

[EC 3.1.1.49 created 1984]

EC 3.1.1.50

Accepted name: wax-ester hydrolase

**Reaction:** a wax ester  $+ H_2O = a$  long-chain alcohol + a long-chain carboxylate

Other name(s): jojoba wax esterase; WEH
Systematic name: wax-ester acylhydrolase

**Comments:** Also acts on long-chain acylglycerol, but not diacyl- or triacylglycerols.

**References:** [1201, 1963]

[EC 3.1.1.50 created 1984]

EC 3.1.1.51

Accepted name: phorbol-diester hydrolase

**Reaction:** phorbol 12,13-dibutanoate +  $H_2O$  = phorbol 13-butanoate + butanoate

Other name(s): diacylphorbate 12-hydrolase; diacylphorbate 12-hydrolase; phorbol-12,13-diester 12-ester hydrolase;

**PDEH** 

**Systematic name:** 12,13-diacylphorbate 12-acylhydrolase

**Comments:** Hydrolyses the 12-ester bond in a variety of 12,13-diacylphorbols (phorbol is a diterpenoid); this re-

action inactivates the tumour promotor 12-O-tetradecanoylphorbol-13-acetate from croton oil.

**References:** [2647]

[EC 3.1.1.51 created 1984]

EC 3.1.1.52

Accepted name: phosphatidylinositol deacylase

**Reaction:** 1-phosphatidyl-D-myo-inositol +  $H_2O = 1$ -acylglycerophosphoinositol + a carboxylate

**Other name(s):** phosphatidylinositol phospholipase A<sub>2</sub>; phospholipase A<sub>2</sub>

**Systematic name:** 1-phosphatidyl-D-*myo*-inositol 2-acylhydrolase

**References:** [970, 969]

# [EC 3.1.1.52 created 1984]

EC 3.1.1.53

**Accepted name:** sialate *O*-acetylesterase

**Reaction:** N-acetyl-O-acetylneuraminate +  $H_2O = N$ -acetylneuraminate + acetate **Other name(s):** N-acetylneuraminate acetyltransferase; sialate 9(4)-O-acetylesterase; sialidase

**Systematic name:** *N*-acyl-*O*-acetylneuraminate *O*-acetylhydrolase

**Comments:** Acts on free and glycosidically bound N-acetyl- or N-glycoloyl-neuraminic acid; acts mainly on the

4-O- and 9-O-acetyl groups. Also acts on some other O-acetyl esters, both cyclic and acyclic com-

pounds, which are not sialic acids.

**References:** [877, 2648]

[EC 3.1.1.53 created 1984]

EC 3.1.1.54

Accepted name: acetoxybutynylbithiophene deacetylase

**Reaction:** 5-(4-acetoxybut-1-ynyl)-2,2'-bithiophene + H<sub>2</sub>O = <math>5-(4-hydroxybut-1-ynyl)-2,2'-bithiophene + ac-bithiophene +

etate

**Other name(s):** acetoxybutynylbithiophene esterase; 5-(4-acetoxy-1-butynyl)-2,2'-bithiophene:acetate esterase

**Systematic name:** 5-(4-acetoxybut-1-ynyl)-2,2'-bithiophene *O*-acetylhydrolase

**Comments:** The enzyme is highly specific.

**References:** [2807]

[EC 3.1.1.54 created 1986]

EC 3.1.1.55

Accepted name: acetylsalicylate deacetylase

**Reaction:** acetylsalicylate +  $H_2O$  = salicylate + acetate

Other name(s): aspirin esterase; aspirin esterase; acetylsalicylic acid esterase; aspirin hydrolase

**Systematic name:** acetylsalicylate *O*-acetylhydrolase

Comments: Not identical with EC 3.1.1.1 (carboxylesterase), EC 3.1.1.2 (arylesterase), EC 3.1.1.7 (acetyl-

cholinesterase) or EC 3.1.1.8 (cholinesterase). The activity of the liver cytosol enzyme is highest with acetyl esters of aryl alcohols, and thioesters are also hydrolysed; the microsomal enzyme also hydrolyses some other negatively charged esters, with highest activity on esters of salicylate with long-chain

alcohols.

**References:** [35, 1444, 3173]

[EC 3.1.1.55 created 1986, modified 1989]

EC 3.1.1.56

Accepted name: methylumbelliferyl-acetate deacetylase

**Reaction:** 4-methylumbelliferyl acetate  $+ H_2O = 4$ -methylumbelliferone + acetate

**Other name(s):** esterase D

**Systematic name:** 4-methylumbelliferyl-acetate acylhydrolase

Comments: Acts on short-chain acyl esters of 4-methylumbelliferone, but not on naphthyl, indoxyl or thiocholine

esters.

**References:** [1174]

[EC 3.1.1.56 created 1986]

EC 3.1.1.57

**Accepted name:** 2-pyrone-4,6-dicarboxylate lactonase

**Reaction:** 2-oxo-2H-pyran-4,6-dicarboxylate +  $H_2O = (1E)$ -4-oxobut-1-ene-1,2,4-tricarboxylate **Other name(s):** 2-pyrone-4,6-dicarboxylate hydrolase; 2-pyrone-4,6-dicarboxylate lactonohydrolase

**Systematic name:** 2-oxo-2*H*-pyran-4,6-dicarboxylate lactonohydrolase

**Comments:** The product is most likely the keto-form of 4-oxalomesaconate (as shown in the reaction) [1422,

1827]. It can be converted to the enol-form, 4-hydroxybuta-1,3-diene-1,2,4-trioate, either sponta-

neously or by EC 5.3.2.8, 4-oxalomesaconate tautomerase [2103].

**References:** [1422, 1827, 2103]

[EC 3.1.1.57 created 1986, modified 2010]

EC 3.1.1.58

Accepted name: N-acetylgalactosaminoglycan deacetylase

**Reaction:** N-acetyl-D-galactosaminoglycan +  $H_2O$  = D-galactosaminoglycan + acetate

Other name(s): polysaccharide deacetylase (misleading); Vi-polysaccharide deacetylase; N-acetyl galactosaminogly-

can deacetylase

**Systematic name:** *N*-acetyl-D-galactosaminoglycan acetylhydrolase

**References:** [1339]

[EC 3.1.1.58 created 1986]

EC 3.1.1.59

Accepted name: juvenile-hormone esterase

**Reaction:** (1) juvenile hormone  $I + H_2O =$  juvenile hormone I acid + methanol

(2) juvenile hormone III +  $H_2O$  = juvenile hormone III acid + methanol

Other name(s): JH-esterase; juvenile hormone analog esterase; juvenile hormone carboxyesterase; methyl-(2E,6E)-

(10R,11S)-10,11-epoxy-3,7,11-trimethyltrideca-2,6-dienoate acylhydrolase

**Systematic name:** methyl-(2*E*,6*E*,10*R*)-10,11-epoxy-3,7,11-trimethyltrideca-2,6-dienoate acylhydrolase

Comments: Demethylates the insect juvenile hormones JH1 and JH3, but does not hydrolyse the analogous ethyl

or isopropyl esters.

**References:** [558, 1932]

[EC 3.1.1.59 created 1989, modified 2015]

EC 3.1.1.60

**Accepted name:** bis(2-ethylhexyl)phthalate esterase

**Reaction:** bis(2-ethylhexyl)phthalate +  $H_2O = 2$ -ethylhexyl phthalate + 2-ethylhexan-1-ol

**Other name(s):** DEHP esterase

**Systematic name:** bis(2-ethylhexyl)phthalate acylhydrolase

 $\textbf{Comments:} \quad \text{Also acts on 4-nitrophenyl esters, with optimum chain-length $C_6$ to $C_8$.}$ 

**References:** [1011]

[EC 3.1.1.60 created 1989]

EC 3.1.1.61

**Accepted name:** protein-glutamate methylesterase

**Reaction:** protein L-glutamate  $O^5$ -methyl ester +  $H_2O$  = protein L-glutamate + methanol

Other name(s): chemotaxis-specific methylesterase; methyl-accepting chemotaxis protein methyl-esterase; CheB

methylesterase; methylesterase CheB; protein methyl-esterase; protein carboxyl methylesterase;

PME; protein methylesterase; protein-L-glutamate-5-O-methyl-ester acylhydrolase

**Systematic name:** protein-L-glutamate- $O^5$ -methyl-ester acylhydrolase

**Comments:** Hydrolyses the products of EC 2.1.1.77 (protein-L-isoaspartate(D-aspartate) *O*-methyltransferase), EC

2.1.1.78 (isoorientin 3'-O-methyltransferase), EC 2.1.1.80 (protein-glutamate O-methyltransferase)

and EC 2.1.1.100 (protein-S-isoprenylcysteine O-methyltransferase).

**References:** [868, 1411]

# [EC 3.1.1.61 created 1989, modified 2002]

[3.1.1.62 Deleted entry. N-acetyldiaminopimelate deacylase. Now listed as EC 3.5.1.47, N-acetyldiaminopimelate deacetylase]

[EC 3.1.1.62 created 1989, deleted 1992]

#### EC 3.1.1.63

**Accepted name:** 11-cis-retinyl-palmitate hydrolase

**Reaction:** 11-cis-retinyl palmitate +  $H_2O = 11$ -cis-retinol + palmitate

Other name(s): 11-*cis*-retinol palmitate esterase; RPH Systematic name: 11-*cis*-retinyl-palmitate acylhydrolase

**Comments:** Activated by bile salts.

**References:** [238, 239]

[EC 3.1.1.63 created 1989]

#### EC 3.1.1.64

Accepted name: retinoid isomerohydrolase

**Reaction:** an *all-trans*-retinyl ester +  $H_2O = 11$ -*cis*-retinol + a fatty acid

Other name(s): all-trans-retinyl-palmitate hydrolase (ambiguous); retinol isomerase (ambiguous); all-trans-retinol

isomerase:hydrolase (ambiguous); all-trans-retinylester 11-cis isomerohydrolase; RPE65 (gene name)

**Systematic name:** *all-trans*-retinyl ester acylhydrolase, 11-*cis* retinol forming

Comments: This enzyme, which operates in the retinal pigment epithelium (RPE), catalyses the cleavage and iso-

merization of *all-trans*-retinyl fatty acid esters to 11-*cis*-retinol, a key step in the regeneration of the visual chromophore in the vertebrate visual cycle [1946]. Interaction of the enzyme with the mem-

brane is critical for its enzymic activity [942].

**References:** [238, 203, 288, 1946, 2086, 942]

[EC 3.1.1.64 created 1989 (EC 5.2.1.7 created 1989, incorporated 2011), modified 2011]

# EC 3.1.1.65

Accepted name: L-rhamnono-1,4-lactonase

**Reaction:** L-rhamnono-1,4-lactone +  $H_2O$  = L-rhamnonate

**Other name(s):** L-rhamno-γ-lactonase; L-rhamnono-γ-lactonase; L-rhamnonate dehydratase

**Systematic name:** L-rhamnono-1,4-lactone lactonohydrolase

**References:** [2426]

[EC 3.1.1.65 created 1989]

#### EC 3.1.1.66

**Accepted name:** 5-(3,4-diacetoxybut-1-ynyl)-2,2'-bithiophene deacetylase

**Reaction:**  $5-(3,4-\text{diacetoxybut-1-ynyl})-2,2'-\text{bithiophene} + H_2O = 5-(3-\text{hydroxy-4-acetoxybut-1-ynyl})-2,2'-$ 

bithiophene + acetate

**Other name(s):** diacetoxybutynylbithiophene acetate esterase; 3,4-diacetoxybutinylbithiophene:4-acetate esterase

**Systematic name:** 5-(3,4-diacetoxybut-1-ynyl)-2,2'-bithiophene acetylhydrolase

**Comments:** A highly specific enzyme from *Tagetes patula*.

**References:** [2256]

[EC 3.1.1.66 created 1989]

#### EC 3.1.1.67

Accepted name: fatty-acyl-ethyl-ester synthase

**Reaction:** a long-chain-fatty-acyl ethyl ester +  $H_2O$  = a long-chain-fatty acid + ethanol

Other name(s): FAEES

**Systematic name:** long-chain-fatty-acyl-ethyl-ester acylhydrolase

**Comments:** The reaction, forms ethyl esters from fatty acids and ethanol in the absence of coenzyme A or ATP.

Best substrates are unsaturated octadecanoic acids; palmitate, stearate and arachidonate also act, but

more slowly.

**References:** [1945]

[EC 3.1.1.67 created 1989]

#### EC 3.1.1.68

**Accepted name:** xylono-1,4-lactonase

**Reaction:** D-xylono-1,4-lactone +  $H_2O$  = D-xylonate **Other name(s):** xylono- $\gamma$ -lactonase; xylonolactonase **Systematic name:** D-xylono-1,4-lactone lactonohydrolase

References: [324]

[EC 3.1.1.68 created 1990]

[3.1.1.69 Transferred entry. N-acetylglucosaminylphosphatidylinositol deacetylase. Now EC 3.5.1.89, N-acetylglucosaminylphosphatid deacetylase. Previously classified erroneously as an enzyme that hydrolysed an ester and not an amide]

[EC 3.1.1.69 created 1992, deleted 2002]

#### EC 3.1.1.70

Accepted name: cetraxate benzylesterase

**Reaction:** cetraxate benzyl ester +  $H_2O$  = cetraxate + benzyl alcohol

**Systematic name:** cetraxate-benzyl-ester benzylhydrolase

Comments: Acts on a number of benzyl esters of substituted phenyl propanoates, and on the benzyl esters of

phenylalanine and tyrosine.

**References:** [1571]

[EC 3.1.1.70 created 1992]

# EC 3.1.1.71

Accepted name: acetylalkylglycerol acetylhydrolase

**Reaction:** 2-acetyl-1-alkyl-sn-glycerol + H<sub>2</sub>O = 1-alkyl-sn-glycerol + acetate

**Other name(s):** alkylacetylglycerol acetylhydrolase

**Systematic name:** 2-acetyl-1-alkyl-*sn*-glycerol acetylhydrolase

Comments: Hydrolysis of the acetyl group from the 1-alkyl-2-acetyl and 1-alkyl-3-acetyl substrates occurs at

apparently identical rates. The enzyme from Erlich ascites cells is membrane-bound. It differs from lipoprotein lipase (EC 3.1.1.34) since 1,2-diacetyl-sn-glycerols are not substrates. It also differs from

EC 3.1.1.47, 1-acetyl-2-alkyl-glycerophosphocholine esterase.

**References:** [241]

[EC 3.1.1.71 created 1999]

#### EC 3.1.1.72

**Accepted name:** acetylxylan esterase

**Reaction:** Deacetylation of xylans and xylo-oligosaccharides

**Systematic name:** acetylxylan esterase

**Comments:** Catalyses the hydrolysis of acetyl groups from polymeric xylan, acetylated xylose, acetylated glucose,

 $\alpha$ -napthyl acetate, p-nitrophenyl acetate but not from triacetylglycerol. Does not act on acetylated

mannan or pectin.

**References:** [2802, 2319, 1811]

[EC 3.1.1.72 created 1999]

EC 3.1.1.73

Accepted name: feruloyl esterase

**Reaction:** feruloyl-polysaccharide +  $H_2O$  = ferulate + polysaccharide

Other name(s): ferulic acid esterase, hydroxycinnamoyl esterase, hemicellulase accessory enzymes; FAE-III, cin-

namoyl ester hydrolase, FAEA, cinnAE, FAE-I, FAE-II

**Systematic name:** 4-hydroxy-3-methoxycinnamoyl-sugar hydrolase

**Comments:** Catalyses the hydrolysis of the 4-hydroxy-3-methoxycinnamoyl (feruloyl) group from an esterified

sugar, which is usually arabinose in "natural" substrates. *p*-Nitrophenol acetate and methyl ferulate are poorer substrates. All microbial ferulate esterases are secreted into the culture medium. They are sometimes called hemicellulase accessory enzymes, since they help xylanases and pectinases to break

down plant cell wall hemicellulose.

**References:** [745, 746, 1538, 597, 379]

[EC 3.1.1.73 created 2000]

EC 3.1.1.74

Accepted name: cutinase

**Reaction:**  $cutin + H_2O = cutin monomers$ 

Systematic name: cutin hydrolase

**Comments:** Cutin, a polymeric structural component of plant cuticles, is a polymer of hydroxy fatty acids that are

usually  $C_{16}$  or  $C_{18}$  and contain up to three hydroxy groups. The enzyme from several fungal sources also hydrolyses the p-nitrophenyl esters of hexadecanoic acid. It is however inactive towards several

esters that are substrates for non-specific esterases.

**References:** [876, 2331, 2330]

[EC 3.1.1.74 created 2000]

EC 3.1.1.75

**Accepted name:** poly(3-hydroxybutyrate) depolymerase

**Reaction:**  $[(R)-3-hydroxybutanoate]_n + H_2O = [(R)-3-hydroxybutanoate]_{n-x} + [(R)-3-hydroxybutanoate]_x; x =$ 

1–5

**Other name(s):** PHB depolymerase; poly(3HB) depolymerase; poly[(R)-hydroxyalkanoic acid] depolymerase;

poly(HA) depolymerase; poly(HA<sub>SCL</sub>) depolymerase; poly[(R)-3-hydroxybutyrate] hydrolase

**Systematic name:** poly[(R)-3-hydroxybutanoate] hydrolase

**Comments:** Reaction also occurs with esters of other short-chain-length  $(C_1-C_5)$  hydroxyalkanoic acids (HA).

There are two types of polymers: native (intracellular) granules are amorphous and have an intact surface layer; denatured (extracellular) granules either have no surface layer or a damaged surface layer

and are partially crystalline.

**References:** [1316, 874]

[EC 3.1.1.75 created 2001]

EC 3.1.1.76

**Accepted name:** poly(3-hydroxyoctanoate) depolymerase

**Reaction:** Hydrolyses the polyester polyoxycarbonyl[(R)-2-pentylethylene] to oligomers

**Other name(s):** PHO depolymerase; poly(3HO) depolymerase; poly[(R)-hydroxyalkanoic acid] depolymerase;

poly(HA) depolymerase; poly(HA $_{MCL}$ ) depolymerase; poly[(R)-3-hydroxyoctanoate] hydrolase

**Systematic name:** polyoxycarbonyl[(R)-2-pentylethylene] hydrolase

**Comments:** The main product after prolonged incubation is the dimer [2566]. Besides hydrolysing polymers of 3-

hydroxyoctanoic acid, the enzyme also hydrolyses other polymers derived from medium-chain-length  $(C_6-C_{12})$  hydroxyalkanoic acids and copolymers of mixtures of these. It also hydrolyses p-nitrophenyl esters of fatty acids. Polymers of short-chain-length hydroxyalkanoic acids such as poly[(R)-3-

hydroxybutanoic acid] and poly[(R)-3-hydroxypentanoic acid] are not hydrolysed.

**References:** [1316, 874, 2566]

[EC 3.1.1.76 created 2001, modified 2005]

EC 3.1.1.77

**Accepted name:** acyloxyacyl hydrolase

**Reaction:** 3-(acyloxy)acyl group of bacterial toxin +  $H_2O = 3$ -hydroxyacyl group of bacterial toxin + a fatty acid **Comments:** The substrate is lipid A on the reducing end of the toxic lipopolysaccharide (LPS) of *Salmonella ty*-

*phimurium* and related organisms. It consists of diglucosamine, β-D-GlcN-(1 $\rightarrow$  6)-D-GlcN, attached by glycosylation on O-6 of its non-reducing residue, phosphorylated on O-4 of this residue and on O-1 of its potentially reducing residue. Both residues carry 3-(acyloxy)acyl groups on *N*-2 and O-3. The enzyme from human leucocytes detoxifies the lipid by hydrolysing the secondary acyl groups from O-3 of the 3-hydroxyacyl groups on the disaccharide (LPS). It also possesses a wide range of phospholipase and acyltransferase activities [e.g. EC 3.1.1.4 (phospholipase A<sub>2</sub>), EC 3.1.1.5 (lysophospholipase), EC 3.1.1.32 (phospholipase A<sub>1</sub>) and EC 3.1.1.52 (phosphatidylinositol deacylase)], hydrolysing diacylglycerol and phosphatidyl compounds, but not triacylglycerols. It has a preference for saturated

 $C_{12}$ - $C_{16}$  acyl groups.

**References:** [715, 1021, 2002]

[EC 3.1.1.77 created 2001]

EC 3.1.1.78

**Accepted name:** polyneuridine-aldehyde esterase

**Reaction:** polyneuridine aldehyde +  $H_2O = 16$ -epivellosimine +  $CO_2$  + methanol

**Other name(s):** polyneuridine aldehyde esterase; PNAE

**Systematic name:** polyneuridine aldehyde hydrolase (decarboxylating)

**Comments:** Following hydrolysis of this indole alkaloid ester the carboxylic acid decarboxylates spontaneously

giving the sarpagan skeleton. The enzyme also acts on akuammidine aldehyde (the 16-epimer of

polyneuridine aldehyde).

**References:** [2270, 2271, 629, 1845]

[EC 3.1.1.78 created 2002]

EC 3.1.1.79

**Accepted name:** hormone-sensitive lipase

**Reaction:** (1) diacylglycerol +  $H_2O$  = monoacylglycerol + a carboxylate

(2) triacylglycerol +  $H_2O$  = diacylglycerol + a carboxylate

(3) monoacylglycerol +  $H_2O$  = glycerol + a carboxylate

Other name(s): HSL

**Systematic name:** diacylglycerol acylhydrolase

**Comments:** This enzyme is a serine hydrolase. Compared with other lipases, hormone-sensitive lipase has a

uniquely broad substrate specificity. It hydrolyses all acylglycerols (triacylglycerol, diacylglycerol and monoacylglycerol) [2,3,4] as well as cholesteryl esters [804, 2207], steroid fatty acid esters [1623], retinyl esters [3148] and *p*-nitrophenyl esters [2207, 2981]. It exhibits a preference for the 1- or 3-ester bond of its acylglycerol substrate compared with the 2-ester bond [3282]. The enzyme shows little preference for the fatty acids in the triacylglycerol, although there is some increase in activity with decreasing chain length. The enzyme activity is increased in response to hormones that

elevate intracellular levels of cAMP.

**References:** [1163, 804, 3056, 2207, 1623, 3148, 2981, 3282]

EC 3.1.1.80

Accepted name: acetylajmaline esterase

**Reaction:** (1) 17-*O*-acetylajmaline +  $H_2O$  = ajmaline + acetate

(2) 17-O-acetylnorajmaline +  $H_2O$  = norajmaline + acetate

Other name(s): AAE;  $2\beta(R)$ -17-O-acetylajmalan:acetylesterase; acetylajmalan esterase

**Systematic name:** 17-*O*-acetylajmaline *O*-acetylhydrolase

**Comments:** This plant enzyme is responsible for the last stages in the biosynthesis of the indole alkaloid ajmaline.

The enzyme is highly specific for the substrates 17-*O*-acetylajmaline and 17-*O*-acetylnorajmaline as the structurally related acetylated alkaloids vinorine, vomilenine, 1,2-dihydrovomilenine and 1,2-dihydroraucaffricine cannot act as substrates [2480]. This is a novel member of the GDSL family of

serine esterases/lipases.

**References:** [2308, 2480]

[EC 3.1.1.80 created 2006]

EC 3.1.1.81

**Accepted name:** quorum-quenching *N*-acyl-homoserine lactonase

**Reaction:** an *N*-acyl-L-homoserine lactone +  $H_2O$  = an *N*-acyl-L-homoserine

Other name(s): acyl homoserine degrading enzyme; acyl-homoserine lactone acylase; AHL lactonase; AHL-

degrading enzyme; AHL-inactivating enzyme; AHLase; AhlD; AhlK; AiiA; AiiA lactonase; AiiA-like protein; AiiB; AiiC; AttM; delactonase; lactonase-like enzyme; *N*-acyl homoserine lactonase; *N*-acyl-homoserine lactone lactonase; *N*-acyl-homoserine lactone hydrolase; quorum-quenching lactonase; quorum-quenching *N*-acyl homoserine lactone hydrolase; quorum-quenching lactonase; quorum-quenching *N*-acyl homoserine lactone hydrolase;

drolase

**Systematic name:** *N*-acyl-L-homoserine-lactone lactonohydrolase

**Comments:** Acyl-homoserine lactones (AHLs) are produced by a number of bacterial species and are used by

them to regulate the expression of virulence genes in a process known as quorum-sensing. Each bacterial cell has a basal level of AHL and, once the population density reaches a critical level, it triggers AHL-signalling which, in turn, initiates the expression of particular virulence genes [637]. Plants or animals capable of degrading AHLs would have a therapeutic advantage in avoiding bacterial infection as they could prevent AHL-signalling and the expression of virulence genes in quorum-sensing bacteria [637]. *N*-(3-Oxohexanoyl)-L-homoserine lactone, *N*-(3-oxododecanoyl)-L-homoserine lactone, *N*-butanoyl-L-homoserine lactone and *N*-(3-oxooctanoyl)-L-homoserine lactone can act as sub-

strates [637].

**References:** [2915, 636, 3116, 638, 637, 1630, 2236, 3008, 1453, 1718, 3258]

[EC 3.1.1.81 created 2007]

EC 3.1.1.82

**Accepted name:** pheophorbidase

**Reaction:** pheophorbide  $a + H_2O = \text{pyropheophorbide } a + \text{methanol} + \text{CO}_2 \text{ (overall reaction)}$ 

(1a) pheophorbide  $a + H_2O = C-13^2$ -carboxypyropheophorbide a + methanol

(1b) C-13<sup>2</sup>-carboxypyropheophorbide a = pyropheophorbide a + CO<sub>2</sub> (spontaneous)

Other name(s): phedase; PPD

**Systematic name:** pheophorbide-*a* hydrolase

**Comments:** This enzyme forms part of the chlorophyll degradation pathway, and is found in higher plants and in

> algae. In higher plants it participates in de-greening processes such as fruit ripening, leaf senescence, and flowering. The enzyme exists in two forms: type 1 is induced by senescence whereas type 2 is constitutively expressed [2817, 2815]. The enzyme is highly specific for pheophorbide as substrate (with a preference for pheophorbide a over pheophorbide b) as other chlorophyll derivatives such as protochlorophyllide a, pheophytin a and c, chlorophyll a and b, and chlorophyllide a cannot act as substrates [2815]. Another enzyme, called pheophorbide demethoxycarbonylase (PDC), produces pyropheophorbide a from pheophorbide a without forming an intermediate although the precise reaction is not yet known [2817].

**References:** [2817, 2815, 1180]

[EC 3.1.1.82 created 2007]

EC 3.1.1.83

Accepted name: monoterpene ε-lactone hydrolase

> Reaction: (1) isoprop(en)ylmethyloxepan-2-one + H<sub>2</sub>O = 6-hydroxyisoprop(en)ylmethylhexanoate (general re-

> > action)

(2) 4-isopropenyl-7-methyloxepan-2-one +  $H_2O$  = 6-hydroxy-3-isopropenylheptanoate

(3) 7-isopropyl-4-methyloxepan-2-one +  $H_2O$  = 6-hydroxy-3,7-dimethyloctanoate

Other name(s):

**Systematic name:** isoprop(en)ylmethyloxepan-2-one lactonohydrolase

**Comments:** The enzyme catalyses the ring opening of \varepsilon-lactones which are formed during degradation of dihydro-

carveol by the Gram-positive bacterium Rhodococcus erythropolis DCL14. The enzyme also acts on ethyl caproate, indicating that it is an esterase with a preference for lactones (internal cyclic esters).

The enzyme is not stereoselective.

**References:** [3040]

[EC 3.1.1.83 created 2008]

EC 3.1.1.84

Accepted name: cocaine esterase

cocaine +  $H_2O$  = ecgonine methyl ester + benzoate **Reaction:** Other name(s): CocE; hCE2; hCE-2; human carboxylesterase 2

**Systematic name:** cocaine benzoylhydrolase

> **Comments:** Rhodococcus sp. strain MB1 and Pseudomonas maltophilia strain MB11L can utilize cocaine as sole

> > source of carbon and energy [287, 294].

[871, 287, 294, 1605, 2285] **References:** 

[EC 3.1.1.84 created 2010]

EC 3.1.1.85

Accepted name: pimelyl-[acyl-carrier protein] methyl ester esterase

pimeloyl-[acyl-carrier protein] methyl ester +  $H_2O$  = pimeloyl-[acyl-carrier protein] + methanol **Reaction:** 

Other name(s): **BioH** 

**Systematic name:** pimeloyl-[acyl-carrier protein] methyl ester hydrolase

**Comments:** Involved in biotin biosynthesis in Gram-negative bacteria. The enzyme exhibits carboxylesterase ac-

tivity, particularly toward substrates with short acyl chains [2518, 1644]. Even though the enzyme can interact with coenzyme A thioesters [2933], the in vivo role of the enzyme is to hydrolyse the methyl ester of pimeloyl-[acyl carrier protein], terminating the part of the biotin biosynthesis pathway that is

catalysed by the fatty acid elongation enzymes [1693].

References: [2518, 1644, 2933, 1693]

[EC 3.1.1.85 created 2011]

#### EC 3.1.1.86

Accepted name: rhamnogalacturonan acetylesterase

**Reaction:** Hydrolytic cleavage of 2-O-acetyl- or 3-O-acetyl groups of α-D-galacturonic acid in rhamnogalactur-

onan I.

**Other name(s):** RGAE

**Systematic name:** rhamnogalacturonan 2/3-O-acetyl-α-D-galacturonate O-acetylhydrolase

**Comments:** The degradation of rhamnogalacturonan by rhamnogalacturonases depends on the removal of the

acetyl esters from the substrate [1404].

**References:** [1404, 1950]

[EC 3.1.1.86 created 2011]

# EC 3.1.1.87

Accepted name: fumonisin B1 esterase

**Reaction:** fumonisin B1 + 2  $H_2O$  = aminopentol + 2 propane-1,2,3-tricarboxylate

**Other name(s):** *fumD* (gene name)

**Systematic name:** fumonisin B1 acylhydrolase

**Comments:** The enzyme is involved in degradation of fumonisin B1 [1103].

**References:** [1103]

[EC 3.1.1.87 created 2011]

#### EC 3.1.1.88

**Accepted name:** pyrethroid hydrolase

**Reaction:** trans-permethrin + H<sub>2</sub>O = (3-phenoxyphenyl)methanol + (1S,3R)-3-(2,2-dichloroethenyl)-2,2-

dimethylcyclopropanecarboxylate

**Other name(s):** pyrethroid-hydrolyzing carboxylesterase; pyrethroid-hydrolysing esterase; pyrethroid-hydrolyzing

esterase; pyrethroid-selective esterase; pyrethroid-cleaving enzyme; permethrinase; PytH; EstP

**Systematic name:** pyrethroid-ester hydrolase

**Comments:** The enzyme is involved in degradation of pyrethroid pesticides. The enzymes from *Sphingobium* sp.,

*Klebsiella* sp. and *Aspergillus niger* hydrolyse *cis*-permethrin at approximately equal rate to *trans*-permethrin [3105, 3220, 1677]. The enzyme from mouse hydrolyses *trans*-permethrin at a rate about

22-fold higher than *cis*-permethrin [2778].

**References:** [3105, 3220, 1677, 2778, 1794, 1001]

[EC 3.1.1.88 created 2011]

# EC 3.1.1.89

**Accepted name:** protein phosphatase methylesterase-1

**Reaction:** [phosphatase 2A protein]-leucine methyl ester +  $H_2O$  = [phosphatase 2A protein]-leucine + methanol

**Other name**(s): PME-1; PPME1

**Systematic name:** [phosphatase 2A protein]-leucine ester acylhydrolase

Comments: A key regulator of protein phosphatase 2A. The methyl ester is formed by EC 2.1.1.233 (leucine car-

boxy methyltransferase-1). Occurs mainly in the nucleus.

**References:** [2153, 3229]

[EC 3.1.1.89 created 2011]

## EC 3.1.1.90

**Accepted name:** all-trans-retinyl ester 13-cis isomerohydrolase

**Reaction:** an *all-trans*-retinyl ester  $+ H_2O = 13$ -*cis*-retinol + a fatty acid **Systematic name:** *all-trans*-retinyl ester acylhydrolase, 13-*cis* retinol forming

Comments: All-trans-retinyl esters, which are a storage form of vitamin A, are generated by the activity of EC

2.3.1.135, phosphatidylcholine—retinol *O*-acyltransferase (LRAT). They can be hydrolysed to 11-*cis*-

retinol by EC 3.1.1.64, retinoid isomerohydrolase (RPE65), or to 13-cis-retinol by this enzyme.

**References:** [2849]

[EC 3.1.1.90 created 2011]

EC 3.1.1.91

**Accepted name:** 2-oxo-3-(5-oxofuran-2-ylidene)propanoate lactonase

**Reaction:** 2-oxo-3-(5-oxofuran-2-ylidene)propanoate +  $H_2O$  = maleylpyruvate

**Other name(s):** *naaC* (gene name)

**Systematic name:** 2-oxo-3-(5-oxofuran-2-ylidene)propanoate lactonohydrolase

Comments: This enzyme, characterized from the soil bacterium *Bradyrhizobium* sp. JS329, is involved in the

pathway of 5-nitroanthranilate degradation.

**References:** [2340]

[EC 3.1.1.91 created 2012]

EC 3.1.1.92

**Accepted name:** 4-sulfomuconolactone hydrolase

**Reaction:** 4-sulfomuconolactone +  $H_2O$  = maleylacetate + sulfite

**Systematic name:** 4-sulfomuconolactone sulfohydrolase

Comments: The enzyme was isolated from the bacteria Hydrogenophaga intermedia and Agrobacterium ra-

diobacter S2. It catalyses a step in the degradation of 4-sulfocatechol.

**References:** [1024]

[EC 3.1.1.92 created 2012]

EC 3.1.1.93

**Accepted name:** mycophenolic acid acyl-glucuronide esterase

**Reaction:** mycophenolic acid O-acyl-glucuronide +  $H_2O$  = mycophenolate + D-glucuronate **Other name(s):** mycophenolic acid acyl-glucuronide deglucuronidase; AcMPAG deglucuronidase

**Systematic name:** mycophenolic acid *O*-acyl-glucuronide-ester hydrolase

**Comments:** This liver enzyme deglucuronidates mycophenolic acid *O*-acyl-glucuronide, a metabolite of the im-

munosuppressant drug mycophenolate that is thought to be immunotoxic.

**References:** [1282]

[EC 3.1.1.93 created 2012]

EC 3.1.1.94

Accepted name: versiconal hemiacetal acetate esterase

**Reaction:** (1) versiconal hemiacetal acetate  $+ H_2O = versiconal + acetate$ 

(2) versiconol acetate +  $H_2O$  = versiconol + acetate

Other name(s): VHA esterase

**Systematic name:** versiconal-hemiacetal-acetate *O*-acetylhydrolase

**Comments:** Isolated from the mold *Aspergillus parasiticus*. Involved in a metabolic grid that leads to aflatoxin

biosynthesis.

**References:** [1577, 396]

[EC 3.1.1.94 created 2013]

EC 3.1.1.95

**Accepted name:** aclacinomycin methylesterase

**Reaction:** aclacinomycin T +  $H_2O = 15$ -demethylaclacinomycin T + methanol

Other name(s): RdmC; aclacinomycin methyl esterase Systematic name: aclacinomycin T acylhydrolase

**Comments:** The enzyme is involved in the modification of the aklavinone skeleton in the biosynthesis of anthracy-

clines in Streptomyces species.

**References:** [3123, 1309]

[EC 3.1.1.95 created 2013]

EC 3.1.1.96

Accepted name: D-aminoacyl-tRNA deacylase

**Reaction:** a D-aminoacyl-tRNA +  $H_2O$  = a D-amino acid + tRNA

**Other name(s):** Dtd2; D-Tyr-tRNA(Tyr) deacylase; D-Tyr-tRNA<sup>Tyr</sup> deacylase; D-tyrosyl-tRNA<sup>Tyr</sup> aminoacylhydrolase;

dtdA (gene name)

**Systematic name:** D-aminoacyl-tRNA aminoacylhydrolase

Comments: The enzyme from Escherichia coli can cleave D-tyrosyl-tRNA<sup>Tyr</sup>, D-aspartyl-tRNA<sup>Asp</sup> and D-

tryptophanyl-tRNA<sup>Trp</sup> [2733]. Whereas the enzyme from the archaeon *Pyrococcus abyssi* is a zinc

protein, the enzyme from Escherichia coli does not carry any zinc [760].

**References:** [2733, 760, 759, 3225]

[EC 3.1.1.96 created 2014]

EC 3.1.1.97

**Accepted name:** methylated diphthine methylhydrolase

**Reaction:** diphthine methyl ester-[translation elongation factor 2] + H<sub>2</sub>O = diphthine-[translation elongation

factor 2] + methanol

Other name(s): Dph7; diphthine methylesterase (incorrect)
Systematic name: diphthine methyl ester acylhydrolase
Comments: The protein is only present in eukaryotes.

**References:** [1698]

[EC 3.1.1.97 created 2014, modified 2015]

EC 3.1.1.98

**Accepted name:** [Wnt protein] *O*-palmitoleoyl-L-serine hydrolase

**Reaction:** [Wnt]-O-(9Z)-hexadec-9-enoyl-L-serine + H<sub>2</sub>O = [Wnt]-L-serine + (9Z)-hexadec-9-enoate

**Other name(s):** Notum

**Systematic name:** [Wnt]-O-(9Z)-hexadec-9-enoyl-L-serine acylhydrolase

Comments: The enzyme removes the palmitoleate modification that is introduced to specific L-serine residues in

Wnt proteins by EC 2.3.1.250, [Wnt protein]-O-palmitoleoyl transferase.

**References:** [1359]

[EC 3.1.1.98 created 2015]

EC 3.1.1.99

**Accepted name:** 6-deoxy-6-sulfogluconolactonase

**Reaction:** 6-deoxy-6-sulfo-D-glucono-1,5-lactone +  $H_2O$  = 6-deoxy-6-sulfo-D-gluconate

Other name(s): SGL lactonase

**Systematic name:** 6-deoxy-6-sulfo-D-glucono-1,5-lactone lactonohydrolase

**Comments:** The enzyme, characterized from the bacterium *Pseudomonas putida* SQ1, participates in a sulfo-

quinovose degradation pathway.

**References:** [753]

# [EC 3.1.1.99 created 2016]

EC 3.1.1.100

**Accepted name:** chlorophyllide *a* hydrolase

**Reaction:** chlorophyllide  $a + H_2O = 8$ -ethyl-12-methyl-3-vinyl-bacteriochlorophyllide d + methanol +  $CO_2$ 

**Other name(s):** *bciC* (gene name)

**Systematic name:** chlorophyllide-*a* hydrolase

**Comments:** This enzyme, found in green sulfur bacteria (*Chlorobiaceae*) and green filamentous bacteria (*Chlorobiaceae*)

roflexaceae), catalyses the first committed step in the biosynthesis of bacteriochlorophylls c, d and e, the removal of the C-13<sup>2</sup>-methylcarboxyl group from chlorophyllide a. The reaction is very similar to the conversion of pheophorbide a to pyropheophorbide a during chlorophyll a degradation, which is

catalysed by EC 3.1.1.82, pheophorbidase.

**References:** [1727]

[EC 3.1.1.100 created 2016]

EC 3.1.1.101

**Accepted name:** poly(ethylene terephthalate) hydrolase

**Reaction:** (ethylene terephthalate) $_n + H_2O = (\text{ethylene terephthalate})_{n-1} + 4-[(2-$ 

hydroxyethoxy)carbonyl]benzoate

**Other name(s):** PETase; PET hydrolase

**Systematic name:** poly(ethylene terephthalate) hydrolase

**Comments:** The enzyme, isolated from the bacterium *Ideonella sakaiensis*, also produces small amounts of

terephthalate (cf. EC 3.1.1.102, mono(ethylene terephthalate) hydrolase). The reaction takes place

on PET-film placed in solution.

**References:** [3301]

[EC 3.1.1.101 created 2016]

EC 3.1.1.102

Accepted name: mono(ethylene terephthalate) hydrolase

**Reaction:** 4-[(2-hydroxyethoxy)carbonyl]benzoate + H<sub>2</sub>O = terephthalate + ethylene glycol

**Other name(s):** MHET hydrolase; MHETase

**Systematic name:** 4-[(2-hydroxyethoxy)carbonyl]benzoate acylhydrolase

Comments: The enzyme, isolated from the bacterium *Ideonella sakaiensis*, has no activity with poly(ethylene

terephthalate) PET (cf. EC 3.1.1.101, poly(ethylene terephthalate) hydrolase).

**References:** [3301]

[EC 3.1.1.102 created 2016]

EC 3.1.1.103

**Accepted name:** teichoic acid D-alanine hydrolase

**Reaction:**  $[(4-D-Ala)-(2-GlcNAc)-Rib-ol-P]_n-[Gro-P]_m-ManNAc-GlcNAc-PP-peptidoglycan + <math>n$  H<sub>2</sub>O =  $[(2-D-Ala)-(2-GlcNAc)-Rib-ol-P]_n-[Gro-P]_m-ManNAc-GlcNAc-PP-peptidoglycan + <math>n$  H<sub>2</sub>O =  $[(2-D-Ala)-(2-GlcNAc)-Rib-ol-P]_n-[Gro-P]_$ 

GlcNAc)-Rib-ol-P] $_n$ -[Gro-P] $_m$ -ManNAc-GlcNAc-PP-peptidoglycan + n D-alanine

**Other name(s):** *fmtA* (gene name)

**Systematic name:** teichoic acid D-alanylhydrolase

**Comments:** The enzyme, characterized from the bacterium *Staphylococcus aureus*, removes D-alanine groups

from the teichoic acid produced by this organism, thus modulating the electrical charge of the bacte-

rial surface. The activity greatly increases methicillin resistance in MRSA strains.

**References:** [1501, 2335, 2358]

[EC 3.1.1.103 created 2018]

#### EC 3.1.1.104

**Accepted name:** 5-phospho-D-xylono-1,4-lactonase

**Reaction:** (1) D-xylono-1,4-lactone 5-phosphate  $+ H_2O = 5$ -phospho-D-xylonate

(2) L-arabino-1,4-lactone 5-phosphate +  $H_2O$  = 5-phospho-L-arabinate

**Systematic name:** 5-phospho-D-xylono-1,4-lactone hydrolase

**Comments:** The enzyme, characterized from *Mycoplasma* spp., contains a binuclear metal center with two

zinc cations. The enzyme is specific for the phosphorylated forms, and is unable to hydrolyse non-

phosphorylated 1,4-lactones.

**References:** [1510]

[EC 3.1.1.104 created 2018]

# EC 3.1.2 Thioester hydrolases

#### EC 3.1.2.1

Accepted name: acetyl-CoA hydrolase

**Reaction:** acetyl-CoA +  $H_2O = CoA + acetate$ 

Other name(s): acetyl-CoA deacylase; acetyl-CoA acylase; acetyl coenzyme A hydrolase; acetyl coenzyme A deacy-

lase; acetyl coenzyme A acylase; acetyl-CoA thiol esterase

**Systematic name:** acetyl-CoA hydrolase

**References:** [894]

[EC 3.1.2.1 created 1961]

## EC 3.1.2.2

**Accepted name:** palmitoyl-CoA hydrolase

**Reaction:** palmitoyl-CoA +  $H_2O = CoA + palmitate$ 

Other name(s): long-chain fatty-acyl-CoA hydrolase; palmitoyl coenzyme A hydrolase; palmitoyl thioesterase; palmi-

toyl coenzyme A hydrolase; palmitoyl-CoA deacylase; palmityl thioesterase; palmityl-CoA deacylase;

fatty acyl thioesterase I; palmityl thioesterase I

**Systematic name:** palmitoyl-CoA hydrolase

**Comments:** Also hydrolyses CoA thioesters of other long-chain fatty acids.

**References:** [146, 193, 1939, 2745, 3238]

[EC 3.1.2.2 created 1961]

#### EC 3.1.2.3

Accepted name: succinyl-CoA hydrolase

**Reaction:** succinyl-CoA +  $H_2O$  = CoA + succinate

Other name(s): succinyl-CoA acylase; succinyl coenzyme A hydrolase; succinyl coenzyme A deacylase

Systematic name: succinyl-CoA hydrolase

**References:** [894]

[EC 3.1.2.3 created 1961]

# EC 3.1.2.4

**Accepted name:** 3-hydroxyisobutyryl-CoA hydrolase

**Reaction:** 3-hydroxy-2-methylpropanoyl-CoA +  $H_2O$  = CoA + 3-hydroxy-2-methylpropanoate

**Other name(s):** 3-hydroxy-isobutyryl CoA hydrolase; HIB CoA deacylase

**Systematic name:** 3-hydroxy-2-methylpropanoyl-CoA hydrolase **Comments:** Also hydrolyses 3-hydroxypropanoyl-CoA.

**References:** [2408]

## [EC 3.1.2.4 created 1961]

EC 3.1.2.5

**Accepted name:** hydroxymethylglutaryl-CoA hydrolase

**Reaction:** (S)-3-hydroxy-3-methylglutaryl-CoA +  $H_2O = CoA + 3$ -hydroxy-3-methylglutarate

**Other name(s):** β-hydroxy-β-methylglutaryl coenzyme A hydrolase; β-hydroxy-β-methylglutaryl coenzyme A deacy-

lase; hydroxymethylglutaryl coenzyme A hydrolase; hydroxymethylglutaryl coenzyme A deacylase;

3-hydroxy-3-methylglutaryl-CoA hydrolase

**Systematic name:** (*S*)-3-hydroxy-3-methylglutaryl-CoA hydrolase

**References:** [570]

[EC 3.1.2.5 created 1961]

EC 3.1.2.6

**Accepted name:** hydroxyacylglutathione hydrolase

**Reaction:** S-(2-hydroxyacyl)glutathione +  $H_2O$  = glutathione + a 2-hydroxy carboxylate

Other name(s): glyoxalase II; S-2-hydroxylacylglutathione hydrolase; hydroxyacylglutathione hydrolase; acetoacetyl-

glutathione hydrolase

**Systematic name:** S-(2-hydroxyacyl)glutathione hydrolase

**Comments:** Also hydrolyses *S*-acetoacetylglutathione, but more slowly.

**References:** [2352, 3014, 3015]

[EC 3.1.2.6 created 1961 (EC 3.1.2.8 created 1961, incorporated 1978)]

EC 3.1.2.7

**Accepted name:** glutathione thiolesterase

**Reaction:** S-acylglutathione +  $H_2O$  = glutathione + a carboxylate

**Other name(s):** citryl-glutathione thioesterhydrolase

**Systematic name:** S-acylglutathione hydrolase

**References:** [1438]

[EC 3.1.2.7 created 1961]

[3.1.2.8 Deleted entry. S-acetoacylglutathione hydrolase. Now included with EC 3.1.2.6 hydroxyacylglutathione hydrolase]

[EC 3.1.2.8 created 1961, deleted 1978]

[3.1.2.9 Deleted entry. S-acetoacetylhydrolipoate hydrolase]

[EC 3.1.2.9 created 1961, deleted 1964]

EC 3.1.2.10

**Accepted name:** formyl-CoA hydrolase

**Reaction:** formyl-CoA +  $H_2O = CoA + formate$ 

**Other name(s):** formyl coenzyme A hydrolase

**Systematic name:** formyl-CoA hydrolase

References: [2691]

[EC 3.1.2.10 created 1965]

EC 3.1.2.11

Accepted name: acetoacetyl-CoA hydrolase

**Reaction:** acetoacetyl-CoA +  $H_2O$  = CoA + acetoacetate

Other name(s): acetoacetyl coenzyme A hydrolase; acetoacetyl CoA deacylase; acetoacetyl coenzyme A deacylase

Systematic name: acetoacetyl-CoA hydrolase

**References:** [64, 655]

[EC 3.1.2.11 created 1972]

EC 3.1.2.12

**Accepted name:** *S*-formylglutathione hydrolase

**Reaction:** S-formylglutathione +  $H_2O$  = glutathione + formate

**Systematic name:** *S*-formylglutathione hydrolase

**Comments:** Also hydrolyses *S*-acetylglutathione, but more slowly.

**References:** [3014, 3017, 1056]

[EC 3.1.2.12 created 1978]

EC 3.1.2.13

**Accepted name:** S-succinylglutathione hydrolase

**Reaction:** S-succinylglutathione +  $H_2O$  = glutathione + succinate

**Systematic name:** S-succinylglutathione hydrolase

**References:** [3014, 3016]

[EC 3.1.2.13 created 1978]

EC 3.1.2.14

Accepted name: oleoyl-[acyl-carrier-protein] hydrolase

**Reaction:** an oleoyl-[acyl-carrier protein] +  $H_2O$  = an [acyl-carrier protein] + oleate

Other name(s): acyl-[acyl-carrier-protein] hydrolase; acyl-ACP-hydrolase; acyl-acyl carrier protein hydrolase; oleoyl-

ACP thioesterase; oleoyl-acyl carrier protein thioesterase; oleoyl-[acyl-carrier-protein] hydrolase

**Systematic name:** oleoyl-[acyl-carrier protein] hydrolase

Comments: Acts on acyl-carrier-protein thioesters of fatty acids from  $C_{12}$  to  $C_{18}$ , but the derivative of oleic acid is

hydrolysed much more rapidly than any other compound tested.

**References:** [2162, 2639]

[EC 3.1.2.14 created 1984]

[3.1.2.15 Deleted entry. This activity is covered by EC 3.4.19.12, ubiquitinyl hydrolase 1]

[EC 3.1.2.15 created 1986, deleted 2014]

EC 3.1.2.16

**Accepted name:** citrate-lyase deacetylase

**Reaction:** acetyl-[citrate (pro-3S)-lyase] + H<sub>2</sub>O = holo-[citrate (pro-3S)-lyase] + acetate

Other name(s): [citrate-(pro-3S)-lyase] thiolesterase; acetyl-S-(acyl-carrier protein) enzyme thioester hydrolase; cit-

rate lyase deacetylase; [citrate-(pro-3S)-lyase](acetyl-form) hydrolase

**Systematic name:** acetyl-[citrate-(*pro-3S*)-lyase] hydrolase

Comments: In the proteobacterium Rubrivivax gelatinosus, this enzyme modulates the activity of EC 4.1.3.6, cit-

rate (*pro-3S*)-lyase, by converting it from its active acetyl form into its inactive thiol form by removal of its acetyl groups [915]. The activity of citrate-lyase deacetylase is itself inhibited by L-glutamate

[915].

**References:** [914, 915]

[EC 3.1.2.16 created 1989]

EC 3.1.2.17

**Accepted name:** (S)-methylmalonyl-CoA hydrolase

**Reaction:** (S)-methylmalonyl-CoA +  $H_2O$  = methylmalonate + CoA

**Other name(s):** D-methylmalonyl-coenzyme A hydrolase **Systematic name:** (*S*)-methylmalonyl-CoA hydrolase

**References:** [1524]

[EC 3.1.2.17 created 1989]

EC 3.1.2.18

Accepted name: ADP-dependent short-chain-acyl-CoA hydrolase

**Reaction:**  $acyl-CoA + H_2O = CoA + a carboxylate$ 

Other name(s): short-chain acyl coenzyme A hydrolase; propionyl coenzyme A hydrolase; propionyl-CoA hydrolase;

propionyl-CoA thioesterase; short-chain acyl-CoA hydrolase; short-chain acyl-CoA thioesterase

**Systematic name:** ADP-dependent-short-chain-acyl-CoA hydrolase

**Comments:** Requires ADP; inhibited by NADH. Maximum activity is shown with propanoyl-CoA.

**References:** [32, 33]

[EC 3.1.2.18 created 1992]

EC 3.1.2.19

Accepted name: ADP-dependent medium-chain-acyl-CoA hydrolase

**Reaction:**  $acyl-CoA + H_2O = CoA + a carboxylate$ 

Other name(s): medium-chain acyl coenzyme A hydrolase; medium-chain acyl-CoA hydrolase; medium-chain acyl-

thioester hydrolase; medium-chain hydrolase; myristoyl-CoA thioesterase

Systematic name: ADP-dependent-medium-chain-acyl-CoA hydrolase

Comments: Requires ADP; inhibited by NADH. Maximum activity is shown with nonanoyl-CoA.

**References:** [32]

[EC 3.1.2.19 created 1992]

EC 3.1.2.20

**Accepted name:** acyl-CoA hydrolase

**Reaction:**  $acyl-CoA + H_2O = CoA + a carboxylate$ 

**Other name(s):** acyl coenzyme A thioesterase; acyl-CoA thioesterase; acyl coenzyme A hydrolase; thioesterase B;

thioesterase II; acyl-CoA thioesterase

**Systematic name:** acyl-CoA hydrolase

**Comments:** Broad specificity for medium- to long-chain acyl-CoA. Insensitive to NAD<sup>+</sup> (cf. EC 3.1.2.19 ADP-

dependent medium-chain-acyl-CoA hydrolase)

**References:** [33]

[EC 3.1.2.20 created 1992]

EC 3.1.2.21

Accepted name: dodecanoyl-[acyl-carrier-protein] hydrolase

**Reaction:** a dodecanoyl-[acyl-carrier protein] +  $H_2O$  = an [acyl-carrier protein] + dodecanoate

Other name(s): lauryl-acyl-carrier-protein hydrolase; dodecanoyl-acyl-carrier-protein hydrolase; dodecyl-acyl-carrier

protein hydrolase; dodecanoyl-[acyl-carrier protein] hydrolase; dodecanoyl-[acyl-carrier-protein] hy-

drolase

**Systematic name:** dodecanoyl-[acyl-carrier protein] hydrolase

**Comments:** Acts on the acyl-carrier-protein thioester of  $C_{12}$  and, with a much lower activity,  $C_{14}$  fatty acids. The

derivative of oleic acid is hydrolysed very slowly (cf. EC 3.1.2.14, oleoyl-[acyl-carrier-protein] hydro-

lase).

**References:** [2305, 547]

## [EC 3.1.2.21 created 1999]

#### EC 3.1.2.22

Accepted name: palmitoyl[protein] hydrolase

**Reaction:** palmitoyl[protein] +  $H_2O$  = palmitate + protein

**Other name(s):** palmitoyl-protein thioesterase; palmitoyl-(protein) hydrolase

**Systematic name:** palmitoyl[protein] hydrolase

Comments: Specific for long-chain thioesters of fatty acids. Hydrolyses fatty acids from S-acylated cysteine

residues in proteins, palmitoyl cysteine and palmitoyl-CoA.

**References:** [358, 2584, 3061]

[EC 3.1.2.22 created 1999]

#### EC 3.1.2.23

**Accepted name:** 4-hydroxybenzoyl-CoA thioesterase

**Reaction:** 4-hydroxybenzoyl-CoA +  $H_2O$  = 4-hydroxybenzoate + CoA

**Systematic name:** 4-hydroxybenzoyl-CoA hydrolase

**Comments:** This enzyme is part of the bacterial 2,4-dichlorobenzoate degradation pathway.

**References:** [394, 666]

#### [EC 3.1.2.23 created 1999]

[3.1.2.24 Transferred entry. 2-(2-hydroxyphenyl)benzenesulfinate hydrolase. Now EC 3.13.1.3, 2'-hydroxybiphenyl-2-sulfinate desulfinase. The enzyme was incorrectly classified as a thioester hydrolase when the bond broken is a C-S bond, which is not an ester]

[EC 3.1.2.24 created 2000, deleted 2005]

# EC 3.1.2.25

Accepted name: phenylacetyl-CoA hydrolase

**Reaction:** phenylglyoxylyl-CoA +  $H_2O$  = phenylglyoxylate + CoA

Systematic name: phenylglyoxylyl-CoA hydrolase

Comments: This is the second step in the conversion of phenylacetyl-CoA to phenylglyoxylate, the first step being

carried out by EC 1.17.5.1, phenylacetyl-CoA dehydrogenase.

**References:** [2415, 2576]

[EC 3.1.2.25 created 2004]

[3.1.2.26 Transferred entry. bile-acid-CoA hydrolase. Now EC 2.8.3.25, bile acid CoA transferase]

[EC 3.1.2.26 created 2005, deleted 2016]

#### EC 3.1.2.27

Accepted name: choloyl-CoA hydrolase

**Reaction:** choloyl-CoA +  $H_2O$  = cholate + CoA

Other name(s): PTE-2 (ambiguous); choloyl-coenzyme A thioesterase; chenodeoxycholoyl-coenzyme A thioesterase;

peroxisomal acyl-CoA thioesterase 2

Systematic name: choloyl-CoA hydrolase

**Comments:** Also acts on chenodeoxycholoyl-CoA and to a lesser extent on short- and medium- to long-chain

acyl-CoAs, and other substrates, including trihydroxycoprostanoyl-CoA, hydroxymethylglutaryl-CoA and branched chain acyl-CoAs, all of which are present in peroxisomes. The enzyme is strongly in-

hibited by CoA and may be involved in controlling CoA levels in the peroxisome [1208].

**References:** [1208, 2710, 2481]

## [EC 3.1.2.27 created 2005]

EC 3.1.2.28

**Accepted name:** 1,4-dihydroxy-2-naphthoyl-CoA hydrolase

**Reaction:** 1,4-dihydroxy-2-naphthoyl-CoA +  $H_2O = 1$ ,4-dihydroxy-2-naphthoate + CoA

Other name(s): *menI* (gene name); *ydiL* (gene name)

Systematic name: 1,4-dihydroxy-2-naphthoyl-CoA hydrolase

**Comments:** This enzyme participates in the synthesis of menaquinones [413], phylloquinone [3180], as well as

several plant pigments [1999, 684]. The enzyme from the cyanobacterium Synechocystis sp. PCC

6803 does not accept benzoyl-CoA or phenylacetyl-CoA as substrates [3180].

**References:** [1999, 684, 3180, 413]

[EC 3.1.2.28 created 2010]

EC 3.1.2.29

**Accepted name:** fluoroacetyl-CoA thioesterase

**Reaction:** fluoroacetyl-CoA +  $H_2O$  = fluoroacetate + CoA

Systematic name: fluoroacetyl-CoA hydrolase

Comments: Fluoroacetate is extremely toxic. It reacts with CoA to form fluoroacetyl-CoA, which substitutes

for acetyl CoA and reacts with EC 2.3.3.1 (citrate synthase) to produce fluorocitrate, a metabolite of which binds very tightly to EC 4.2.1.3 (aconitase) and halts the TCA cycle. This enzyme hydrolyses fluoroacetyl-CoA before it can react with citrate synthase, and thus confers fluoroacetate resistance on the organisms that produce it. It has been described in the poisonous plant *Dichapetalum cymosum* 

and the bacterium *Streptomyces cattleya*, both of which are fluoroacetate producers.

**References:** [1895, 1202, 600]

[EC 3.1.2.29 created 2011]

EC 3.1.2.30

**Accepted name:** (3S)-malyl-CoA thioesterase

**Reaction:** (S)-malyl-CoA +  $H_2O = (S)$ -malate + CoA

Other name(s): mcl2 (gene name)
Systematic name: (S)-malyl-CoA hydrolase

**Comments:** Stimulated by  $Mg^{2+}$  or  $Mn^{2+}$ . The enzyme has no activity with (2R,3S)-2-methylmalyl-CoA (cf. EC

4.1.3.24, malyl-CoA lyase) or other CoA esters.

**References:** [710]

[EC 3.1.2.30 created 2014]

EC 3.1.2.31

Accepted name: dihydromonacolin L-[lovastatin nonaketide synthase] thioesterase

**Reaction:** dihydromonacolin L-[lovastatin nonaketide synthase] +  $H_2O$  = holo-[lovastatin nonaketide synthase]

+ dihydromonacolin L acid

Other name(s): LovG

Systematic name: dihydromonacolin L-[lovastatin nonaketide synthase] hydrolase

**Comments:** Dihydromonacolin L acid is synthesized while bound to an acyl-carrier protein domain of the lovas-

tatin nonaketide synthase (EC 2.3.1.161). Since that enzyme lacks a thioesterase domain, release of the dihydromonacolin L acid moiety from the polyketide synthase requires this dedicated enzyme.

**References:** [3235]

[EC 3.1.2.31 created 2015]

Accepted name: 2-aminobenzoylacetyl-CoA thioesterase

**Reaction:** (2-aminobenzoyl)acetyl-CoA +  $H_2O = (2\text{-aminobenzoyl})$ acetate + CoA

**Other name(s):** pqsE (gene name)

Systematic name: (2-aminobenzoyl)acetyl-CoA hydrolase

**Comments:** The enzyme, characterized from the bacterium *Pseudomonas aeruginosa*, participates in the produc-

tion of the signal molecule 2-heptyl-4(1H)-quinolone (HHQ).

**References:** [3310, 651]

[EC 3.1.2.32 created 2016]

# EC 3.1.3 Phosphoric-monoester hydrolases

EC 3.1.3.1

Accepted name: alkaline phosphatase

**Reaction:** a phosphate monoester +  $H_2O$  = an alcohol + phosphate

Other name(s): alkaline phosphomonoesterase; phosphomonoesterase; glycerophosphatase; alkaline phosphohydro-

lase; alkaline phenyl phosphatase; orthophosphoric-monoester phosphohydrolase (alkaline optimum)

**Systematic name:** phosphate-monoester phosphohydrolase (alkaline optimum)

**Comments:** Wide specificity. Also catalyses transphosphorylations. The human placental enzyme is a zinc pro-

tein. Some enzymes hydrolyse diphosphate (cf. EC 3.6.1.1 inorganic diphosphatase)

**References:** [706, 1054, 1785, 1984, 2752]

[EC 3.1.3.1 created 1961]

EC 3.1.3.2

**Accepted name:** acid phosphatase

**Reaction:** a phosphate monoester  $+ H_2O =$  an alcohol + phosphate

Other name(s): acid phosphomonoesterase; phosphomonoesterase; glycerophosphatase; acid monophosphatase; acid

phosphohydrolase; acid phosphomonoester hydrolase; uteroferrin; acid nucleoside diphosphate phos-

phatase; orthophosphoric-monoester phosphohydrolase (acid optimum)

**Systematic name:** phosphate-monoester phosphohydrolase (acid optimum)

**Comments:** Wide specificity. Also catalyses transphosphorylations.

**References:** [1345, 1561, 2976]

[EC 3.1.3.2 created 1961]

EC 3.1.3.3

Accepted name: phosphoserine phosphatase

**Reaction:** *O*-phospho-L(or D)-serine +  $H_2O = L(or D)$ -serine + phosphate

**Systematic name:** *O*-phosphoserine phosphohydrolase

**References:** [266, 344, 2070]

[EC 3.1.3.3 created 1961]

EC 3.1.3.4

**Accepted name:** phosphatidate phosphatase

**Reaction:** a 1,2-diacylglycerol 3-phosphate +  $H_2O$  = a 1,2-diacyl-sn-glycerol + phosphate

Other name(s): phosphatic acid phosphatase; acid phosphatidyl phosphatase; phosphatic acid phosphohydrolase; PAP,

Lipin

**Systematic name:** diacylglycerol-3-phosphate phosphohydrolase

**Comments:** This enzyme catalyses the Mg<sup>2+</sup>-dependent dephosphorylation of a 1,2-diacylglycerol-3-phosphate,

yielding a 1,2-diacyl-*sn*-glycerol (DAG), the substrate for *de novo* lipid synthesis via the Kennedy pathway and for the synthesis of triacylglycerol. In lipid signalling, the enzyme generates a pool of DAG to be used for protein kinase C activation. The mammalian enzymes are known as lipins.

**References:** [2700, 373]

[EC 3.1.3.4 created 1961, modified 2010]

EC 3.1.3.5

**Accepted name:** 5'-nucleotidase

**Reaction:** a 5'-ribonucleotide +  $H_2O$  = a ribonucleoside + phosphate

Other name(s): uridine 5'-nucleotidase; 5'-adenylic phosphatase; adenosine 5'-phosphatase; AMP phosphatase;

adenosine monophosphatase; 5'-mononucleotidase; AMPase; UMPase; snake venom 5'-nucleotidase; thimidine monophosphate nucleotidase; 5'-AMPase; 5'-AMP nucleotidase; AMP phosphohydrolase;

IMP 5'-nucleotidase

**Systematic name:** 5'-ribonucleotide phosphohydrolase

**Comments:** Wide specificity for 5'-nucleotides.

**References:** [999, 1121, 2603]

[EC 3.1.3.5 created 1961]

EC 3.1.3.6

**Accepted name:** 3'-nucleotidase

**Reaction:** a 3'-ribonucleotide +  $H_2O$  = a ribonucleoside + phosphate **Other name(s):** 3'-mononucleotidase; 3'-phosphatase; 3'-ribonucleotidase

**Systematic name:** 3'-ribonucleotide phosphohydrolase **Comments:** Wide specificity for 3'-nucleotides.

**References:** [2650]

[EC 3.1.3.6 created 1961]

EC 3.1.3.7

**Accepted name:** 3'(2'),5'-bisphosphate nucleotidase

**Reaction:** adenosine 3',5'-bisphosphate +  $H_2O = AMP$  + phosphate

Other name(s): phosphoadenylate 3'-nucleotidase; 3'-phosphoadenylylsulfate 3'-p

bisphosphonucleoside 3'(2')-phosphohydrolase

**Systematic name:** adenosine-3'(2'),5'-bisphosphate 3'(2')-phosphohydrolase

**Comments:** Also acts on 3'-phosphoadenylyl sulfate, and on the corresponding 2'-phosphates.

**References:** [318, 740, 2366, 2971]

[EC 3.1.3.7 created 1961]

EC 3.1.3.8

**Accepted name:** 3-phytase

**Reaction:** myo-inositol hexakisphosphate +  $H_2O = 1D$ -myo-inositol 1,2,4,5,6-pentakisphosphate + phosphate

**Other name(s):** 1-phytase; phytate 1-phosphatase; phytate 6-phosphatase

**Systematic name:** *myo*-inositol-hexakisphosphate 3-phosphohydrolase

**References:** [491, 1328, 1255, 492]

[EC 3.1.3.8 created 1961, modified 1976, modified 2002]

EC 3.1.3.9

Accepted name: glucose-6-phosphatase

**Reaction:** D-glucose 6-phosphate  $+ H_2O = D$ -glucose + phosphate

**Other name(s):** glucose 6-phosphate phosphatase

**Systematic name:** D-glucose-6-phosphate phosphohydrolase

**Comments:** Wide distribution in animal tissues. Also catalyses potent transphosphorylations from carbamoyl

phosphate, hexose phosphates, diphosphate, phospho*enol*pyruvate and nucleoside di- and triphosphates, to D-glucose, D-mannose, 3-methyl-D-glucose or 2-deoxy-D-glucose [*cf.* EC 2.7.1.62 (phosphoramidate—hexose phosphotransferase), EC 2.7.1.79 (diphosphate—glycerol phosphotrans-

ferase) and EC 3.9.1.1 (phosphoamidase)].

**References:** [50, 466, 2110, 2111]

[EC 3.1.3.9 created 1961]

EC 3.1.3.10

Accepted name: glucose-1-phosphatase

**Reaction:**  $\alpha$ -D-glucose 1-phosphate + H<sub>2</sub>O = D-glucose + phosphate

**Systematic name:** α-D-glucose-1-phosphate phosphohydrolase

**Comments:** Also acts, more slowly, on D-galactose 1-phosphate.

**References:** [747, 2999]

[EC 3.1.3.10 created 1961]

EC 3.1.3.11

**Accepted name:** fructose-bisphosphatase

**Reaction:** D-fructose 1,6-bisphosphate +  $H_2O$  = D-fructose 6-phosphate + phosphate

Other name(s): hexose diphosphatase; FBPase; fructose 1,6-diphosphatase; fructose 1,6-diphosphate phosphatase; D-

fructose 1,6-diphosphatase; fructose 1,6-bisphosphatase; fructose diphosphatase; fructose diphosphatase; fructose bisphosphatase; fructose 1,6-bisphosphate 1-phosphatase; fructose 1,6-bisphosphate phosphatase; become 1,6-bisphosphate phosphatase; become 1,6-bisphosphate phosphatase

**Systematic name:** D-fructose-1,6-bisphosphate 1-phosphohydrolase

**Comments:** The animal enzyme also acts on sedoheptulose 1,7-bisphosphate.

**References:** [690, 954, 1949, 2310]

[EC 3.1.3.11 created 1961, modified 1976]

EC 3.1.3.12

Accepted name: trehalose-phosphatase

**Reaction:**  $\alpha, \alpha$ -trehalose 6-phosphate + H<sub>2</sub>O =  $\alpha, \alpha$ -trehalose + phosphate

**Other name(s):** trehalose 6-phosphatase; trehalose 6-phosphatase; trehalose-6-phosphate phosphohydrolase

**Systematic name:**  $\alpha, \alpha$ -trehalose-6-phosphate phosphohydrolase

**References:** [348, 365]

[EC 3.1.3.12 created 1961]

[3.1.3.13 Deleted entry. bisphosphoglycerate phosphatase. Recent studies have shown that this is a partial activity of EC 5.4.2.11, phosphoglycerate mutase (2,3-diphosphoglycerate-dependent)]

[EC 3.1.3.13 created 1961, deleted 2016]

EC 3.1.3.14

**Accepted name:** methylphosphothioglycerate phosphatase

**Reaction:** S-methyl-3-phospho-1-thio-D-glycerate + H<sub>2</sub>O = S-methyl-1-thio-D-glycerate + phosphate

**Other name(s):** methylthiophosphoglycerate phosphatase

**Systematic name:** S-methyl-3-phospho-1-thio-D-glycerate phosphohydrolase

**References:** [229]

## [EC 3.1.3.14 created 1961]

EC 3.1.3.15

Accepted name: histidinol-phosphatase

**Reaction:** L-histidinol phosphate  $+ H_2O = L$ -histidinol + phosphate

Other name(s): histidinol phosphate phosphatase; L-histidinol phosphate phosphatase; histidinolphosphate phosphate

phatase; HPpase; histidinolphosphatase

Systematic name: L-histidinol-phosphate phosphohydrolase

**References:** [46]

[EC 3.1.3.15 created 1961]

EC 3.1.3.16

Accepted name: protein-serine/threonine phosphatase

**Reaction:** [a protein]-serine/threonine phosphate  $+ H_2O = [a protein]$ -serine/threonine + phosphate

Other name(s): phosphoprotein phosphatase (ambiguous); protein phosphatase-1; protein phosphatase-2A; protein

phosphatase-2B; protein phosphatase-2C; protein D phosphatase; phosphospectrin phosphatase; casein phosphatase; *Aspergillus awamori* acid protein phosphatase; calcineurin; phosphatase 2A; phosphatase 2B; phosphatase II; phosphatase IB; phosphatase C-II; polycation modulated (PCM-) phosphatase; phosphopyruvate dehydrogenase phosphatase; phosphatase SP; branched-chain α-keto acid dehydrogenase phosphatase; BCKDH phosphatase; 3-hydroxy 3-methylglutaryl coenzymeA reductase phosphatase; HMG-CoA reductase phosphatase; phosphatase H-II; phosphatase III; phosphatase

I; protein phosphatase; phosphatase IV; phosphoprotein phosphohydrolase

**Systematic name:** protein-serine/threonine-phosphate phosphohydrolase

**Comments:** A group of enzymes removing the serine- or threonine-bound phosphate group from a wide range

of phosphoproteins, including a number of enzymes that have been phosphorylated under the action of a kinase (*cf.* EC 3.1.3.48 protein-tyrosine-phosphatase). The spleen enzyme also acts on phenolic

phosphates and phosphamides (cf. EC 3.9.1.1, phosphoamidase).

**References:** [595, 1249, 2801, 2945]

[EC 3.1.3.16 created 1961, modified 1989, modified 2013]

EC 3.1.3.17

**Accepted name:** [phosphorylase] phosphatase

**Reaction:** [phosphorylase a] + 4 H<sub>2</sub>O = 2 [phosphorylase b] + 4 phosphate

**Other name(s):** PR-enzyme; phosphorylase a phosphatase; glycogen phosphorylase phosphatase; protein phosphatase

C; type 1 protein phosphatase

**Systematic name:** [phosphorylase *a*] phosphohydrolase

**References:** [277, 966, 2363]

[EC 3.1.3.17 created 1961]

EC 3.1.3.18

**Accepted name:** phosphoglycolate phosphatase

**Reaction:** 2-phosphoglycolate +  $H_2O$  = glycolate + phosphate

Other name(s): phosphoglycolate hydrolase; 2-phosphoglycolate phosphatase; P-glycolate phosphatase; phosphogly-

collate phosphatase

**Systematic name:** 2-phosphoglycolate phosphohydrolase

**References:** [441]

[EC 3.1.3.18 created 1965]

Accepted name: glycerol-2-phosphatase

**Reaction:** glycerol 2-phosphate +  $H_2O$  = glycerol + phosphate

**Other name(s):** β-glycerophosphatase; β-glycerophosphate phosphatase; 2-glycerophosphatase

**Systematic name:** glycerol-2-phosphate phosphohydrolase

**References:** [2574, 2976]

[EC 3.1.3.19 created 1965]

EC 3.1.3.20

Accepted name: phosphoglycerate phosphatase

**Reaction:** D-glycerate 2-phosphate +  $H_2O$  = D-glycerate + phosphate

**Other name(s):** D-2-phosphoglycerate phosphatase; glycerophosphate phosphatase

**Systematic name:** D-glycerate-2-phosphate phosphohydrolase

**References:** [735]

[EC 3.1.3.20 created 1972]

EC 3.1.3.21

Accepted name: glycerol-1-phosphatase

**Reaction:** glycerol 1-phosphate +  $H_2O$  = glycerol + phosphate

**Other name(s):** α-glycerophosphatase; α-glycerol phosphatase; glycerol 3-phosphatase; glycerol-3-phosphate phos-

phatase; glycerol 3-phosphate phosphohydrolase

**Systematic name:** glycerol-1-phosphate phosphohydrolase

**Comments:** The *Dunaliella* enzyme acts more rapidly on *sn*-glycerol 1-phosphate than on the 3-phosphate. The

enzyme from yeast also acts on propane-1,2-diol 1-phosphate, but not on a variety of other phosphate

esters.

**References:** [2806]

[EC 3.1.3.21 created 1972, modified 1986]

EC 3.1.3.22

**Accepted name:** mannitol-1-phosphatase

**Reaction:** D-mannitol 1-phosphate +  $H_2O$  = D-mannitol + phosphate

**Other name(s):** mannitol-1-phosphate phosphatase

**Systematic name:** D-mannitol-1-phosphate phosphohydrolase

**References:** [2477, 3242]

[EC 3.1.3.22 created 1972]

EC 3.1.3.23

Accepted name: sugar-phosphatase

**Reaction:** sugar phosphate  $+ H_2O = sugar + phosphate$ 

**Systematic name:** sugar-phosphate phosphohydrolase

**Comments:** Has a wide specificity, acting on aldohexose 1-phosphates, ketohexose 1-phosphates, aldohexose 6-

phosphates, ketohexose 6-phosphates, both phosphate ester bonds of fructose 1,6-bisphosphate, phos-

phoric esters of disaccharides, and on pentose and triose phosphates, but at a slower rate.

**References:** [1637]

[EC 3.1.3.23 created 1972]

EC 3.1.3.24

Accepted name: sucrose-phosphate phosphatase

**Reaction:** sucrose  $6^F$ -phosphate +  $H_2O$  = sucrose + phosphate

**Other name(s):** sucrose 6-phosphate hydrolase; sucrose-phosphate hydrolase; sucrose-phosphate phosphohydrolase;

sucrose-6-phosphatase; sucrose phosphatase; sucrose-6-phosphate phosphatase; SPP

**Systematic name:** sucrose-6<sup>F</sup>-phosphate phosphohydrolase

**Comments:** Requires  $Mg^{2+}$  for maximal activity [1754]. This is the final step in the sucrose-biosynthesis pathway.

The enzyme is highly specific for sucrose 6-phosphate, with fructose 6-phosphate unable to act as a substrate [1754]. Belongs in the haloacid dehydrogenase (HAD) superfamily. The F of sucrose 6<sup>F</sup>-

phosphate is used to indicate that the fructose residue of sucrose carries the substituent.

**References:** [1081, 1754, 1755, 767]

[EC 3.1.3.24 created 1972, modified 2008]

#### EC 3.1.3.25

Accepted name: inositol-phosphate phosphatase

**Reaction:** myo-inositol phosphate +  $H_2O = myo$ -inositol + phosphate

**Other name(s):** *myo*-inositol-1(or 4)-monophosphatase; inositol 1-phosphatase; L-*myo*-inositol-1-phosphate phos-

phatase; *myo*-inositol 1-phosphatase; inositol phosphatase; inositol monophosphate phosphatase; inositol-1(or 4)-monophosphatase; *myo*-inositol-1(or 4)-phosphate phosphohydrolase; *myo*-inositol

 $monophosphatase; {\it myo}\mbox{-inositol-1-phosphatase}$ 

**Systematic name:** *myo*-inositol-phosphate phosphohydrolase

**Comments:** Acts on five of the six isomers of *myo*-inositol phosphate, all except *myo*-inositol 2-phosphate, but

does not act on myo-inositol bearing more than one phosphate group. It also acts on adenosine 2'-phosphate (but not the 3'- or 5'- phosphates), sn-glycerol 3-phosphate and glycerol 2-phosphate. Two

isoforms are known [3302].

**References:** [687, 887, 1028, 3302, 3214, 11]

[EC 3.1.3.25 created 1972, modified 1990, modified 2002, modified 2004]

# EC 3.1.3.26

**Accepted name:** 4-phytase

**Reaction:** myo-inositol hexakisphosphate +  $H_2O = 1D$ -myo-inositol 1,2,3,5,6-pentakisphosphate + phosphate **Other name(s):** 6-phytase (name based on 1L-numbering system and not 1D-numbering); phytase; phytate 6-

phosphatase; myo-inositol-hexakisphosphate 6-phosphohydrolase (name based on 1L-numbering sys-

tem and not 1D-numbering)

**Systematic name:** *myo*-inositol-hexakisphosphate 4-phosphohydrolase

**References:** [1328, 2938, 1688, 492]

[EC 3.1.3.26 created 1972, modified 1976, modified 2002]

# EC 3.1.3.27

Accepted name: phosphatidylglycerophosphatase

**Reaction:** phosphatidylglycerophosphate +  $H_2O$  = phosphatidylglycerol + phosphate

Other name(s): phosphatidylglycerol phosphate phosphatase; phosphatidylglycerol phosphatase; PGP phosphatase

**Systematic name:** phosphatidylglycerophosphate phosphohydrolase

**References:** [399]

[EC 3.1.3.27 created 1972]

# EC 3.1.3.28

**Accepted name:** ADP-phosphoglycerate phosphatase

**Reaction:** 3-(ADP)-2-phosphoglycerate +  $H_2O = 3-(ADP)$ -glycerate + phosphate

**Other name(s):** adenosine diphosphate phosphoglycerate phosphatase **Systematic name:** 3-(ADP)-2-phosphoglycerate phosphohydrolase

**Comments:** Also acts on 2,3-bisphosphoglycerate.

**References:** [3317]

[EC 3.1.3.28 created 1972]

EC 3.1.3.29

**Accepted name:** *N*-acylneuraminate-9-phosphatase

**Reaction:** N-acylneuraminate 9-phosphate +  $H_2O = N$ -acylneuraminate + phosphate

Other name(s): acylneuraminate 9-phosphatase; N-acylneuraminic acid 9-phosphate phosphatase; N-acylneuraminic

(sialic) acid 9-phosphatase

**Systematic name:** *N*-acylneuraminate-9-phosphate phosphohydrolase

**References:** [1344]

[EC 3.1.3.29 created 1972]

[3.1.3.30 Deleted entry. 3'-phosphoadenylylsulfate 3'-phosphatase. Now included with EC 3.1.3.31 nucleotidase]

[EC 3.1.3.30 created 1972, deleted 1992]

EC 3.1.3.31

Accepted name: nucleotidase

**Reaction:** a nucleotide +  $H_2O$  = a nucleoside + phosphate

Other name(s): nucleotide phosphatase; nucleotide-specific phosphatase; NSP I; NSP II; deoxyribonucleoside-

activated nucleotidase (DAN); deoxyinosine-activated nucleotidase (DIAN); acid nucleotidase

Systematic name: nucleotide phosphohydrolase

Comments: A wide specificity for 2', 3'- and 5'- nucleotides; also hydrolyses glycerol phosphate and 4-

nitrophenyl phosphate.

**References:** [76]

[EC 3.1.3.31 created 1972 (EC 3.1.3.30 created 1972, incorporated 1992)]

EC 3.1.3.32

**Accepted name:** polynucleotide 3'-phosphatase

**Reaction:** a 3'-phosphopolynucleotide +  $H_2O$  = a polynucleotide + phosphate

**Other name(s):** 2'(3')-polynucleotidase; DNA 3'-phosphatase; deoxyribonucleate 3'-phosphatase; 5'-

polynucleotidekinase 3'-phosphatase

**Systematic name:** polynucleotide 3'-phosphohydrolase

**Comments:** Also hydrolyses nucleoside 2'-, 3'- and 5'-monophosphates, but only 2'- and 3'-

phosphopolynucleotides.

**References:** [180]

[EC 3.1.3.32 created 1972]

EC 3.1.3.33

**Accepted name:** polynucleotide 5'-phosphatase

**Reaction:** a 5'-phosphopolynucleotide +  $H_2O$  = a polynucleotide + phosphate

**Other name(s):** 5'-polynucleotidase

**Systematic name:** polynucleotide 5'-phosphohydrolase

Comments: Does not act on nucleoside monophosphates. Induced in *Escherichia coli* by T-even phages.

**References:** [180]

[EC 3.1.3.33 created 1972]

### EC 3.1.3.34

**Accepted name:** deoxynucleotide 3'-phosphatase

**Reaction:** a 2'-deoxyribonucleoside 3'-phosphate +  $H_2O$  = a 2'-deoxyribonucleoside + phosphate

Other name(s): 3'-deoxynucleotidase; 3'-deoxyribonucleotidase Systematic name: 2'-deoxyribonucleotide 3'-phosphohydrolase

**Comments:** Also catalyses the selective removal of 3'-phosphate groups from DNA and oligodeoxyribonu-

cleotides. Induced in Escherichia coli by T-even phages.

**References:** [180]

[EC 3.1.3.34 created 1972]

#### EC 3.1.3.35

**Accepted name:** thymidylate 5'-phosphatase

**Reaction:** thymidylate  $+ H_2O =$  thymidine + phosphate

Other name(s): thymidylate 5'-nucleotidase; deoxythymidylate 5'-nucleotidase; thymidylate nucleotidase; de-

oxythymidylic 5'-nucleotidase; deoxythymidylate phosphohydrolase; dTMPase

**Systematic name:** thymidylate 5'-phosphohydrolase

**Comments:** Acts on 5-methyl-dCMP and on TMP, but more slowly than on dTMP.

**References:** [63]

[EC 3.1.3.35 created 1972]

### EC 3.1.3.36

**Accepted name:** phosphoinositide 5-phosphatase

**Reaction:** 1-phosphatidyl-1D-myo-inositol 4,5-bisphosphate + H<sub>2</sub>O = 1-phosphatidyl-1D-myo-inositol 4-

phosphate + phosphate

Other name(s): type II inositol polyphosphate 5-phosphatase; triphosphoinositide phosphatase; IP<sub>3</sub> phosphatase; Pt-

 $dIns(4,5)P_2$  phosphatase; triphosphoinositide phosphomonoesterase; diphosphoinositide phosphatase; inositol 1,4,5-triphosphate 5-phosphomonoesterase; inositol triphosphate 5-phosphomonoesterase; phosphatidylinositol-bisphosphatase; phosphatidyl-myo-inositol-4,5-bisphosphate phosphatase; phosphatidylinositol 4,5-bisphosphate phosphatase; polyphosphoinositol lipid 5-phosphatase;

phosphatidyl-inositol-bisphosphate phosphatase

**Systematic name:** phosphatidyl-*myo*-inositol-4,5-bisphosphate 4-phosphohydrolase

**Comments:** These enzymes can also remove the 5-phosphate from  $Ins(1,4,5)P_3$  and/or  $Ins(1,3,4,5)P_4$ . They are a

diverse family of enzymes, with differing abilities to catalyse two or more of the four reactions listed. They are thought to use inositol lipids rather than inositol phosphates as substrates *in vivo*. All of them can use either or both of  $PtdIns(4,5)P_2$  and  $PtdIns(3,4,5)P_3$  as substrates; this is the main prop-

erty that distinguishes them from EC 3.1.3.56, inositol-polyphosphate 5-phosphatase.

**References:** [556, 2432, 3214]

[EC 3.1.3.36 created 1972, modified 2002]

## EC 3.1.3.37

**Accepted name:** sedoheptulose-bisphosphatase

**Reaction:** sedoheptulose 1,7-bisphosphate + H<sub>2</sub>O = sedoheptulose 7-phosphate + phosphate

Other name(s): SBPase; sedoheptulose 1,7-diphospate phosphatase; sedoheptulose 1,7-diphosphatase; sedoheptulose

diphosphatase; sedoheptulose bisphosphatase; sedoheptulose 1,7-bisphosphatase

**Systematic name:** sedoheptulose-1,7-bisphosphate 1-phosphohydrolase

**References:** [2353, 2960]

[EC 3.1.3.37 created 1976]

EC 3.1.3.38

**Accepted name:** 3-phosphoglycerate phosphatase

**Reaction:** D-glycerate 3-phosphate  $+ H_2O = D$ -glycerate + phosphate **Other name(s):** D-3-Phosphoglycerate phosphatase; 3-PGA phosphatase

**Systematic name:** D-glycerate-3-phosphate phosphohydrolase

**Comments:** Wide specificity, but 3-phosphoglycerate is the best substrate.

**References:** [2369]

[EC 3.1.3.38 created 1976]

EC 3.1.3.39

**Accepted name:** streptomycin-6-phosphatase

**Reaction:** streptomycin 6-phosphate +  $H_2O$  = streptomycin + phosphate

Other name(s): streptomycin 6-phosphate phosphatese; streptomycin 6-phosphate phosphohydrolase; streptomycin-6-

P phosphohydrolase

**Systematic name:** streptomycin-6-phosphate phosphohydrolase

**Comments:** Also acts on dihydrostreptomycin  $3'\alpha$ ,6-bisphosphate and streptidine 6-phosphate.

**References:** [3096, 3097]

[EC 3.1.3.39 created 1976]

EC 3.1.3.40

**Accepted name:** guanidinodeoxy-scyllo-inositol-4-phosphatase

**Reaction:** 1-guanidino-1-deoxy-scyllo-inositol 4-phosphate + H<sub>2</sub>O = 1-guanidino-1-deoxy-scyllo-inositol +

phosphate

Other name(s): 1-guanidino-scyllo-inositol 4-phosphatase; 1-guanidino-1-deoxy-scyllo-inositol-4-P phosphohydro-

lase

**Systematic name:** 1-guanidino-1-deoxy-*scyllo*-inositol-4-phosphate 4-phosphohydrolase

**References:** [3097]

[EC 3.1.3.40 created 1976]

EC 3.1.3.41

**Accepted name:** 4-nitrophenylphosphatase

**Reaction:** 4-nitrophenyl phosphate  $+ H_2O = 4$ -nitrophenol + phosphate

**Other name(s):** nitrophenyl phosphatase; *p*-nitrophenylphosphatase; para-nitrophenyl phosphatase; K-pNPPase;

NPPase; PNPPase; Ecto-p-nitrophenyl phosphatase; p-nitrophenylphosphate phosphohydrolase

**Systematic name:** 4-nitrophenylphosphate phosphohydrolase

**Comments:** A number of other substances, including phenyl phosphate, 4-nitrophenyl sulfate, acetyl phosphate

and glycerol phosphate, are not substrates.

**References:** [93, 94]

[EC 3.1.3.41 created 1976]

EC 3.1.3.42

Accepted name: [glycogen-synthase-D] phosphatase

**Reaction:** [glycogen-synthase D] +  $H_2O$  = [glycogen-synthase I] + phosphate

Other name(s): uridine diphosphoglucose-glycogen glucosyltransferase phosphatase; UDP-glycogen glucosyltrans-

ferase phosphatase; UDPglucose-glycogen glucosyltransferase phosphatase; glycogen glucosyltransferase phosphatase; glycogen synthese phosphatase; glycogen synthase phosphatase; glycogen synthase D phosphatase; Mg<sup>2+</sup> dependent glycogen synthase phosphatase; phosphatase type 2°C

**Systematic name:** [UDP-glucose:glycogen 4-α-D-glucosyltransferase-D] phosphohydrolase

**Comments:** The product is EC 2.4.1.11 glycogen(starch) synthase.

**References:** [6]

### [EC 3.1.3.42 created 1976]

EC 3.1.3.43

**Accepted name:** [pyruvate dehydrogenase (acetyl-transferring)]-phosphatase

**Reaction:** [pyruvate dehydrogenase (acetyl-transferring)] phosphate  $+ H_2O = [pyruvate dehydrogenase (acetyl-transferring)]$ 

transferring)] + phosphate

**Other name(s):** pyruvate dehydrogenase phosphatase; phosphopyruvate dehydrogenase phosphatase; [pyruvate dehydrogenase phosphatase]

drogenase (lipoamide)]-phosphatase; [pyruvate dehydrogenase (lipoamide)]-phosphate phosphohy-

drolase

**Systematic name:** [pyruvate dehydrogenase (acetyl-transferring)]-phosphate phosphohydrolase

Comments: A mitochondrial enzyme associated with EC 1.2.4.1 pyruvate dehydrogenase (acetyl-transferring), in

the pyruvate dehydrogenase complex.

**References:** [1708, 2395]

[EC 3.1.3.43 created 1978]

EC 3.1.3.44

**Accepted name:** [acetyl-CoA carboxylase]-phosphatase

**Reaction:** [acetyl-CoA carboxylase] phosphate  $+ H_2O = [acetyl-CoA carboxylase] + phosphate$ **Systematic name:** [acetyl-CoA:carbon-dioxide ligase (ADP-forming)]-phosphate phosphohydrolase

**Comments:** Simultaneously dephosphorylates and activates EC 6.4.1.2 acetyl-CoA carboxylase. Acts similarly

on EC 1.1.1.88 (hydroxymethylglutaryl-CoA reductase), EC 2.4.1.1 (phosphorylase), EC 2.4.1.11 [glycogen(starch) synthase], and dephosphorylates phosphoprotamine and 4-nitrophenyl phosphate. Not identical to EC 3.1.3.17 ([phosphorylase] phosphatase) or EC 3.1.3.43 [pyruvate dehydrogenase

(acetyl-transferring)]-phosphatase.

**References:** [1527]

[EC 3.1.3.44 created 1983]

EC 3.1.3.45

Accepted name: 3-deoxy-manno-octulosonate-8-phosphatase

**Reaction:** 3-deoxy-D-manno-octulosonate 8-phosphate +  $H_2O = 3$ -deoxy-D-manno-octulosonate + phosphate

**Systematic name:** 3-deoxy-D-*manno*-octulosonate-8-phosphate 8-phosphohydrolase

**References:** [2385]

[EC 3.1.3.45 created 1983]

EC 3.1.3.46

**Accepted name:** fructose-2,6-bisphosphate 2-phosphatase

**Reaction:**  $\beta$ -D-fructose 2,6-bisphosphate +  $H_2O$  = D-fructose 6-phosphate + phosphate **Other name(s):** fructose-2,6-bisphosphatase; D-fructose-2,6-bisphosphate 2-phosphohydrolase

**Systematic name:**  $\beta$ -D-fructose-2,6-bisphosphate 2-phosphohydrolase

Comments: The enzyme copurifies with EC 2.7.1.105 6-phosphofructo-2-kinase. (cf. EC 3.1.3.54 fructose-2,6-

bisphosphate 6-phosphatase).

**References:** [2553]

[EC 3.1.3.46 created 1984]

EC 3.1.3.47

Accepted name: [hydroxymethylglutaryl-CoA reductase (NADPH)]-phosphatase

**Reaction:** [hydroxymethylglutaryl-CoA reductase (NADPH)] phosphate  $+ H_2O = [hydroxymethylglutaryl-CoA$ 

reductase (NADPH)] + phosphate

Other name(s): reductase phosphatase

**Systematic name:** [hydroxymethylglutaryl-CoA reductase (NADPH)]-phosphate phosphohydrolase

**Comments:** Acts on the product of the reaction catalysed by EC 2.7.11.31 [hydroxymethylglutaryl-CoA

reductase (NADPH)] kinase, simultaneously dephosphorylating and activating EC 1.1.1.34

hydroxymethylglutaryl-CoA reductase (NADPH).

**References:** [917, 918]

[EC 3.1.3.47 created 1984]

EC 3.1.3.48

Accepted name: protein-tyrosine-phosphatase

> [a protein]-tyrosine phosphate +  $H_2O$  = [a protein]-tyrosine + phosphate **Reaction:**

Other name(s): phosphotyrosine phosphatase; phosphoprotein phosphatase (phosphotyrosine); phosphotyrosine his-

> tone phosphatase; protein phosphotyrosine phosphatase; tyrosylprotein phosphatase; phosphotyrosine protein phosphatase; phosphotyrosylprotein phosphatase; tyrosine O-phosphate phosphatase; PPT-

phosphatase; PTPase; [phosphotyrosine]protein phosphatase; PTP-phosphatase

**Systematic name:** protein-tyrosine-phosphate phosphohydrolase

Dephosphorylates O-phosphotyrosine groups in phosphoproteins, such as the products of EC 2.7.10.2, **Comments:** 

non-specific protein-tyrosine kinase.

References: [798, 870]

[EC 3.1.3.48 created 1984]

EC 3.1.3.49

Accepted name: [pyruvate kinase]-phosphatase

> Reaction: [pyruvate kinase] phosphate +  $H_2O$  = [pyruvate kinase] + phosphate

Other name(s): pyruvate kinase phosphatase

**Systematic name:** [ATP:pyruvate 2-O-phosphotransferase]-phosphate phosphohydrolase

**Comments:** Simultaneously dephosphorylates and activates EC 2.7.1.40 pyruvate kinase, that has been inactivated

by protein kinase.

**References:** [1321]

[EC 3.1.3.49 created 1984]

EC 3.1.3.50

Accepted name: sorbitol-6-phosphatase

> sorbitol 6-phosphate +  $H_2O$  = sorbitol + phosphate **Reaction:**

Other name(s): sorbitol-6-phosphate phosphatase sorbitol-6-phosphate phosphohydrolase Systematic name: **Comments:** Acts, very slowly, on hexose 6-phosphates.

**References:** [964]

[EC 3.1.3.50 created 1984]

EC 3.1.3.51

Accepted name: dolichyl-phosphatase

> **Reaction:** dolichyl phosphate +  $H_2O$  = dolichol + phosphate

Other name(s): dolichol phosphate phosphatase; dolichol phosphatase; dolichol monophosphatase; dolichyl

monophosphate phosphatase; dolichyl phosphate phosphatase; polyisoprenyl phosphate phosphatase;

polyprenylphosphate phosphatase; Dol-P phosphatase

dolichyl-phosphate phosphohydrolase **Systematic name:** 

> References: [17, 2429, 3146]

### [EC 3.1.3.51 created 1984]

EC 3.1.3.52

Accepted name: [3-methyl-2-oxobutanoate dehydrogenase (2-methylpropanoyl-transferring)]-phosphatase

**Reaction:** [3-methyl-2-oxobutanoate dehydrogenase (2-methylpropanoyl-transferring)] phosphate  $+ H_2O = [3-methyl-2-oxobutanoate dehydrogenase (2-methylpropanoyl-transferring)]$ 

methyl-2-oxobutanoate dehydrogenase (2-methylpropanoyl-transferring)] + phosphate

Other name(s): branched-chain oxo-acid dehydrogenase phosphatase; branched-chain 2-keto acid dehydroge-

> nase phosphatase; branched-chain α-keto acid dehydrogenase phosphatase; BCKDH; [3-methyl-2oxobutanoate dehydrogenase (lipoamide)]-phosphatase; [3-methyl-2-oxobutanoate dehydrogenase

(lipoamide)]-phosphate phosphohydrolase

**Systematic name:** [3-methyl-2-oxobutanoate dehydrogenase (2-methylpropanoyl-transferring)]-phosphate phosphohy-

drolase

**Comments:** A mitochondrial enzyme associated with the 3-methyl-2-oxobutanoate dehydrogenase complex. Si-

multaneously dephosphorylates and activates EC 1.2.4.4 3-methyl-2-oxobutanoate dehydrogenase

(2-methylpropanoyl-transferring), that has been inactivated by phosphorylation.

**References:** [742, 2395]

[EC 3.1.3.52 created 1986]

EC 3.1.3.53

Accepted name: [myosin-light-chain] phosphatase

> Reaction: [myosin light-chain] phosphate +  $H_2O$  = [myosin light-chain] + phosphate

myosin light chain kinase phosphatase; myosin phosphatase; myosin phosphatase; protein phos-Other name(s):

phatase 2A; myosin-light-chain-phosphatase

**Systematic name:** [myosin-light-chain]-phosphate phosphohydrolase

**Comments:** The enzyme is composed of three subunits. The holoenzyme dephosphorylates myosin light chains

and EC 2.7.11.18, myosin-light-chain kinase, but not myosin; the catalytic subunit acts on all three

substrates.

References: [2243]

[EC 3.1.3.53 created 1986]

EC 3.1.3.54

Accepted name: fructose-2,6-bisphosphate 6-phosphatase

> $\beta$ -D-fructose 2,6-bisphosphate + H<sub>2</sub>O =  $\beta$ -D-fructofuranose 2-phosphate + phosphate **Reaction:**

fructose 2,6-bisphosphate-6-phosphohydrolase; fructose-2,6-bisphosphate 6-phosphohydrolase; D-Other name(s):

fructose-2,6-bisphosphate 6-phosphohydrolase

**Systematic name:** β-D-fructose-2,6-bisphosphate 6-phosphohydrolase **Comments:** 

cf. EC 3.1.3.46 fructose-2,6-bisphosphate 2-phosphatase.

**References:** [2332, 2333]

[EC 3.1.3.54 created 1989]

EC 3.1.3.55

Accepted name: caldesmon-phosphatase

> caldesmon phosphate  $+ H_2O = caldesmon + phosphate$ **Reaction:**

Other name(s): SMP-I; smooth muscle caldesmon phosphatase **Systematic name:** caldesmon-phosphate phosphohydrolase

**Comments:** Dephosphorylation activates the calmodulin- and actin-binding ability of the protein caldesmon.

**References:** [2078]

[EC 3.1.3.55 created 1989]

## EC 3.1.3.56

**Accepted name:** inositol-polyphosphate 5-phosphatase

**Reaction:** (1) D-myo-inositol 1,4,5-trisphosphate + H<sub>2</sub>O = myo-inositol 1,4-bisphosphate + phosphate

(2) 1D-myo-inositol 1,3,4,5-tetrakisphosphate + H<sub>2</sub>O = 1D-myo-inositol 1,3,4-trisphosphate + phos-

phate

**Other name(s):** type I inositol-polyphosphate phosphatase; inositol trisphosphate phosphomonoesterase;

Ins $P_3$ /Ins(1,3,4,5) $P_4$  5-phosphatase; inosine triphosphatase; D-myo-inositol 1,4,5-triphosphate 5-phosphatase; D-myo-inositol 1,4,5-trisphosphate-monoesterase; inositol phosphate 5-phosphomonoesterase; inositol-1,4,5-trisphosphate/1,3,4,5-tetrakisphosphate 5-phosphatase; Ins(1,4,5) $P_3$  5-phosphatase; D-myo-inositol(1,4,5)/(1,3,4,5)-polyphosphate 5-phosphatase; inositol 1,4,5-trisphosphate phosphatase; inositol polyphosphate-5-phosphatase; myo-inositol-1,4,5-trisphosphate 5-phosphatase; inositol-1,4,5-trisphosphate 5-phosphatase; myo-inositol-1,4,5-trisphosphate 5-phosphatase; myo-inositol-1,4,5-trisphosphate

phosphatase

**Systematic name:** 1D-myo-inositol-1,4,5-trisphosphate 5-phosphohydrolase

**Comments:** One mammalian isoform is known. This enzyme is distinguished from the family of enzymes classi-

fied under EC 3.1.3.36, phosphoinositide 5-phosphatase, by its inability to dephosphorylate inositol

lipids.

**References:** [643, 714, 3214, 3060]

[EC 3.1.3.56 created 1989, modified 2002]

#### EC 3.1.3.57

**Accepted name:** inositol-1,4-bisphosphate 1-phosphatase

**Reaction:** 1D-myo-inositol 1,4-bisphosphate + H<sub>2</sub>O = 1D-myo-inositol 4-phosphate + phosphate

**Other name(s):** inositol-polyphosphate 1-phosphatase

**Systematic name:** 1D-*myo*-inositol-1,4-bisphosphate 1-phosphohydrolase

**Comments:** The enzyme acts on inositol 1,4-bisphosphate and inositol 1,3,4-trisphosphate (forming inositol 3,4-

bisphosphate) with similar  $V_{max}$  values for both substrates, but with a five-times higher affinity for the bisphosphate. Does not act on inositol 1-phosphate, inositol 1,4,5-trisphosphate or inositol 1,3,4,5-

tetrakisphosphate.

**References:** [204, 481, 1251]

[EC 3.1.3.57 created 1989, modified 2002]

# EC 3.1.3.58

Accepted name: sugar-terminal-phosphatase

**Reaction:** D-glucose 6-phosphate  $+ H_2O = D$ -glucose + phosphate

**Other name(s):** xylitol-5-phosphatase

**Systematic name:** sugar-ω-phosphate phosphohydrolase

Comments: Acts on sugars and polyols phosphorylated on the terminal carbon, with a preference for sugars with

a D-erythro-configuration, e.g. good substrates are glucose 6-phosphate, mannose 6-phosphate, 6-

phosphogluconate, erythrose 4-phosphate and xylitol 5-phosphate.

**References:** [1734]

[EC 3.1.3.58 created 1989]

### EC 3.1.3.59

**Accepted name:** alkylacetylglycerophosphatase

**Reaction:** 1-alkyl-2-acetyl-sn-glycero-3-phosphate +  $H_2O$  = 1-alkyl-2-acetyl-sn-glycerol + phosphate

**Other name(s):** 1-alkyl-2-lyso-sn-glycero-3-P:acetyl-CoA acetyltransferase; alkylacetylglycerophosphate phosphatase

**Systematic name:** 1-alkyl-2-acetyl-*sn*-glycero-3-phosphate phosphohydrolase

**Comments:** Involved in the biosynthesis of thrombocyte activating factor in animal tissues.

**References:** [1632]

### [EC 3.1.3.59 created 1989]

#### EC 3.1.3.60

**Accepted name:** phospho*enol*pyruvate phosphatase

**Reaction:** phospho*enol*pyruvate +  $H_2O$  = pyruvate + phosphate

**Other name(s):** PEP phosphatase

**Systematic name:** phospho*enol*pyruvate phosphohydrolase

**Comments:** Also acts, but more slowly, on a wide range of other monophosphates.

**References:** [664, 1788, 1789]

[EC 3.1.3.60 created 1992]

[3.1.3.61 Deleted entry. inositol-1,4,5-trisphosphate 1-phosphatase, as its existence has not been established]

[EC 3.1.3.61 created 1992, deleted 2002]

#### EC 3.1.3.62

**Accepted name:** multiple inositol-polyphosphate phosphatase

**Reaction:** myo-inositol hexakisphosphate +  $H_2O = myo$ -inositol pentakisphosphate (mixed isomers) + phosphate

**Other name(s):** inositol (1,3,4,5)-tetrakisphosphate 3-phosphatase; inositol 1,3,4,5-tetrakisphosphate 3-

phosphomonoesterase; inositol 1,3,4,5-tetrakisphosphate-5-phosphomonoesterase; inositol tetrakisphosphate phosphomonoesterase; inositol-1,3,4,5-tetrakisphosphate 3-phosphatase; MIPP

**Systematic name:** 1D-*myo*-inositol-hexakisphosphate 5-phosphohydrolase

**Comments:** This enzyme exists in two isoforms. It also acts on  $Ins(1,3,4,5)P_4$  to yield  $Ins(1,4,5)P_3$ .

**References:** [515, 501]

[EC 3.1.3.62 created 1992, modified 2002]

## EC 3.1.3.63

**Accepted name:** 2-carboxy-D-arabinitol-1-phosphatase

**Reaction:** 2-carboxy-D-arabinitol 1-phosphate +  $H_2O = 2$ -carboxy-D-arabinitol + phosphate

Other name(s): 2-carboxyarabinitol 1-phosphatase; 2-carboxy-D-arabinitol 1-phosphate phosphohydrolase

**Systematic name:** 2-carboxy-D-arabinitol-1-phosphate 1-phosphohydrolase

**References:** [2513]

[EC 3.1.3.63 created 1992]

## EC 3.1.3.64

**Accepted name:** phosphatidylinositol-3-phosphatase

**Reaction:** 1-phosphatidyl-1D-myo-inositol 3-phosphate + H<sub>2</sub>O = 1-phosphatidyl-1D-myo-inositol + phosphate

Other name(s): inositol-1,3-bisphosphate 3-phosphatase; inositol 1,3-bisphosphate phosphatase; inositol-

polyphosphate 3-phosphatase; D-myo-inositol-1,3-bisphosphate 3-phosphohydrolase; phosphatidyl-

3-phosphate 3-phosphohydrolase

**Systematic name:** 1-phosphatidyl-1D-*myo*-inositol-3-phosphate 3-phosphohydrolase

**Comments:** This enzyme still works when the 2,3-bis(acyloxy)propyl group is removed, i.e., it hydrolyses

 $Ins(1,3)P_2$  to Ins-1-P.

**References:** [1711, 355]

[EC 3.1.3.64 created 1992, [EC 3.1.3.65 created 1992, incorporated 2002], modified 2002]]

[3.1.3.65 Deleted entry. inositol-1,3-bisphosphate 3-phosphatase. Now included with EC 3.1.3.64, phosphatidylinositol-3-phosphatase]

[EC 3.1.3.65 created 1992, deleted 2002]

# EC 3.1.3.66

**Accepted name:** phosphatidylinositol-3,4-bisphosphate 4-phosphatase

**Reaction:** 1-phosphatidyl-myo-inositol 3,4-bisphosphate +  $H_2O$  = 1-phosphatidyl-1D-myo-inositol 3-phosphate

+ phosphate

**Other name(s):** inositol-3,4-bisphosphate 4-phosphatase; D-myo-inositol-3,4-bisphosphate 4-phosphohydrolase;

phosphoinositide 4-phosphatase; inositol polyphosphate 4-phosphatase; inositol polyphosphate 4-

phosphatase type II

**Systematic name:** 1-phosphatidyl-1D-*myo*-inositol-3,4-bisphosphate 4-phosphohydrolase

**Comments:** Mg<sup>2+</sup>-independent. This enzyme still works when the 2,3-bis(acyloxy)propyl group is removed, i.e.,

it hydrolyses  $Ins(1,3,4)P_3$  to  $Ins(1,3)P_2$ . It also converts  $Ins(3,4)P_2$  into Ins-3-P.

**References:** [1194, 2114, 2113]

[EC 3.1.3.66 created 1992, modified 2002]

#### EC 3.1.3.67

**Accepted name:** phosphatidylinositol-3,4,5-trisphosphate 3-phosphatase

**Reaction:** 1-phosphatidyl-1D-myo-inositol 3,4,5-trisphosphate + H<sub>2</sub>O = 1-phosphatidyl-1D-myo-inositol 4,5-

bisphosphate + phosphate

Other name(s): PTEN; MMAC1; phosphatidylinositol-3,4,5-trisphosphate 3-phosphohydrolase 1-phosphatidyl-1D-*myo*-inositol-3,4,5-trisphosphate 3-phosphohydrolase

**Comments:** Requires Mg<sup>2+</sup>. Does not dephosphorylate inositol 4,5-bisphosphate. This enzyme still works when

the 2,3-bis(acyloxy) propyl group is removed, i.e., it hydrolyses  $Ins(1,3,4,5)P_4$  to  $Ins(1,4,5)P_3$ 

**References:** [1351, 2824]

[EC 3.1.3.67 created 1999, modified 2002]

### EC 3.1.3.68

**Accepted name:** 2-deoxyglucose-6-phosphatase

**Reaction:** 2-deoxy-D-glucose 6-phosphate  $+ H_2O = 2$ -deoxy-D-glucose + phosphate

**Other name(s):** 2-deoxyglucose-6-phosphate phosphatase

**Systematic name:** 2-deoxy-D-glucose-6-phosphate phosphohydrolase

**Comments:** Also active towards fructose 1-phosphate

**References:** [1332, 2371]

[EC 3.1.3.68 created 1999]

## EC 3.1.3.69

Accepted name: glucosylglycerol 3-phosphatase

**Reaction:** 2-O-( $\alpha$ -D-glucosyl)-sn-glycerol-3-phosphate + H<sub>2</sub>O = 2-O-( $\alpha$ -D-glucopyranosyl)glycerol + phosphate Other name(s): salt tolerance protein A; StpA; 2-( $\beta$ -D-glucosyl)-sn-glycerol-3-phosphate phosphohydrolase (incor-

rect)

**Systematic name:**  $2-O-(\alpha-D-glucopyranosyl)-sn-glycerol-3-phosphate phosphohydrolase$ 

**Comments:** Acts with EC 2.4.1.213 (glucosylglycerol-phosphate synthase) to form glucosylglycerol, an osmolyte

that endows cyanobacteria with resistance to salt.

**References:** [1018, 1019, 1020]

[EC 3.1.3.69 created 2001, modified 2015]

# EC 3.1.3.70

**Accepted name:** mannosyl-3-phosphoglycerate phosphatase

**Reaction:** 2-O-( $\alpha$ -D-mannosyl)-3-phosphoglycerate + H<sub>2</sub>O = 2-O-( $\alpha$ -D-mannosyl)-D-glycerate + phosphate

**Systematic name:** 2-O-( $\alpha$ -D-mannosyl)-3-phosphoglycerate phosphohydrolase

Comments: Requires  $Mg^{2+}$ . The enzyme from *Pyrococcus horikoshii* is specific for  $\alpha$ -D-mannosyl-3-

phosphoglycerate and forms part of the pathway for the synthesis of mannosylglycerate.

**References:** [701]

[EC 3.1.3.70 created 2002]

EC 3.1.3.71

**Accepted name:** 2-phosphosulfolactate phosphatase

**Reaction:** (2R)-2-phospho-3-sulfolactate +  $H_2O = (2R)$ -3-sulfolactate + phosphate **Other name(s):** (2R)-phosphosulfolactate phosphohydrolase; ComB phosphatase

**Systematic name:** (*R*)-2-phospho-3-sulfolactate phosphohydrolase

**Comments:** Requires Mg<sup>2+</sup>. The enzyme from *Methanococcus jannaschii* acts on both stereoisoimers of the sub-

strate and also hydrolyses a number of phosphate monoesters of (S)-2-hydroxycarboxylic acids, including 2-phosphomalate, 2-phospholactate and 2-phosphoglycolate. This enzyme can also hydrolyse phosphate monoesters of (R)-2-hydroxycarboxylic acids such as (S)-2-phospho-3-sulfolactate and (R)-

2-phosphomalate, which, presumably, bind to the enzyme in opposite orientations.

**References:** [962]

[EC 3.1.3.71 created 2002]

EC 3.1.3.72

**Accepted name:** 5-phytase

**Reaction:** myo-inositol hexakisphosphate +  $H_2O = 1L$ -myo-inositol 1,2,3,4,6-pentakisphosphate + phosphate

**Systematic name:** *myo*-inositol-hexakisphosphate 5-phosphohydrolase

**Comments:** The enzyme attacks the product of the above reaction more slowly to yield  $Ins(1,2,3)P_3$ .

**References:** [163]

[EC 3.1.3.72 created 2002]

EC 3.1.3.73

**Accepted name:** adenosylcobalamin/α-ribazole phosphatase

**Reaction:** (1) adenosylcobalamin 5'-phosphate +  $H_2O$  = coenzyme  $B_{12}$  + phosphate

(2)  $\alpha$ -ribazole 5'-phosphate + H<sub>2</sub>O =  $\alpha$ -ribazole + phosphate

Other name(s): CobC; adenosylcobalamin phosphatase; α-ribazole phosphatase systematic name: adenosylcobalamin/α-ribazole-5'-phosphate phosphohydrolase

**Comments:** This enzyme catalyses the last step in the anaerobic (early cobalt insertion) pathway of adenosyl-

cobalamin biosynthesis, characterized in Salmonella enterica [3318]. It also participates in a salvage

pathway that recycles cobinamide into adenosylcobalamin [2213].

**References:** [2213, 3131, 3318]

[EC 3.1.3.73 created 2004, modified 2011]

EC 3.1.3.74

**Accepted name:** pyridoxal phosphatase

**Reaction:** pyridoxal 5'-phosphate +  $H_2O$  = pyridoxal + phosphate

Other name(s): vitamine B<sub>6</sub> (pyridoxine) phosphatase; PLP phosphatase; vitamin B<sub>6</sub>-phosphate phosphatase; PNP

phosphatase

**Systematic name:** pyridoxal-5'-phosphate phosphohydrolase

**Comments:** Requires Mg<sup>2+</sup>. This enzyme is specific for phosphorylated vitamin B<sub>6</sub> compounds: it acts not only

on pyridoxal phosphate (PLP), but also on pyridoxine phosphate (PNP), pyridoxamine phosphate (PMP), 4-pyridoxic acid phosphate and 4-deoxypyridoxine phosphate. This reaction can also be carried out by EC 3.1.3.1 (alkaline phosphatase) and EC 3.1.3.2 (acid phosphatase), but these enzymes

have very broad substrate specificities.

**References:** [794, 795, 1307]

### [EC 3.1.3.74 created 2004]

EC 3.1.3.75

**Accepted name:** phosphoethanolamine/phosphocholine phosphatase

**Reaction:** (1) *O*-phosphoethanolamine +  $H_2O$  = ethanolamine + phosphate

(2) phosphocholine +  $H_2O$  = choline + phosphate

Other name(s): PHOSPHO1; 3X11A

**Systematic name:** phosphoethanolamine phosphohydrolase

**Comments:** Requires active site  $Mg^{2+}$  but also works, to a lesser extent, with  $Co^{2+}$  and  $Mn^{2+}$ . The enzyme is

highly specific for phosphoethanolamine and phosphocholine.

**References:** [1189, 2771, 2438]

[EC 3.1.3.75 created 2004]

EC 3.1.3.76

Accepted name: lipid-phosphate phosphatase

 $\textbf{Reaction:} \hspace{0.2in} (9S,10S)-10-\text{hydroxy-}9-(\text{phosphooxy}) octade can oate + H_2O = (9S,10S)-9,10-\text{dihydroxyoctade can oate} + H_2O = (9S,10S)-9,10-\text{di$ 

+ phosphate

**Other name(s):** hydroxy fatty acid phosphatase; dihydroxy fatty acid phosphatase; hydroxy lipid phos-

phatase; sEH (ambiguous); soluble epoxide hydrolase (ambiguous); (9S,10S)-10-hydroxy-9-

(phosphonooxy)octadecanoate phosphohydrolase

**Systematic name:** (9*S*,10*S*)-10-hydroxy-9-(phosphooxy)octadecanoate phosphohydrolase

**Comments:** Requires Mg<sup>2+</sup> for maximal activity. The enzyme from mammals is a bifunctional enzyme: the N-

terminal domain exhibits lipid-phosphate-phosphatase activity and the C-terminal domain has the activity of EC 3.3.2.10, soluble epoxide hydrolase (sEH) [2074]. The best substrates for this enzyme are 10-hydroxy-9-(phosphooxy)octadecanoates, with the *threo*- form being a better substrate than the *erythro*- form [2074]. The phosphatase activity is not found in plant sEH or in EC 3.3.2.9, microsomal

epoxide hydrolase, from mammals [2074].

**References:** [2074, 507, 1977, 2959, 2073, 2749, 952]

[EC 3.1.3.76 created 2006]

EC 3.1.3.77

Accepted name: acireductone synthase

**Reaction:** 5-(methylsulfanyl)-2,3-dioxopentyl phosphate  $+ H_2O = 1,2$ -dihydroxy-5-(methylsulfanyl)pent-1-en-3-

one + phosphate (overall reaction)

(1a) 5-(methylsulfanyl)-2,3-dioxopentyl phosphate = 2-hydroxy-5-(methylsulfanyl)-3-oxopent-1-enyl

phosphate (probably spontaneous)

(1b) 2-hydroxy-5-(methylsulfanyl)-3-oxopent-1-enyl phosphate +  $H_2O$  = 1,2-dihydroxy-5-

(methylsulfanyl)pent-1-en-3-one + phosphate

Other name(s): E1; E-1 enolase-phosphatase; 5-(methylthio)-2,3-dioxopentyl-phosphate phosphohydrolase (isomeriz-

ing)

**Systematic name:** 5-(methylsulfanyl)-2,3-dioxopentyl-phosphate phosphohydrolase (isomerizing)

**Comments:** This bifunctional enzyme first enolizes the substrate to form the intermediate 2-hydroxy-5-

(methylsulfanyl)-3-oxopent-1-enyl phosphate, which is then dephosphorylated to form the acire-ductone 1,2-dihydroxy-5-(methylsulfanyl)pent-1-en-3-one [3215]. The acireductone represents a branch point in the methione-salvage pathway as it is used in the formation of formate, CO and 3-(methylsulfanyl)propanoate by EC 1.13.11.53 [acireductone dioxygenase (Ni<sup>2+</sup>-requiring)] and of formate and 4-(methylsulfanyl)-2-oxobutanoate either by a spontaneous reaction under aerobic condi-

tions or by EC 1.13.11.54 acireductone dioxygenase [iron(II)-requiring] [2016, 3215].

**References:** [2016, 3215, 3109]

[EC 3.1.3.77 created 2006]

## EC 3.1.3.78

**Accepted name:** phosphatidylinositol-4,5-bisphosphate 4-phosphatase

**Reaction:** 1-phosphatidyl-1D-myo-inositol 4,5-bisphosphate + H<sub>2</sub>O = 1-phosphatidyl-1D-myo-inositol 5-

phosphate + phosphate

Other name(s): phosphatidylinositol-4,5-bisphosphate 4-phosphatase I; phosphatidylinositol-4,5-bisphosphate 4-phosphatase I;

phosphatase II; type I PtdIns-4,5-P<sub>2</sub> 4-Ptase; type II PtdIns-4,5-P<sub>2</sub> 4-Ptase; IpgD; PtdIns-4,5-P<sub>2</sub> 4-phosphatase type I; PtdIns-4,5-P<sub>2</sub> 4-phosphatase type II; type I phosphatidylinositol-4,5-bisphosphate

4-phosphatase; type 1 4-phosphatase

**Systematic name:** 1-phosphatidyl-1D-*myo*-inositol-4,5-bisphosphate 4-phosphohydrolase

**Comments:** Two pathways exist in mammalian cells to degrade 1-phosphatidyl-1D-myo-inositol 4,5-bisphosphate

[PtdIns(4,5) $P_2$ ] [3013]. One is catalysed by this enzyme and the other by EC 3.1.3.36, phosphoinositide 5-phosphatase, where the product is PtdIns4P. The enzyme from human is specific for PtdIns(4,5) $P_2$  as substrate, as it cannot use PtdIns(3,4,5) $P_3$ , PtdIns(3,4) $P_2$ , PtdIns(3,5) $P_2$ , PtdIns5 $P_3$ , PtdIns4 $P_3$  or PtdIns3 $P_3$  [3013]. In humans, the enzyme is localized to late endosomal/lysosomal membranes [3013]. It can control nuclear levels of PtdIns5 $P_3$  and thereby control p53-dependent apoptosis

[3349].

**References:** [2082, 3013, 3349, 1833]

[EC 3.1.3.78 created 2008]

## EC 3.1.3.79

Accepted name: mannosylfructose-phosphate phosphatase

**Reaction:**  $\beta$ -D-fructofuranosyl- $\alpha$ -D-mannopyranoside  $6^F$ -phosphate +  $H_2O = \beta$ -D-fructofuranosyl- $\alpha$ -D-

mannopyranoside + phosphate

**Other name(s):** mannosylfructose-6-phosphate phosphatase; MFPP

Systematic name: β-D-fructofuranosyl-α-D-mannopyranoside-6F-phosphate phosphohydrolase

**Comments:** This enzyme, from the soil proteobacterium and plant pathogen *Agrobacterium tumefaciens* strain

C58, requires  $Mg^{2+}$  for activity. Mannosylfructose is the major endogenous osmolyte produced by several  $\alpha$ -proteobacteria in response to osmotic stress and is synthesized by the sequential action of EC 2.4.1.246 (mannosylfructose-phosphate synthase) followed by this enzyme. While mannosylfructose 6-phosphate is the physiological substrate, the enzyme can use sucrose 6-phosphate very efficiently. The F in mannosylfructose  $6^F$ -phosphate is used to indicate that the fructose residue of

sucrose carries the substituent.

**References:** [2948]

[EC 3.1.3.79 created 2009]

### EC 3.1.3.80

**Accepted name:** 2,3-bisphosphoglycerate 3-phosphatase

**Reaction:** 2,3-bisphospho-D-glycerate +  $H_2O$  = 2-phospho-D-glycerate + phosphate

Other name(s): MIPP1; 2,3-BPG 3-phosphatase

**Systematic name:** 2,3-bisphospho-D-glycerate 3-phosphohydrolase

**Comments:** This reaction is a shortcut in the Rapoport-Luebering shunt. It bypasses the reactions of EC

5.4.2.11/EC 5.4.2.12 [phosphoglycerate mutases (2,3-diphosphoglycerate-dependent and independent)] and directly forms 2-phospho-D-glycerate by removing the 3-phospho-group of 2,3-diphospho-D-glycerate [436]. The MIPP1 protein also catalyses the reaction of EC 3.1.3.62 (multiple inositol-

polyphosphate phosphatase).

**References:** [436]

[EC 3.1.3.80 created 2010]

#### EC 3.1.3.81

**Accepted name:** diacylglycerol diphosphate phosphatase

**Reaction:** 1,2-diacyl-sn-glycerol 3-diphosphate + H<sub>2</sub>O = 1,2-diacyl-sn-glycerol 3-phosphate + phosphate

Other name(s): DGPP phosphatase; DGPP phosphohydrolase; DPP1; DPPL1; DPPL2; PAP2; pyrophosphate phos-

phatase

**Systematic name:** 1,2-diacyl-*sn*-glycerol 3-phosphate phosphohydrolase

**Comments:** The bifunctional enzyme catalyses the dephosphorylation of diacylglycerol diphosphate to phos-

phatidate and the subsequent dephosphorylation of phosphatidate to diacylglycerol (*cf.* phosphatidate phosphatase (EC 3.1.3.4)). It regulates intracellular levels of diacylglycerol diphosphate and phosphatidate, phospholipid molecules believed to play a signalling role in stress response [1040]. The phosphatase activity of the bifunctional enzyme is Mg<sup>2+</sup>-independent and N-ethylmaleimide-insensitive and is distinct from the Mg<sup>2+</sup>-dependent and N-ethylmaleimide-sensitive enzyme EC 3.1.3.4 (phosphatidate phosphatase) [372]. The diacylglycerol pyrophosphate phosphatase activity in *Saccharomyces cerevisiae* is induced by zinc depletion, by inositol supplementation, and when cells

enter the stationary phase [2206].

**References:** [615, 614, 3223, 2206, 372, 1040]

[EC 3.1.3.81 created 2010]

EC 3.1.3.82

**Accepted name:** D-*glycero*-β-D-*manno*-heptose 1,7-bisphosphate 7-phosphatase

**Reaction:** D-glycero- $\beta$ -D-manno-heptose 1,7-bisphosphate + H<sub>2</sub>O = D-glycero- $\beta$ -D-manno-heptose 1-phosphate

+ phosphate

**Other name(s):** *gmhB* (gene name); *yaeD* (gene name)

**Systematic name:** D-*glycero*-β-D-*manno*-heptose 1,7-bisphosphate 7-phosphohydrolase

**Comments:** The enzyme is involved in biosynthesis of ADP-L-*glycero*-β-D-*manno*-heptose, which is utilized for

assembly of the lipopolysaccharide inner core in Gram-negative bacteria. In vitro the catalytic effi-

ciency with the  $\beta$ -anomer is 100-200-fold higher than with the  $\alpha$ -anomer [3113].

**References:** [1482, 3031, 3113]

[EC 3.1.3.82 created 2010]

EC 3.1.3.83

**Accepted name:** D-glycero-α-D-manno-heptose 1,7-bisphosphate 7-phosphatase

**Reaction:** D-glycero- $\alpha$ -D-manno-heptose 1,7-bisphosphate + H<sub>2</sub>O = D-glycero- $\alpha$ -D-manno-heptose 1-

phosphate + phosphate

**Other name(s):** *gmhB* (gene name)

**Systematic name:** D-*glycero*-α-D-*manno*-heptose 1,7-bisphosphate 7-phosphohydrolase

**Comments:** The enzyme is involved in biosynthesis of GDP-D-glycero-α-D-manno-heptose, which is required for

assembly of S-layer glycoprotein in some Gram-positive bacteria. The *in vitro* catalytic efficiency of the enzyme from *Bacteroides thetaiotaomicron* is 6-fold higher with the  $\alpha$ -anomer than with the  $\beta$ -

anomer [3113].

**References:** [3113]

[EC 3.1.3.83 created 2010]

EC 3.1.3.84

**Accepted name:** ADP-ribose 1"-phosphate phosphatase

**Reaction:** ADP-D-ribose 1"-phosphate +  $H_2O$  = ADP-D-ribose + phosphate

**Other name(s):** POA1; Appr1p phosphatase; Poa1p; ADP-ribose 1"-phosphate phosphohydrolase

**Systematic name:** ADP-D-ribose 1"-phosphate phosphohydrolase

**Comments:** The enzyme is highly specific for ADP-D-ribose 1"-phosphate. Involved together with EC 3.1.4.37,

2',3'-cyclic-nucleotide 3'-phosphodiesterase, in the breakdown of adenosine diphosphate ribose 1",2"-

cyclic phosphate (Appr¿p), a by-product of tRNA splicing.

**References:** [2649]

[EC 3.1.3.84 created 2011]

### EC 3.1.3.85

**Accepted name:** glucosyl-3-phosphoglycerate phosphatase

**Reaction:** 2-O-( $\alpha$ -D-glucopyranosyl)-3-phospho-D-glycerate +  $H_2O$  = 2-O-( $\alpha$ -D-glucopyranosyl)-D-glycerate +

phosphate

**Other name**(s): GpgP protein

**Systematic name:** α-D-glucosyl-3-phospho-D-glycerate phosphohydrolase

**Comments:** The enzyme is involved in biosynthesis of 2-*O*-(α-D-glucopyranosyl)-D-glycerate via the two-step

pathway in which EC 2.4.1.266 (glucosyl-3-phosphoglycerate synthase) catalyses the conversion of GDP-glucose and 3-phospho-D-glycerate into 2-O-( $\alpha$ -D-glucopyranosyl)-3-phospho-D-glycerate, which is then converted to 2-O-( $\alpha$ -D-glucopyranosyl)-D-glycerate by glucosyl-3-phosphoglycerate phosphatase. *In vivo* the enzyme catalyses the dephosphorylation of 2-O-( $\alpha$ -D-mannopyranosyl)-3-phospho-D-glycerate with lower efficiency [493, 494]. Divalent metal ions (Mg<sup>2+</sup>, Mn<sup>2+</sup> or Co<sup>2+</sup>)

stimulate activity [493, 494].

**References:** [493, 494, 1885]

[EC 3.1.3.85 created 2011]

#### EC 3.1.3.86

**Accepted name:** phosphatidylinositol-3,4,5-trisphosphate 5-phosphatase

**Reaction:** 1-phosphatidyl-1D-myo-inositol 3,4,5-trisphosphate + H<sub>2</sub>O = 1-phosphatidyl-1D-myo-inositol 3,4-

bisphosphate + phosphate

Other name(s): SHIP1; SHIP2; SHIP; p150Ship

**Systematic name:** 1-phosphatidyl-1D-*myo*-inositol-3,4,5-trisphosphate 5-phosphohydrolase

Comments: This enzyme hydroylses 1-phosphatidyl-1D-myo-inositol 3,4,5-trisphosphate (PtdIns(3,4,5) $P_3$ ) to pro-

duce  $PtdIns(3,4)P_2$ , thereby negatively regulating the PI3K (phosphoinositide 3-kinase) pathways. The enzyme also shows activity toward ( $PtdIns(1,3,4,5)P_4$ ) [2258]. The enzyme is involved in several

signal transduction pathways in the immune system leading to an adverse range of effects.

**References:** [1709, 533, 925, 650, 2258]

[EC 3.1.3.86 created 2011]

## EC 3.1.3.87

**Accepted name:** 2-hydroxy-3-keto-5-methylthiopentenyl-1-phosphate phosphatase

**Reaction:** 2-hydroxy-5-(methylsulfanyl)-3-oxopent-1-enyl phosphate + H<sub>2</sub>O = 1,2-dihydroxy-5-

(methylsulfanyl)pent-1-en-3-one + phosphate

Other name(s): HK-MTPenyl-1-P phosphatase; MtnX; YkrX; 2-hydroxy-5-(methylthio)-3-oxopent-1-enyl phosphate

phosphohydrolase

**Systematic name:** 2-hydroxy-5-(methylsulfanyl)-3-oxopent-1-enyl phosphate phosphohydrolase

**Comments:** The enzyme participates in the methionine salvage pathway in *Bacillus subtilis* [86]. In some species

a single bifunctional enzyme, EC 3.1.3.77, acireductone synthase, catalyses both this reaction and EC

5.3.2.5, 2,3-diketo-5-methylthiopentyl-1-phosphate enolase [2016].

**References:** [2016, 86]

[EC 3.1.3.87 created 2012]

## EC 3.1.3.88

**Accepted name:** 5"-phosphoribostamycin phosphatase

**Reaction:** 5''-phosphoribostamycin +  $H_2O$  = ribostamycin + phosphate

Other name(s): *btrP* (gene name); *neoI* (gene name)

Systematic name: 5"-phosphoribostamycin phosphohydrolase

Comments: Involved in the biosynthetic pathways of several clinically important aminocyclitol antibiotics, includ-

ing ribostamycin, neomycin and butirosin. No metal is required for activity.

**References:** [1543]

### [EC 3.1.3.88 created 2012]

EC 3.1.3.89

**Accepted name:** 5'-deoxynucleotidase

**Reaction:** a 2'-deoxyribonucleoside 5'-monophosphate +  $H_2O$  = a 2'-deoxyribonucleoside + phosphate

**Other name(s):** *yfbR* (gene name)

**Systematic name:** 2'-deoxyribonucleoside 5'-monophosphate phosphohydrolase

**Comments:** The enzyme, characterized from the bacterium *Escherichia coli*, shows strict specificity towards de-

oxyribonucleoside 5'-monophosphates and does not dephosphorylate 5'-ribonucleotides or ribonucleoside 3'-monophosphates. A divalent metal cation is required for activity, with cobalt providing the

highest activity.

**References:** [2327, 3342]

[EC 3.1.3.89 created 2013]

EC 3.1.3.90

**Accepted name:** maltose 6'-phosphate phosphatase

**Reaction:** maltose 6'-phosphate +  $H_2O$  = maltose + phosphate **Other name(s):** maltose 6'-P phosphatase; mapP (gene name) **Systematic name:** maltose 6'-phosphate phosphohydrolase

Comments: The enzyme from the bacterium Enterococcus faecalis also has activity with the sucrose isomer tura-

nose 6'-phosphate ( $\alpha$ -D-glucopyranosyl-( $1\rightarrow 3$ )-D-fructose 6-phosphate).

**References:** [1947]

[EC 3.1.3.90 created 2013]

EC 3.1.3.91

Accepted name: 7-methylguanosine nucleotidase

**Reaction:** (1)  $N^7$ -methyl-GMP + H<sub>2</sub>O =  $N^7$ -methyl-guanosine + phosphate

(2)  $CMP + H_2O = cytidine + phosphate$ 

Other name(s): cytosolic nucleotidase III-like; cNIII-like;  $N^7$ -methylguanylate 5'-phosphatase

**Systematic name:**  $N^7$ -methyl-GMP phosphohydrolase

**Comments:** The enzyme also has low activity with  $N^7$ -methyl-GDP, producing  $N^7$ -methyl-GMP. Does not accept

AMP or GMP, and has low activity with UMP.

**References:** [334]

[EC 3.1.3.91 created 2013]

EC 3.1.3.92

**Accepted name:** kanosamine-6-phosphate phosphatase

**Reaction:** kanosamine 6-phosphate  $+ H_2O = \text{kanosamine} + \text{phosphate}$ 

**Other name(s):** *ntdB* (gene name)

Systematic name: kanosamine-6-phosphate phosphohydrolase

Comments: The enzyme, found in the bacterium *Bacillus subtilis*, is involved in a kanosamine biosynthesis path-

way.

**References:** [3064]

[EC 3.1.3.92 created 2013]

EC 3.1.3.93

**Accepted name:** L-galactose 1-phosphate phosphatase

**Reaction:**  $\beta$ -L-galactose 1-phosphate +  $H_2O$  = L-galactose + phosphate

**Other name(s):** VTC4 (gene name) (ambiguous); IMPL2 (gene name) (ambiguous)

**Systematic name:** β-L-galactose-1-phosphate phosphohydrolase

**Comments:** The enzyme from plants also has the activity of EC 3.1.3.25, inositol-phosphate phosphatase. The en-

zymes have very low activity with D-galactose 1-phosphate (cf. EC 3.1.3.94, D-galactose 1-phosphate

phosphatase).

**References:** [1590, 2947, 2262]

[EC 3.1.3.93 created 2014]

EC 3.1.3.94

**Accepted name:** D-galactose 1-phosphate phosphatase

**Reaction:**  $\alpha$ -D-galactose 1-phosphate + H<sub>2</sub>O = D-galactose + phosphate

**Systematic name:** α-D-galactose-1-phosphate phosphohydrolase

**Comments:** The human enzyme also has the activity of EC 3.1.3.25, inositol-phosphate phosphatase. The enzyme

has very low activity with L-galactose 1-phosphate (cf. EC 3.1.3.93, L-galactose 1-phosphate phos-

phatase).

**References:** [2241]

[EC 3.1.3.94 created 2014]

EC 3.1.3.95

**Accepted name:** phosphatidylinositol-3,5-bisphosphate 3-phosphatase

**Reaction:** 1-phosphatidyl-1D-myo-inositol 3,5-bisphosphate + H<sub>2</sub>O = 1-phosphatidyl-1D-myo-inositol 5-

phosphate + phosphate

Other name(s): MTMR; PtdIns-3,5-P<sub>2</sub> 3-Ptase

**Systematic name:** 1-phosphatidyl-1D-*myo*-inositol-3,5-bisphosphate 3-phosphohydrolase

**Comments:** The enzyme is found in both plants and animals. It also has the activity of EC 3.1.3.64

(phosphatidylinositol-3-phosphatase).

**References:** [3093, 194, 619]

[EC 3.1.3.95 created 2014]

EC 3.1.3.96

**Accepted name:** pseudouridine 5'-phosphatase

**Reaction:** pseudouridine 5'-phosphate + H<sub>2</sub>O = pseudouridine + phosphate **Other name(s):** pseudouridine 5'-monophosphatase; 5'-PsiMPase; HDHD1

**Systematic name:** pseudouridine 5'-phosphohydrolase

**Comments:** Requires  $Mg^{2+}$  for activity.

**References:** [2324]

[EC 3.1.3.96 created 2014]

EC 3.1.3.97

**Accepted name:** 3',5'-nucleoside bisphosphate phosphatase

**Reaction:** nucleoside 3',5'-bisphosphate +  $H_2O$  = nucleoside 5'-phosphate + phosphate

**Systematic name:** nucleoside-3',5'-bisphosphate 3'-phosphohydrolase

**Comments:** The enzyme, characterized from the bacterium *Chromobacterium violaceum*, has similar cat-

alytic efficiencies with all the bases. The enzyme has similar activity with ribonucleoside and 2'-deoxyribonucleoside 3',5'-bisphosphates, but shows no activity with nucleoside 2',5'-bisphosphates

(cf. EC 3.1.3.7, 3'(2'), 5'-bisphosphate nucleotidase).

**References:** [516]

[EC 3.1.3.97 created 2015]

[3.1.3.98 Transferred entry. geranyl diphosphate phosphohydrolase, transferred to EC 3.6.1.68, geranyl diphosphate phosphohydrolase]

#### [EC 3.1.3.98 created 2015, deleted 2016]

## EC 3.1.3.99

Accepted name: IMP-specific 5'-nucleotidase Reaction: IMP +  $H_2O$  = inosine + phosphate

Other name(s): ISN1 (gene name)

**Systematic name:** inosine 5'-phosphate phosphohydrolase

**Comments:** The enzyme, isolated from the yeast *Saccharomyces cerevisiae*, is highly specific for inosine 5'-

phosphate, and has no detectable activity with other purine and pyrimidine nucleotides. Requires di-

valent metals, such as Mg<sup>2+</sup>, Co<sup>2+</sup> or Mn<sup>2+</sup>.

**References:** [1270, 1271]

[EC 3.1.3.99 created 2016]

#### EC 3.1.3.100

Accepted name: thiamine phosphate phosphatase

**Reaction:** thiamine phosphate  $+ H_2O =$  thiamine + phosphate

Systematic name: thiamine phosphate phosphohydrolase

**Comments:** The enzyme participates in the thiamine biosynthesis pathway in eukaryotes and a few prokaryotes.

These organisms lack EC 2.7.4.16, thiamine-phosphate kinase, and need to convert thiamine phosphate to thiamine diphosphate, the active form of the vitamin, by first removing the phosphate group,

followed by diphosphorylation by EC 2.7.6.2, thiamine diphosphokinase.

**References:** [2517, 1504, 2596, 1997, 1500, 1920]

[EC 3.1.3.100 created 2016]

### EC 3.1.3.101

**Accepted name:** validoxylamine A 7'-phosphate phosphatase

**Reaction:** validoxylamine A 7'-phosphate +  $H_2O$  = validoxylamine A + phosphate

**Other name(s):** *vldH* (gene name)

**Systematic name:** validoxylamine-A 7'-phosphate phosphohydrolase

Comments: The enzyme, characterized from the bacterium Streptomyces hygroscopicus subsp. limoneus, is in-

volved in the biosynthesis of the antifungal agent validamycin A.

**References:** [81]

[EC 3.1.3.101 created 2016]

## EC 3.1.3.102

Accepted name: FMN hydrolase

Reaction: FMN + H<sub>2</sub>O = riboflavin + phosphate
Other name(s): FMN phosphatase; AtcpFHy1
Systematic name: FMN phosphohydrolase

**Comments:** Requires  $Mg^{2+}$ . The enzyme, found in many isoforms purified from both bacteria and plants, is a

member of the haloacid dehalogenase superfamily. Most of the isoforms have a wide substrate speci-

ficity [1579], but isoforms specific for FMN also exist [2381].

**References:** [2516, 1579, 2381]

[EC 3.1.3.102 created 2016]

EC 3.1.3.103

Accepted name: 3-deoxy-D-glycero-D-galacto-nonulopyranosonate 9-phosphatase

**Reaction:** 3-deoxy-D-glycero-D-galacto-non-2-ulopyranosonate 9-phosphate + H<sub>2</sub>O = 3-deoxy-D-glycero-D-

*galacto*-non-2-ulopyranosonate + phosphate

Other name(s): 3-deoxy-D-glycero-D-galacto-non-2-ulopyranosonate-9-phosphate phosphatase **Systematic name:** 3-deoxy-D-glycero-D-galacto-non-2-ulopyranosonate 9-phosphohydrolase

**Comments:** The enzyme, characterized from the bacterium *Bacteroides thetaiotaomicron*, is part of the biosyn-

> thesis pathway of the sialic acid 3-deoxy-D-glycero-D-galacto-non-2-ulopyranosonate (Kdn). Kdn is abundant in extracellular glycoconjugates of lower vertebrates such as fish and amphibians, but is also

found in the capsular polysaccharides of bacteria that belong to the *Bacteroides* genus.

**References:** [3114, 1747]

[EC 3.1.3.103 created 2016]

EC 3.1.3.104

Accepted name: 5-amino-6-(5-phospho-D-ribitylamino)uracil phosphatase

Reaction: 5-amino-6-(5-phospho-D-ribitylamino)uracil + H<sub>2</sub>O = 5-amino-6-(D-ribitylamino)uracil + phosphate

Other name(s): 5-amino-6-ribitylamino-2,4(1*H*,3*H*)-pyrimidinedione 5'-phosphate phosphatase

**Systematic name:** 5-amino-6-(5-phospho-D-ribitylamino)uracil phosphohydrolase

Requires Mg<sup>2+</sup>. The enzyme, which is found in plants and bacteria, is part of a pathway for riboflavin **Comments:** 

biosynthesis. Most forms of the enzyme has a broad substrate specificity [1013, 2529].

References: [1013, 1735, 2529]

[EC 3.1.3.104 created 2016]

EC 3.1.3.105

Accepted name: *N*-acetyl-D-muramate 6-phosphate phosphatase

Reaction: N-acetyl-D-muramate 6-phosphate +  $H_2O = N$ -acetyl-D-muramate + phosphate

Other name(s): mupP (gene name)

**Systematic name:** *N*-acetyl-D-muramate 6-phosphate phosphohydrolase

**Comments:** The enzyme, characterized from Pseudomonas species, participates in a peptidoglycan salvage path-

way.

**References:** [265]

[EC 3.1.3.105 created 2017]

## EC 3.1.4 Phosphoric-diester hydrolases

EC 3.1.4.1

Accepted name: phosphodiesterase I

> Reaction: Hydrolytically removes 5'-nucleotides successively from the 3'-hydroxy termini of 3'-hydroxy-

> > terminated oligonucleotides

5'-exonuclease; 5'-phosphodiesterase; 5'-nucleotide phosphodiesterase; oligonucleate 5'-Other name(s):

> nucleotidohydrolase; 5' nucleotide phosphodiesterase/alkaline phosphodiesterase I; 5'-NPDase; 5'-PDase; 5'-PDE; 5'NPDE; alkaline phosphodiesterase; nucleotide pyrophosphatase/phosphodiesterase

I; orthophosphoric diester phosphohydrolase; PDE I; phosphodiesterase (ambiguous); exonuclease I

oligonucleotide 5'-nucleotidohydrolase **Systematic name:** 

**Comments:** Hydrolyses both ribonucleotides and deoxyribonucleotides. Has low activity towards polynucleotides.

A 3'-phosphate terminus on the substrate inhibits hydrolysis.

References: [1435]

[EC 3.1.4.1 created 1961]

### EC 3.1.4.2

Accepted name: glycerophosphocholine phosphodiesterase

**Reaction:** sn-glycero-3-phosphocholine +  $H_2O$  = choline + sn-glycerol 3-phosphate

Other name(s): glycerophosphinicocholine diesterase; glycerylphosphorylcholinediesterase; sn-glycero-3-

phosphorylcholine diesterase; glycerolphosphorylcholine phosphodiesterase; glycerophosphohydro-

lase

**Systematic name:** *sn*-glycero-3-phosphocholine glycerophosphohydrolase

**Comments:** Also acts on *sn*-glycero-3-phosphoethanolamine.

**References:** [550, 1084, 3144]

[EC 3.1.4.2 created 1961, modified 1976]

### EC 3.1.4.3

Accepted name: phospholipase C

**Reaction:** a phosphatidylcholine +  $H_2O = 1,2$ -diacyl-sn-glycerol + phosphocholine

Other name(s): lipophosphodiesterase I; lecithinase C; Clostridium welchii  $\alpha$ -toxin; Clostridium oedematiens  $\beta$ - and

 $\gamma$ -toxins; lipophosphodiesterase C; phosphatidase C; heat-labile hemolysin;  $\alpha$ -toxin

**Systematic name:** phosphatidylcholine cholinephosphohydrolase

**Comments:** The bacterial enzyme, which is a zinc protein, also acts on sphingomyelin and phosphatidylinositol;

that from seminal plasma does not act on phosphatidylinositol.

**References:** [656, 1712, 2625, 2846]

[EC 3.1.4.3 created 1961]

### EC 3.1.4.4

**Accepted name:** phospholipase D

**Reaction:** a phosphatidylcholine  $+ H_2O =$  choline + a phosphatidate **Other name(s):** lipophosphodiesterase II; lecithinase D; choline phosphatase

**Systematic name:** phosphatidylcholine phosphatidohydrolase **Comments:** Also acts on other phosphatidyl esters.

**References:** [91, 686, 1043, 2946]

### [EC 3.1.4.4 created 1961]

[3.1.4.5	Transferred entry	dearyribanuclease	Now FC 3 1 21 1	, deoxyribonuclease I	7
13.1.4.3	Transferrea emry	. aeoxyriboniaciease.	NOW EC 3.1.21.1	, aeoxyribonuciease i	1

[EC 3.1.4.5 created 1961, deleted 1978]

[3.1.4.6 Transferred entry. deoxyribonuclease II. Now EC 3.1.22.1, deoxyribonuclease II]

[EC 3.1.4.6 created 1961, deleted 1978]

[3.1.4.7 Transferred entry, micrococcal nuclease, Now EC 3.1.31.1, micrococcal nuclease]

[EC 3.1.4.7 created 1961, deleted 1978]

[3.1.4.8 Transferred entry. Aspergillus oryzae ribonuclease. Now EC 3.1.27.3, ribonuclease T<sub>1</sub>]

[EC 3.1.4.8 created 1961, transferred 1965 to EC 2.7.7.26, reinstated 1972, deleted 1978]

[3.1.4.9 Transferred entry. nucleate endonuclease. Now EC 3.1.30.2, Serratia marcescens nuclease]

[EC 3.1.4.9 created 1965, deleted 1978]

[3.1.4.10 Transferred entry. 1-phosphatidylinositol phosphodiesterase. Now EC 4.6.1.13, phosphatidylinositol diacylglycerollyase. As there is no hydrolysis of the inositol 1,2-cyclic phosphate formed, previous classification of the enzyme as a hydrolase was incorrect]

[EC 3.1.4.10 created 1972, modified 1976, deleted 2002]

## EC 3.1.4.11

Accepted name: phosphoinositide phospholipase C

**Reaction:** 1-phosphatidyl-1D-myo-inositol 4,5-bisphosphate + H<sub>2</sub>O = 1D-myo-inositol 1,4,5-trisphosphate +

diacylglycerol

Other name(s): triphosphoinositide phosphodiesterase; phosphoinositidase C; 1-phosphatidylinositol-4,5-

bisphosphate phosphodiesterase; monophosphatidylinositol phosphodiesterase; phosphatidylinositol phospholipase C; PI-PLC; 1-phosphatidyl-D-*myo*-inositol-4,5-bisphosphate inositoltrisphosphohydro-

lase

**Systematic name:** 1-phosphatidyl-1D-*myo*-inositol-4,5-bisphosphate inositoltrisphosphohydrolase

**Comments:** These enzymes form some of the cyclic phosphate  $Ins(cyclic 1, 2)P(4,5)P_2$  as well as  $Ins(1,4,5)P_3$ .

They show activity towards phosphatidylinositol, i.e., the activity of EC 4.6.1.13, phosphatidylinositol diacylglycerol-lyase, in vitro at high [Ca<sup>2+</sup>]. Four  $\beta$ -isoforms regulated by G-proteins, two  $\gamma$ -forms regulated by tyrosine kinases, four  $\delta$ -forms regulated at least in part by calcium and an  $\epsilon$ -form, proba-

bly regulated by the oncogene ras, have been found.

**References:** [642, 2917, 2414]

[EC 3.1.4.11 created 1972, modified 2002]

#### EC 3.1.4.12

Accepted name: sphingomyelin phosphodiesterase

**Reaction:** a sphingomyelin +  $H_2O$  = a ceramide + phosphocholine

**Other name(s):** neutral sphingomyelinase

Systematic name: sphingomyelin cholinephosphohydrolase
Comments: Has very little activity on phosphatidylcholine.

**References:** [149, 404, 1107, 1383]

[EC 3.1.4.12 created 1972]

## EC 3.1.4.13

**Accepted name:** serine-ethanolaminephosphate phosphodiesterase

**Reaction:** serine phosphoethanolamine  $+ H_2O =$  serine + ethanolamine phosphate **Other name(s):** serine ethanolamine phosphodiester phosphodiesterase; SEP diesterase

**Systematic name:** serine-phosphoethanolamine ethanolaminephosphohydrolase

**Comments:** Acts only on those phosphodiesters that have ethanolamine as a component part of the molecule.

**References:** [1022]

[EC 3.1.4.13 created 1972, modified 1976]

## EC 3.1.4.14

Accepted name: [acyl-carrier-protein] phosphodiesterase

**Reaction:** holo-[acyl-carrier protein]  $+ H_2O = 4'$ -phosphopantetheine + apo-[acyl-carrier protein] **Other name(s):** ACP hydrolyase; ACP phosphodiesterase; AcpH; [acyl-carrier-protein] 4'-pantetheine-

phosphohydrolase; holo-[acyl-carrier-protein] 4'-pantetheine-phosphohydrolase

**Systematic name:** holo-[acyl-carrier protein] 4'-pantetheine-phosphohydrolase

**Comments:** The enzyme cleaves acyl-[acyl-carrier-protein] species with acyl chains of 6-16 carbon atoms al-

though it appears to demonstrate a preference for the unacylated acyl-carrier protein (ACP) and short-chain ACPs over the medium- and long-chain species [2913]. Deletion of the gene encoding this enzyme abolishes ACP prosthetic-group turnover in vivo [2913]. Activation of apo-ACP to form the

holoenzyme is carried out by EC 2.7.8.7, holo-[acyl-carrier-protein] synthase.

**References:** [2706, 3027, 2913]

[EC 3.1.4.14 created 1972, modified 2006]

[3.1.4.15] Transferred entry, adenylyl-[glutamateammonia ligase] hydrolase. As it has been shown that the enzyme catalyses

a transfer of the adenylyl group to phosphate, the enzyme has been transferred to EC 2.7.7.89, adenylyl-[glutamateammonia ligase] phosphorylase]

#### [EC 3.1.4.15 created 1972, deleted 2015]

## EC 3.1.4.16

**Accepted name:** 2',3'-cyclic-nucleotide 2'-phosphodiesterase

**Reaction:** nucleoside 2',3'-cyclic phosphate +  $H_2O$  = nucleoside 3'-phosphate

Other name(s): ribonucleoside 2',3'-cyclic phosphate diesterase; 2',3 '-cyclic AMP phosphodiesterase; 2',3'-cyclic

nucleotidase; cyclic 2',3'-nucleotide 2'-phosphodiesterase; cyclic 2',3'-nucleotide phosphodiesterase; 2',3'-cyclic nucleoside monophosphate phosphodiesterase; 2',3'-cyclic AMP 2'-phosphohydrolase; cyclic phosphodiesterase:3'-nucleotidase; 2',3'-cyclic nucleotide phosphohydrolase; 2':3'-cyclic phosphohydrolas

phodiesterase; 2':3'-cyclic nucleotide phosphodiesterase:3'-nucleotidase

**Systematic name:** nucleoside-2',3'-cyclic-phosphate 3'-nucleotidohydrolase

**Comments:** Also hydrolyses 3'-nucleoside monophosphates and bis-4-nitrophenyl phosphate, but not 3'-

deoxynucleotides. Similar reactions are carried out by EC 3.1.27.3 (ribonuclease T<sub>1</sub>) and EC 3.1.27.5

(pancreatic ribonuclease).

**References:** [58, 59, 385, 2191, 3011]

[EC 3.1.4.16 created 1972, modified 1976]

## EC 3.1.4.17

**Accepted name:** 3',5'-cyclic-nucleotide phosphodiesterase

**Reaction:** nucleoside 3',5'-cyclic phosphate +  $H_2O$  = nucleoside 5'-phosphate

**Other name(s):** cyclic 3',5'-mononucleotide phosphodiesterase; PDE; cyclic 3',5'-nucleotide phosphodiesterase;

cyclic 3',5'-phosphodiesterase; 3',5'-nucleotide phosphodiesterase; 3':5'-cyclic nucleotide 5'-nucleotidohydrolase; 3',5'-cyclonucleotide phosphodiesterase; cyclic nucleotide phosphodiesterase; 3', 5'-cyclic nucleoside monophosphate phosphodiesterase; 3': 5'-monophosphate phosphodiesterase (cyclic CMP); cyclic 3',5-nucleotide monophosphate phosphodiesterase; nucleoside 3',5'-cyclic phosphate diesterase; nucleoside-3',5-

monophosphate phosphodiesterase

**Systematic name:** 3',5'-cyclic-nucleotide 5'-nucleotidohydrolase

Comments: Acts on 3',5'-cyclic AMP, 3',5'-cyclic dAMP, 3',5'-cyclic IMP, 3',5'-cyclic GMP and 3',5'-cyclic

CMP.

**References:** [773, 2029]

[EC 3.1.4.17 created 1972, modified 1976]

[2 1 4 10 T		1 1 1 1 1 1	II NI FO	2 1 1/1	spleen exonuclease]
[3.1.4.18]	ransterrea entry	ημοςημοσιρςτργαςρ Ι	I NOW E.C.	3 / /D /	ςπιρρη ργωημειράςρι

[EC 3.1.4.18 created 1972, deleted 1978]

[3.1.4.19 Transferred entry. oligonucleotidase. Now EC 3.1.13.3, oligonucleotidase]

[EC 3.1.4.19 created 1972, deleted 1978]

[3.1.4.20 Transferred entry. exoribonuclease. Now EC 3.1.13.1, exoribonuclease II]

[EC 3.1.4.20 created 1972, deleted 1978]

[3.1.4.21 Transferred entry. single-stranded-nucleate endonuclease. Now EC 3.1.30.1, Aspergillus nuclease  $S_1$ ]

[EC 3.1.4.21 created 1972, deleted 1978]

[3.1.4.22 Transferred entry. ribonuclease I. Now EC 3.1.27.5, pancreatic ribonuclease]

[EC 3.1.4.22 created 1972, deleted 1978]

[3.1.4.23 Transferred entry. ribonuclease II. Now EC 3.1.27.1, ribonuclease  $T_2$ ]

	[EC 3.1.4.23 created 1972, deleted 1978]			
[3.1.4.24	Deleted entry. endoribonuclease III]			
	[EC 3.1.4.24 created 1972, deleted 1978]			
[3.1.4.25	Transferred entry. exodeoxyribonuclease I. Now EC 3.1.11.1, exodeoxyribonuclease I]			
	[EC 3.1.4.25 created 1972, deleted 1978]			
[3.1.4.26	Deleted entry. exodeoxyribonuclease II]			
	[EC 3.1.4.26 created 1972, deleted 1978]			
[3.1.4.27	Transferred entry. exodeoxyribonuclease III. Now EC 3.1.11.2, exodeoxyribonuclease III]			
	[EC 3.1.4.27 created 1972, deleted 1978]			
[3.1.4.28	Transferred entry. exodeoxyribonuclease IV. Now EC 3.1.11.3, exodeoxyribonuclease (lambda-induced)]			
	[EC 3.1.4.28 created 1972, deleted 1978]			
[3.1.4.29	Deleted entry. oligodeoxyribonucleate exonuclease]			
	[EC 3.1.4.29 created 1972, deleted 1978]			
[3.1.4.30	Transferred entry. endodeoxyribonuclease. Now EC 3.1.21.2, deoxyribonuclease IV (phage- $T_4$ -induced)]			
	[EC 3.1.4.30 created 1972, deleted 1978]			
[3.1.4.31	Transferred entry. DNA 5'-dinucleotidohydrolase. Now EC 3.1.11.4, exodeoxyribonuclease (phage SP <sub>3</sub> -induced)]			

[3.1.4.32 Deleted entry. endodeoxyribonuclease (ATP- and S-adenosylmethionine-dependent). See EC 3.1.21.3 type 1 site-specific deoxyribonuclease and EC 3.1.21.5 type III site-specific deoxyribonuclease]

[EC 3.1.4.31 created 1972, deleted 1978]

[EC 3.1.4.32 created 1972, deleted 1978]

[3.1.4.33 Deleted entry. endodeoxyribonuclease (ATP-hydrolysing). See EC 3.1.21.3 type 1 site-specific deoxyribonuclease and EC 3.1.21.5 type III site-specific deoxyribonuclease]

[EC 3.1.4.33 created 1972, deleted 1978]

[3.1.4.34 Deleted entry. hybrid nuclease. See subclasses EC 3.1.15, EC 3.1.16, EC 3.1.30 and EC 3.1.31]

[EC 3.1.4.34 created 1972, deleted 1978]

## EC 3.1.4.35

**Accepted name:** 3',5'-cyclic-GMP phosphodiesterase

**Reaction:** guanosine 3',5'-cyclic phosphate +  $H_2O = GMP$ 

Other name(s): guanosine cyclic 3',5'-phosphate phosphodiesterase; cyclic GMP phosphodiesterase; cyclic 3',5'-GMP

phosphodiesterase; cyclic guanosine 3',5'-monophosphate phosphodiesterase; cyclic guanosine 3',5'-

phosphate phosphodiesterase; cGMP phosphodiesterase; cGMP-PDE

**Systematic name:** 3',5'-cyclic-GMP 5'-nucleotidohydrolase

**References:** [1815]

[EC 3.1.4.35 created 1976]

[3.1.4.36 Deleted entry. 1,2-cyclic-inositol-phosphate phosphodiesterase. Now included with EC 3.1.4.43, glycerophosphoinositol inositolphosphodiesterase]

[EC 3.1.4.36 created 1976, deleted 2002]

EC 3.1.4.37

**Accepted name:** 2',3'-cyclic-nucleotide 3'-phosphodiesterase

**Reaction:** nucleoside 2',3'-cyclic phosphate +  $H_2O$  = nucleoside 2'-phosphate

Other name(s): cyclic-CMP phosphodiesterase; 2',3'-cyclic AMP phosphodiesterase; cyclic 2',3'-nucleotide 3'-

phosphodiesterase; cyclic 2',3'-nucleotide phosphodiesterase; 2',3'-cyclic nucleoside monophosphate phosphodiesterase; 2',3'-cyclic nucleotide 3'-phosphohydrolase; CNPase; 2',3'-cyclic nucleotide phosphodiesterase; 2',3'-cyclic nucleotide p

phohydrolase; 2':3'-cyclic nucleotide 3'-phosphodiesterase; 2':3'-CNMP-3'-ase

**Systematic name:** nucleoside-2',3'-cyclic-phosphate 2'-nucleotidohydrolase

**Comments:** The brain enzyme acts on 2',3'-cyclic AMP more rapidly than on the UMP or CMP derivatives. An

enzyme from liver acts on 2', 3'-cyclic CMP more rapidly than on the purine derivatives; it also hydrolyses the corresponding 3', 5'-cyclic phosphates, but more slowly. This latter enzyme has been

called cyclic-CMP phosphodiesterase.

**References:** [654, 1105, 1106, 1568, 2098]

[EC 3.1.4.37 created 1976]

EC 3.1.4.38

**Accepted name:** glycerophosphocholine cholinephosphodiesterase

**Reaction:** sn-glycero-3-phosphocholine +  $H_2O$  = glycerol + phosphocholine

Other name(s): L-3-glycerylphosphinicocholine cholinephosphohydrolase Systematic name: sn-glycero-3-phosphocholine cholinephosphohydrolase

**Comments:** No activity on *sn*-3-glycerophosphoethanolamine.

**References:** [7]

[EC 3.1.4.38 created 1976]

EC 3.1.4.39

**Accepted name:** alkylglycerophosphoethanolamine phosphodiesterase

**Reaction:** 1-alkyl-sn-glycero-3-phosphoethanolamine +  $H_2O$  = 1-alkyl-sn-glycerol 3-phosphate + ethanolamine

**Other name(s):** lysophospholipase D

**Systematic name:** 1-alkyl-*sn*-glycero-3-phosphoethanolamine ethanolaminehydrolase

**Comments:** Also acts on acyl and choline analogues.

**References:** [3226]

[EC 3.1.4.39 created 1976]

EC 3.1.4.40

**Accepted name:** CMP-*N*-acylneuraminate phosphodiesterase

**Reaction:** CMP-N-acylneuraminate +  $H_2O = CMP + N$ -acylneuraminate

Other name(s): CMP-sialate hydrolase; CMP-sialic acid hydrolase; CMP-N-acylneuraminic acid hydrolase; cytidine

monophosphosialic hydrolase; cytidine monophosphosialate hydrolase; cytidine monophosphate-N-

acetylneuraminic acid hydrolase; CMP-N-acetylneuraminate hydrolase

**Systematic name:** CMP-*N*-acylneuraminate *N*-acylneuraminohydrolase

**References:** [1409]

[EC 3.1.4.40 created 1976]

EC 3.1.4.41

**Accepted name:** sphingomyelin phosphodiesterase D

**Reaction:** sphingomyelin +  $H_2O$  = ceramide phosphate + choline

Other name(s): sphingomyelinase D

**Systematic name:** sphingomyelin ceramide-phosphohydrolase

**Comments:** Does not act on phosphatidylcholine, but hydrolyses 2-lysophosphatidylcholine to choline and 2-

lysophosphatidate.

**References:** [375, 2731]

[EC 3.1.4.41 created 1978]

EC 3.1.4.42

**Accepted name:** glycerol-1,2-cyclic-phosphate 2-phosphodiesterase

**Reaction:** glycerol 1,2-cyclic phosphate + H<sub>2</sub>O = glycerol 1-phosphate **Other name(s):** rac-glycerol 1:2-cyclic phosphate 2-phosphodiesterase **Systematic name:** rac-glycerol-1,2-cyclic-phosphate 2-glycerophosphohydrolase

**Comments:** Acts on both stereoisomers of the substrate and also, more slowly, on 3',5'-cyclic AMP and on 2',3'-

cyclic AMP.

**References:** [459]

[EC 3.1.4.42 created 1984]

EC 3.1.4.43

**Accepted name:** glycerophosphoinositol inositolphosphodiesterase

**Reaction:** 1-(*sn*-glycero-3-phospho)-1D-*myo*-inositol + H<sub>2</sub>O = glycerol + 1D-*myo*-inositol 1-phosphate **Other name(s):** 1,2-cyclic-inositol-phosphate phosphodiesterase; D-*myo*-inositol 1:2-cyclic phosphate 2-

phosphohydrolase; D-inositol 1,2-cyclic phosphate 2-phosphohydrolase; D-*myo*-inositol 1,2-cyclic phosphate 2-phosphohydrolase; 1-D-*myo*-inositol-1,2-cyclic-phosphate 2-inositolphosphohydrolase;

inositol-1,2-cyclic-phosphate 2-inositolphosphohydrolase

**Systematic name:** 1-(*sn*-glycero-3-phospho)-1D-*myo*-inositol inositolphosphohydrolase

**Comments:** This enzyme also hydrolyses Ins(cyclic1,2)*P* to Ins-1-*P* 

**References:** [554, 552, 553, 2463]

[EC 3.1.4.43 created 1984, (EC 3.1.4.36 created 1976, incorporated 2002), modified 2002]

EC 3.1.4.44

Accepted name: glycerophosphoinositol glycerophosphodiesterase

**Reaction:** 1-(sn-glycero-3-phospho)-1D-myo-inositol +  $H_2O = myo$ -inositol + sn-glycerol 3-phosphate **Other name(s):** sn-glycero(3)phosphoinositol glycerophosphohydrolase; sn-glycero-3-phospho-1-inositol glycerophosphohydrolase; sn-glycero-3-phosphohydrolase; sn-glycero-3-phosphohydrolas

erophosphohydrolase

**Systematic name:** 1-(sn-glycero-3-phospho)-1D-myo-inositol glycerophosphohydrolase

**References:** [555]

[EC 3.1.4.44 created 1984, modified 2002]

EC 3.1.4.45

**Accepted name:** *N*-acetylglucosamine-1-phosphodiester α-*N*-acetylglucosaminidase

**Reaction:** glycoprotein N-acetyl-D-glucosaminyl-phospho-D-mannose +  $H_2O = N$ -acetyl-D-glucosamine + gly-

coprotein phospho-D-mannose

Other name(s):  $\alpha$ -N-acetylglucosaminyl phosphodiesterase; lysosomal  $\alpha$ -N-acetylglucosaminidase; phosphodiester

glycosidase;  $\alpha$ -N-acetyl-D-glucosamine-1-phosphodiester N-acetylglucosaminidase; 2-acetamido-2-

deoxy-α-D-glucose 1-phosphodiester acetamidodeoxyglucohydrolase

**Systematic name:** glycoprotein-*N*-acetyl-D-glucosaminyl-phospho-D-mannose *N*-acetyl-D-

glucosaminylphosphohydrolase

**Comments:** Acts on a variety of compounds in which N-acetyl-D-glucosamine is  $\alpha$ -linked to a phosphate group,

including the biosynthetic intermediates of the high mannose oligosaccharide components of some lysosomal enzymes and the products of EC 2.7.8.17 UDP-*N*-acetylglucosamine—lysosomal-enzyme

*N*-acetylglucosaminephosphotransferase.

**References:** [581, 3037, 3039, 3083]

### [EC 3.1.4.45 created 1984]

EC 3.1.4.46

Accepted name: glycerophosphodiester phosphodiesterase

**Reaction:** a glycerophosphodiester +  $H_2O$  = an alcohol + sn-glycerol 3-phosphate

Other name(s): gene hpd protein; glycerophosphoryl diester phosphodiesterase; IgD-binding protein D

**Systematic name:** glycerophosphodiester glycerophosphohydrolase

**Comments:** Broad specificity for glycerophosphodiesters; glycerophosphocholine, glycerophosphoethanolamine,

glycerophosphoglycerol and bis(glycerophospho)-glycerol are hydrolysed.

**References:** [1606]

[EC 3.1.4.46 created 1986]

[3.1.4.47 Transferred entry. variant-surface-glycoprotein phospholipase C. Now EC 4.6.1.14, glycosylphosphatidylinositol diacylglycerol-lyase]

[EC 3.1.4.47 created 1989, deleted 2002]

EC 3.1.4.48

Accepted name: dolichylphosphate-glucose phosphodiesterase

**Reaction:** dolichyl  $\beta$ -D-glucosyl phosphate + H<sub>2</sub>O = dolichyl phosphate + D-glucose **Other name(s):** dolichol phosphoglucose phosphodiesterase; Dol-*P*-Glc phosphodiesterase

Systematic name: dolichyl-β-D-glucosyl-phosphate dolichylphosphohydrolase

**References:** [502]

[EC 3.1.4.48 created 1989]

EC 3.1.4.49

Accepted name: dolichylphosphate-mannose phosphodiesterase

**Reaction:** dolichyl  $\beta$ -D-mannosyl phosphate + H<sub>2</sub>O = dolichyl phosphate + D-mannose

**Other name(s):** mannosylphosphodolichol phosphodiesterase

**Systematic name:** dolichyl-β-D-mannosyl-phosphate dolichylphosphohydrolase

**References:** [2936]

[EC 3.1.4.49 created 1990]

EC 3.1.4.50

**Accepted name:** glycosylphosphatidylinositol phospholipase D

**Reaction:**  $6-(\alpha-D-glucosaminyl)-1-phosphatidyl-1D-myo-inositol + H<sub>2</sub>O = <math>6-(\alpha-D-glucosaminyl)-1D-myo-inositol$ 

inositol + 3-sn-phosphatidate

Other name(s): GPI-PLD; glycoprotein phospholipase D; phosphatidylinositol phospholipase D;

phosphatidylinositol-specific phospholipase D

Systematic name: glycoprotein-phosphatidylinositol phosphatidohydrolase

**Comments:** This enzyme is also active when *O*-4 of the glucosamine is substituted by carrying the oligosaccharide

that can link a protein to the structure. It therefore cleaves proteins from the lipid part of the glycosylphosphatidylinositol (GPI) anchors, but does so by hydrolysis, whereas glycosylphosphatidylinositol diacylglycerol-lyase (EC 4.6.1.14) does so by elimination. It acts on plasma membranes only after solubilization of the substrate with detergents or solvents, but it may act on intracellular membranes.

**References:** [1742, 1791, 1665, 567]

[EC 3.1.4.50 created 1990, modified 2002]

EC 3.1.4.51

Accepted name: glucose-1-phospho-D-mannosylglycoprotein phosphodiesterase

**Reaction:** 6-(D-glucose-1-phospho)-D-mannosylglycoprotein +  $H_2O = \alpha$ -D-glucose 1-phosphate + D-

mannosylglycoprotein

Other name(s):  $\alpha$ -glucose-1-phosphate phosphodiesterase

**Systematic name:** 6-(D-glucose-1-phospho)-D-mannosylglycoprotein glucose-1-phosphohydrolase

**Comments:** The enzyme is specific for the product of EC 2.7.8.19 UDP-glucose—glycoprotein glucose phospho-

transferase.

**References:** [2748]

[EC 3.1.4.51 created 1992]

EC 3.1.4.52

Accepted name: cyclic-guanylate-specific phosphodiesterase

**Reaction:** cyclic di-3',5'-guanylate +  $H_2O = 5'$ -phosphoguanylyl(3' $\rightarrow$ 5')guanosine

Other name(s): cyclic bis $(3' \rightarrow 5')$  diguanylate phosphodiesterase; c-di-GMP-specific phosphodiesterase; c-di-GMP

phosphodiesterase; phosphodiesterase (misleading); phosphodiesterase A1; PDEA1; VieA

**Systematic name:** cyclic bis $(3' \rightarrow 5')$ diguanylate 3'-guanylylhydrolase

**Comments:** Requires  $Mg^{2+}$  or  $Mn^{2+}$  for activity and is inhibited by  $Ca^{2+}$  and  $Zn^{2+}$ . Contains a heme unit. This

enzyme linearizes cyclic di-3',5'-guanylate, the product of EC 2.7.7.65, diguanylate cyclase and an allosteric activator of EC 2.4.1.12, cellulose synthase (UDP-forming), rendering it inactive [392]. It is

the balance between these two enzymes that determines the cellular level of c-di-GMP [392].

**References:** [392, 442, 2570, 2866]

[EC 3.1.4.52 created 2008]

EC 3.1.4.53

**Accepted name:** 3',5'-cyclic-AMP phosphodiesterase

**Reaction:** adenosine 3',5'-cyclic phosphate +  $H_2O = AMP$ 

**Other name(s):** cAMP-specific phosphodiesterase; cAMP-specific PDE; PDE1; PDE2A; PDE2B; PDE4; PDE7;

PDE8: PDEB1: PDEB2

**Systematic name:** 3',5'-cyclic-AMP 5'-nucleotidohydrolase

**Comments:** Requires  $Mg^{2+}$  or  $Mn^{2+}$  for activity [117]. This enzyme is specific for 3',5'-cAMP and does not hy-

drolyse other nucleoside 3',5'-cyclic phosphates such as cGMP (*cf.* EC 3.1.4.17, 3,5-cyclic-nucleotide phosphodiesterase and EC 3.1.4.35, 3,5-cyclic-GMP phosphodiesterase). It is involved in modulation of the levels of cAMP, which is a mediator in the processes of cell transformation and proliferation

[2376].

**References:** [39, 117, 2376, 1324, 1751, 1240]

[EC 3.1.4.53 created 2008, modified 2011]

EC 3.1.4.54

**Accepted name:** N-acetylphosphatidylethanolamine-hydrolysing phospholipase D

**Reaction:** N-acylphosphatidylethanolamine +  $H_2O = N$ -acylethanolamine + a 1,2-diacylglycerol 3-phosphate **Other name(s):** NAPE-PLD; anandamide-generating phospholipase D; N-acyl phosphatidylethanolamine phospholi-

pase D; NAPE-hydrolyzing phospholipase D

**Systematic name:** *N*-acetylphosphatidylethanolamine phosphatidohydrolase

Comments: This enzyme is involved in the biosynthesis of anandamide. It does not hydrolyse phosphatidylcholine

and phosphatidylethanolamine [2184]. No transphosphatidation [2184]. The enzyme contains Zn<sup>2+</sup>

and is activated by  $Mg^{2+}$  or  $Ca^{2+}$  [3111].

**References:** [2184, 3111]

[EC 3.1.4.54 created 2011]

## EC 3.1.4.55

**Accepted name:** phosphoribosyl 1,2-cyclic phosphate phosphodiesterase

**Reaction:** 5-phospho- $\alpha$ -D-ribose 1,2-cyclic phosphate + H<sub>2</sub>O =  $\alpha$ -D-ribose 1,5-bisphosphate

**Other name(s):** *phnP* (gene name)

Systematic name: 5-phospho-α-D-ribose 1,2-cyclic phosphate 2-phosphohydrolase (α-D-ribose 1,5-bisphosphate-

forming)

**Comments:** Binds  $Mn^{2+}$  and  $Zn^{2+}$ . Isolated from the bacterium *Escherichia coli*, where it participates in the

degradation of methylphosphonate.

**References:** [2296, 1190, 1091]

[EC 3.1.4.55 created 2013]

## EC 3.1.4.56

**Accepted name:** 7,8-dihydroneopterin 2′,3′-cyclic phosphate phosphodiesterase

**Reaction:** (1) 7,8-dihydroneopterin 2',3'-cyclic phosphate + H<sub>2</sub>O = 7,8-dihydroneopterin 3'-phosphate

(2) 7,8-dihydroneopterin 2',3'-cyclic phosphate +  $H_2O = 7$ ,8-dihydroneopterin 2'-phosphate

**Other name**(s): MptB

**Systematic name:** 7,8-dihydroneopterin 2',3'-cyclic phosphate 2'/3'-phosphodiesterase

**Comments:** Contains one zinc atom and one iron atom per subunit of the dodecameric enzyme. It hydrolyses 7,8-

dihydroneopterin 2',3'-cyclic phosphate, a step in tetrahydromethanopterin biosynthesis. *In vitro* the enzyme forms 7,8-dihydroneopterin 2'-phosphate and 7,8-dihydroneopterin 3'-phosphate at a ratio of

4:1.

References: [1832]

[EC 3.1.4.56 created 2013]

#### EC 3.1.4.57

**Accepted name:** phosphoribosyl 1,2-cyclic phosphate 1,2-diphosphodiesterase

**Reaction:** (1) 5-phospho- $\alpha$ -D-ribose 1,2-cyclic phosphate + H<sub>2</sub>O = D-ribofuranose 2,5-bisphosphate

(2) D-ribofuranose 2,5-bisphosphate +  $H_2O$  = D-ribofuranose 5-phosphate + phosphate

**Other name(s):** cyclic phosphate dihydrolase; *phnPP* (gene name)

**Systematic name:** 5-phospho-α-D-ribose 1,2-cyclic phosphate 1,2-diphosphophosphohydrolase

**Comments:** The enzyme, characterized from the bacterium *Eggerthella lenta*, is involed in degradation of

methylphosphonate.

**References:** [904]

[EC 3.1.4.57 created 2014]

#### EC 3.1.4.58

**Accepted name:** RNA 2',3'-cyclic 3'-phosphodiesterase

**Reaction:** (ribonucleotide) $_{n}$ -2',3'-cyclic phosphate + H<sub>2</sub>O = (ribonucleotide) $_{n}$ -2'-phosphate

Other name(s): thpR (gene name); ligT (gene name)

**Systematic name:** (ribonucleotide)n-2', 3'-cyclic phosphate 3'-nucleotidohydrolase

**Comments:** The enzyme hydrolyses RNA 2',3'-cyclic phosphodiester to an RNA 2'-phosphomonoester. *In vitro* 

the enzyme can also ligate tRNA molecules with 2',3'-cyclic phosphate to tRNA with 5'-hydroxyl termini, forming a 2'-5' phosphodiester linkage. However, the ligase activity is unlikely to be relevant

in vivo.

**References:** [1376, 2407]

[EC 3.1.4.58 created 2017]

# EC 3.1.5 Triphosphoric-monoester hydrolases

### EC 3.1.5.1

Accepted name: dGTPase

**Reaction:**  $dGTP + H_2O = deoxyguanosine + triphosphate$ 

Other name(s): deoxy-GTPase; deoxyguanosine 5-triphosphate triphosphohydrolase; deoxyguanosine triphosphatase;

deoxyguanosine triphosphate triphosphohydrolase

**Systematic name:** dGTP triphosphohydrolase

**Comments:** Also acts on GTP.

**References:** [1514]

[EC 3.1.5.1 created 1961]

# EC 3.1.6 Sulfuric-ester hydrolases

## EC 3.1.6.1

Accepted name: arylsulfatase

**Reaction:** a phenol sulfate  $+ H_2O = a$  phenol + sulfate

Other name(s): sulfatase; nitrocatechol sulfatase; phenolsulfatase; phenylsulfatase; p-nitrophenyl sulfatase; arylsulfo-

hydrolase; 4-methylumbelliferyl sulfatase; estrogen sulfatase

**Systematic name:** aryl-sulfate sulfohydrolase

**Comments:** A group of enzymes with rather similar specificities.

**References:** [625, 2471, 2472, 3141]

[EC 3.1.6.1 created 1961, modified 2011]

### EC 3.1.6.2

Accepted name: steryl-sulfatase

**Reaction:**  $3\beta$ -hydroxyandrost-5-en-17-one 3-sulfate +  $H_2O = 3\beta$ -hydroxyandrost-5-en-17-one + sulfate

Other name(s): arylsulfatase; steroid sulfatase; steroid sulfatase; dehydroepiandrosterone sulfate sulfatase; arylsulfatase

C; steroid 3-sulfatase; steroid sulfate sulfohydrolase; dehydroepiandrosterone sulfatase; pregnenolone

sulfatase; phenolic steroid sulfatase;  $3-\beta$ -hydroxysteroid sulfate sulfatase

**Systematic name:** steryl-sulfate sulfohydrolase

**Comments:** Also acts on some related steryl sulfates.

**References:** [2470, 2471, 2772]

[EC 3.1.6.2 created 1961]

## EC 3.1.6.3

Accepted name: glycosulfatase

**Reaction:** D-glucose 6-sulfate +  $H_2O$  = D-glucose + sulfate

Other name(s): glucosulfatase

Systematic name: sugar-sulfate sulfohydrolase

**Comments:** Also acts on other sulfates of monosaccharides and disaccharides and on adenosine 5'-sulfate.

**References:** [624, 679, 2471]

[EC 3.1.6.3 created 1961]

### EC 3.1.6.4

**Accepted name:** *N*-acetylgalactosamine-6-sulfatase

Reaction: Hydrolysis of the 6-sulfate groups of the N-acetyl-D-galactosamine 6-sulfate units of chondroitin sul-

fate and of the D-galactose 6-sulfate units of keratan sulfate

Other name(s): chondroitin sulfatase; chondroitinase; galactose-6-sulfate sulfatase; acetylgalactosamine 6-sulfatase;

N-acetylgalactosamine-6-sulfate sulfatase; N-acetylgalactosamine 6-sulfatase

**Systematic name:** *N*-acetyl-D-galactosamine-6-sulfate 6-sulfohydrolase

**References:** [709, 931, 1687, 2726, 3316]

[EC 3.1.6.4 created 1961]

[3.1.6.5 Deleted entry. sinigrin sulfohydrolase; myrosulfatase]

[EC 3.1.6.5 created 1961, deleted 1964]

EC 3.1.6.6

Accepted name: choline-sulfatase

**Reaction:** choline sulfate  $+ H_2O =$ choline +sulfate

**Systematic name:** choline-sulfate sulfohydrolase

**References:** [2854]

[EC 3.1.6.6 created 1965]

EC 3.1.6.7

Accepted name: cellulose-polysulfatase

**Reaction:** Hydrolysis of the 2- and 3-sulfate groups of the polysulfates of cellulose and charonin

Systematic name: cellulose-sulfate sulfohydrolase

**References:** [2842]

[EC 3.1.6.7 created 1965]

EC 3.1.6.8

**Accepted name:** cerebroside-sulfatase

**Reaction:** a cerebroside 3-sulfate  $+ H_2O = a$  cerebroside + sulfate

Other name(s): arylsulfatase A; cerebroside sulfatase Systematic name: cerebroside-3-sulfate 3-sulfohydrolase

Comments: Hydrolyses galactose-3-sulfate residues in a number of lipids. Also hydrolyses ascorbate 2-sulfate and

many phenol sulfates.

**References:** [1877, 2472]

[EC 3.1.6.8 created 1972]

EC 3.1.6.9

Accepted name: chondro-4-sulfatase

**Reaction:** 4-deoxy- $\beta$ -D-gluc-4-enuronosyl- $(1\rightarrow 3)$ -N-acetyl-D-galactosamine 4-sulfate + H<sub>2</sub>O = 4-deoxy- $\beta$ -D-

gluc-4-enuronosyl- $(1\rightarrow 3)$ -N-acetyl-D-galactosamine + sulfate

Other name(s): chondroitin-4-sulfatase; 4-deoxy-β-D-gluc-4-enuronosyl-(1,3)-N-acetyl-D-galactosamine-4-sulfate

4-sulfohydrolase

**Systematic name:** 4-deoxy- $\beta$ -D-gluc-4-enuronosyl- $(1\rightarrow 3)$ -N-acetyl-D-galactosamine-4-sulfate 4-sulfohydrolase

**Comments:** Also acts on the saturated analogue but not on higher oligosaccharides, nor any 6-sulfates.

**References:** [1104, 2472, 3244]

[EC 3.1.6.9 created 1972]

EC 3.1.6.10

**Accepted name:** chondro-6-sulfatase

**Reaction:** 4-deoxy- $\beta$ -D-gluc-4-enuronosyl- $(1\rightarrow 3)$ -N-acetyl-D-galactosamine 6-sulfate + H<sub>2</sub>O = 4-deoxy- $\beta$ -D-

gluc-4-enuronosyl- $(1\rightarrow 3)$ -N-acetyl-D-galactosamine + sulfate

Other name(s): 4-deoxy-β-D-gluc-4-enuronosyl-(1,3)-*N*-acetyl-D-galactosamine-6-sulfate 6-sulfohydrolase

**Systematic name:** 4-deoxy- $\beta$ -D-gluc-4-enuronosyl- $(1\rightarrow 3)$ -N-acetyl-D-galactosamine-6-sulfate 6-sulfohydrolase

**Comments:** Also acts on the saturated analogue and *N*-acetyl-D-galactosamine 4,6-disulfate, but not higher

oligosaccharides, nor any 4-sulfate

**References:** [3244]

[EC 3.1.6.10 created 1972]

EC 3.1.6.11

Accepted name: disulfoglucosamine-6-sulfatase

**Reaction:** 2-N,6-O-disulfo-D-glucosamine +  $H_2O$  = 2-N-sulfo-D-glucosamine + sulfate

**Other name(s):** N-sulfoglucosamine-6-sulfatase; 6,N-disulfoglucosamine 6-O-sulfohydrolase; N,6-O-disulfo-D-

glucosamine 6-sulfohydrolase

**Systematic name:** 2-N,6-*O*-disulfo-D-glucosamine 6-sulfohydrolase

**Comments:** May be identical with EC 3.1.6.14 *N*-acetylglucosamine-6-sulfatase.

**References:** [609]

[EC 3.1.6.11 created 1972, modified 1989]

EC 3.1.6.12

**Accepted name:** *N*-acetylgalactosamine-4-sulfatase

**Reaction:** Hydrolysis of the 4-sulfate groups of the *N*-acetyl-D-galactosamine 4-sulfate units of chondroitin sul-

fate and dermatan sulfate

Other name(s): chondroitinsulfatase; chondroitinase; arylsulfatase B; acetylgalactosamine 4-sulfatase; N-

acetylgalactosamine 4-sulfate sulfohydrolase

**Systematic name:** *N*-acetyl-D-galactosamine-4-sulfate 4-sulfohydrolase

**Comments:** Acts also on *N*-acetylglucosamine 4-sulfate.

**References:** [739, 959, 2979]

[EC 3.1.6.12 created 1984]

EC 3.1.6.13

**Accepted name:** iduronate-2-sulfatase

Reaction: Hydrolysis of the 2-sulfate groups of the L-iduronate 2-sulfate units of dermatan sulfate, heparan sul-

fate and heparin

Other name(s): chondroitinsulfatase; idurono-2-sulfatase; iduronide-2-sulfate sulfatase; L-iduronosulfatase; L-idurono

sulfatase; iduronate sulfatase; sulfo-L-iduronate sulfatase; L-iduronate 2-sulfate sulfatase; sulfoiduronate sulfohydrolase; 2-sulfo-L-iduronate 2-sulfatase; iduronate 2-sulfatase; iduronate

sulfate sulfatase

**Systematic name:** L-iduronate-2-sulfate 2-sulfohydrolase

**References:** [71, 113, 616, 3315]

[EC 3.1.6.13 created 1984]

EC 3.1.6.14

**Accepted name:** *N*-acetylglucosamine-6-sulfatase

**Reaction:** Hydrolysis of the 6-sulfate groups of the *N*-acetyl-D-glucosamine 6-sulfate units of heparan sulfate

and keratan sulfate

**Other name(s):** chondroitinsulfatase; O,N-disulfate O-sulfohydrolase; acetylglucosamine 6-sulfatase; N-

acetylglucosamine 6-sulfate sulfatase; acetylglucosamine 6-sulfatase; 2-acetamido-2-deoxy-D-glucose

6-sulfate sulfatase

**Systematic name:** *N*-acetyl-D-glucosamine-6-sulfate 6-sulfohydrolase

**Comments:** May be identical with EC 3.1.6.11 disulfoglucosamine-6-sulfatase.

**References:** [169, 1534, 3156]

## [EC 3.1.6.14 created 1984]

EC 3.1.6.15

**Accepted name:** *N*-sulfoglucosamine-3-sulfatase

**Reaction:** Hydrolysis of the 3-sulfate groups of the *N*-sulfo-D-glucosamine 3-*O*-sulfate units of heparin

Other name(s): chondroitinsulfatase

**Systematic name:** *N*-sulfo-3-sulfoglucosamine 3-sulfohydrolase

**Comments:** The enzyme from *Flavobacterium heparinum* also hydrolyses *N*-acetyl-D-glucosamine 3-*O*-sulfate;

the mammalian enzyme acts only on the disulfated residue.

**References:** [315, 1616]

[EC 3.1.6.15 created 1984, modified 1989]

EC 3.1.6.16

**Accepted name:** monomethyl-sulfatase

**Reaction:** monomethyl sulfate  $+ H_2O = methanol + sulfate$ 

**Systematic name:** monomethyl-sulfate sulfohydrolase

Comments: Highly specific; does not act on monoethyl sulfate, monoisopropyl sulfate or monododecyl sulfate.

**References:** [903]

[EC 3.1.6.16 created 1989]

EC 3.1.6.17

Accepted name: D-lactate-2-sulfatase

**Reaction:** (*R*)-2-*O*-sulfolactate +  $H_2O = (R)$ -lactate + sulfate

**Other name(s):** (S)-2-O-sulfolactate 2-sulfohydrolase (incorrect stereochemistry)

**Systematic name:** (R)-2-O-sulfolactate 2-sulfohydrolase

**Comments:** Highly specific.

**References:** [504]

[EC 3.1.6.17 created 1989]

EC 3.1.6.18

**Accepted name:** glucuronate-2-sulfatase

**Reaction:** Hydrolysis of the 2-sulfate groups of the 2-O-sulfo-D-glucuronate residues of chondroitin sulfate,

heparin and heparitin sulfate

**Other name(s):** glucurono-2-sulfatase

**Systematic name:** polysaccharide-2-*O*-sulfo-D-glucuronate 2-sulfohydrolase

**Comments:** Does not act on iduronate 2-sulfate residues (*cf.* EC 3.1.6.13 iduronate-2-sulfatase)

**References:** [2617]

[EC 3.1.6.18 created 1989]

EC 3.1.6.19

**Accepted name:** (*R*)-specific secondary-alkylsulfatase

**Reaction:** an (R)-secondary-alkyl sulfate +  $H_2O$  = an (S)-secondary-alcohol + sulfate

Other name(s): S3 secondary alkylsulphohydrolase; Pisa1; secondary alkylsulphohydrolase; (R)-specific sec-

alkylsulfatase; sec-alkylsulfatase

**Systematic name:** (R)-secondary-alkyl sulfate sulfohydrolase [(S)-secondary-alcohol forming]

**Comments:** The enzyme from *Rhodococcus ruber* is involved in the biodegradation of alkyl sulfate esters used as

detergents and released into the environment. The prefered substrates are linear secondary-alkyl sulfate esters, particularly octan-2-yl, octan-3-yl, and octan-4-yl sulfates [2299]. The enzyme from *Pseudomonas* sp. DSM6611 utilizes a range of secondary-alkyl sulfate esters bearing aromatic, olefinic and acetylenic moieties. Perfect enantioselectivities are obtained for substrates bearing groups of different size adjacent to the sulfate moiety [2579]. The enzymic hydrolysis proceeds through inversion of the configuration at the stereogenic carbon atom [2299, 2579]. The enzyme contains a Zn<sup>2+</sup> ion

[1481].

**References:** [2299, 3100, 1481, 2579]

[EC 3.1.6.19 created 2013]

EC 3.1.6.20

**Accepted name:** S-sulfosulfanyl-L-cysteine sulfohydrolase

**Reaction:** (1) [SoxY protein]-S-sulfosulfanyl-L-cysteine +  $H_2O$  = [SoxY protein]-S-sulfanyl-L-cysteine + sulfate

(2) [SoxY protein]-S-(2-sulfodisulfanyl)-L-cysteine +  $H_2O = [SoxY protein]$ -S-disulfanyl-L-cysteine +

sulfate

Other name(s): SoxB

**Systematic name:** [Sox Y protein]-S-sulfosulfanyl-L-cysteine sulfohydrolase

**Comments:** Contains  $Mn^{2+}$ . The enzyme is part of the Sox enzyme system, which participates in a bacterial thio-

sulfate oxidation pathway that produces sulfate. It catalyses two reactions in the pathway. In both cases the enzyme hydrolyses a sulfonate moiety that is bound (either directly or via a sulfane) to a cysteine residue of a SoxY protein, releasing sulfate. The enzyme from *Paracoccus pantotrophus* con-

tains a pyroglutamate (cycloglutamate) at its N-terminus.

**References:** [2343, 822, 2344, 707, 1118, 961]

[EC 3.1.6.20 created 2018]

# EC 3.1.7 Diphosphoric-monoester hydrolases

EC 3.1.7.1

**Accepted name:** prenyl-diphosphatase

**Reaction:** prenyl diphosphate +  $H_2O$  = prenol + diphosphate

**Other name(s):** prenyl-pyrophosphatase; prenol pyrophosphatase; prenylphosphatase

Systematic name: prenyl-diphosphate diphosphohydrolase

**Comments:** Farnesyl diphosphate is the best substrate tested to date.

**References:** [2969]

[EC 3.1.7.1 created 1972]

EC 3.1.7.2

**Accepted name:** guanosine-3',5'-bis(diphosphate) 3'-diphosphatase

**Reaction:** guanosine 3',5'-bis(diphosphate) +  $H_2O = GDP$  + diphosphate

Other name(s): guanosine-3',5'-bis(diphosphate) 3'-pyrophosphatase; PpGpp-3'-pyrophosphohydrolase; PpGpp phos-

phohydrolase

**Systematic name:** guanosine-3',5'-bis(diphosphate) 3'-diphosphohydrolase

**References:** [1102, 2422]

[EC 3.1.7.2 created 1980]

EC 3.1.7.3

Accepted name: monoterpenyl-diphosphatase

**Reaction:** a monoterpenyl diphosphate  $+ H_2O = a$  monoterpenyl + diphosphate **Other name(s):** bornyl pyrophosphate hydrolase; monoterpenyl-pyrophosphatase

**Systematic name:** monoterpenyl-diphosphate diphosphohydrolase

Comments: A group of enzymes with varying specificity for the monoterpenol moiety. One has the highest activ-

ity on sterically hindered compounds such as (+)-bornyl diphosphate; another has highest activity on

the diphosphates of primary allylic alcohols such as geraniol.

**References:** [510]

[EC 3.1.7.3 created 1984]

[3.1.7.4 Deleted entry. Now recognized as two enzymes EC 4.2.1.133, copal-8-ol diphosphate synthase and EC 4.2.3.141 sclareol synthase]

[EC 3.1.7.4 created 2008, deleted 2013]

EC 3.1.7.5

**Accepted name:** geranylgeranyl diphosphate diphosphatase

**Reaction:** geranylgeranyl diphosphate  $+ H_2O = geranylgeraniol + diphosphate$ 

Other name(s): geranylgeranyl diphosphate phosphatase Systematic name: geranyl-diphosphate diphosphohydrolase

**Comments:** Involved in the biosynthesis of plaunotol. There are two isoenzymes with different ion requirements.

Neither require Mg<sup>2+</sup> but in addition PII is inhibited by Zn<sup>2+</sup>, Mn<sup>2+</sup> and Co<sup>2+</sup>. It is not known which

isoenzyme is involved in plaunotol biosynthesis.

**References:** [2119]

[EC 3.1.7.5 created 2009]

EC 3.1.7.6

**Accepted name:** farnesyl diphosphatase

**Reaction:** (2E,6E)-farnesyl diphosphate +  $H_2O = (2E,6E)$ -farnesol + diphosphate

**Other name(s):** FPP phosphatase

**Systematic name:** (2E,6E)-farnesyl-diphosphate diphosphohydrolase

**Comments:** The enzyme is involved in the biosynthesis of acyclic sesquiterpenoids [2717].

**References:** [2717, 2969]

[EC 3.1.7.6 created 2010]

[3.1.7.7 Transferred entry. (-)-drimenol synthase. Now EC 4.2.3.194, (-)-drimenol synthase]

[EC 3.1.7.7 created 2011, deleted 2017]

EC 3.1.7.8

**Accepted name:** tuberculosinol synthase

**Reaction:** tuberculosinyl diphosphate +  $H_2O$  = tuberculosinol + diphosphate

Other name(s): Rv3378c

Systematic name: tuberculosinyl diphosphate diphosphohydrolase (tuberculosinol forming)

**Comments:** Only found in species of *Mycobacterium* that cause tuberculosis. In addition, it also gives isotubercu-

losinol in 1:1 mixture, cf. EC 3.1.7.9, isotuberculosinol synthase.

**References:** [2049, 1184]

[EC 3.1.7.8 created 2011]

EC 3.1.7.9

Accepted name: isotuberculosinol synthase

**Reaction:** tuberculosinyl diphosphate +  $H_2O = (13S)$ -isotuberculosinol + diphosphate

Other name(s): Rv3378c

**Systematic name:** tuberculosinyl diphosphate diphosphohydrolase (isotuberculosinol forming)

Comments: Only found in species of *Mycobacterium* that cause tuberculosis. In addition, it also gives tubercu-

losinol in 1:1 mixture, cf. EC 3.1.7.8, tuberculosinol synthase. The isotuberculosinol form was a 3:1

mixture of the 13S and 13R forms, respectively.

**References:** [2049, 1184]

[EC 3.1.7.9 created 2011]

EC 3.1.7.10

**Accepted name:** (13*E*)-labda-7,13-dien-15-ol synthase

**Reaction:** geranylgeranyl diphosphate +  $H_2O = (13E)$ -labda-7,13-dien-15-ol + diphosphate

**Other name(s):** labda-7,13*E*-dien-15-ol synthase

**Systematic name:** geranylgeranyl-diphosphate diphosphohydrolase [(13E)-labda-7,13-dien-15-ol-forming]

**Comments:** The enzyme from the lycophyte Selaginella moellendorffii is bifunctional, initially forming (13E)-

labda-7,13-dien-15-yl diphosphate, which is hydrolysed to the alcohol.

**References:** [1770]

[EC 3.1.7.10 created 2012]

EC 3.1.7.11

**Accepted name:** geranyl diphosphate diphosphatase

**Reaction:** geranyl diphosphate +  $H_2O$  = geraniol + diphosphate

**Other name(s):** geraniol synthase; geranyl pyrophosphate pyrophosphatase; GES; CtGES

Systematic name: geranyl-diphosphate diphosphohydrolase

Comments: Isolated from Ocimum basilicum (basil) and Cinnamomum tenuipile (camphor tree). Requires Mg<sup>2+</sup>

or  $Mn^{2+}$ . Geraniol is labelled when formed in the presence of [ $^{18}O$ ] $H_2O$ . Thus mechanism involves a geranyl cation [ $^{1225}$ ]. Neryl diphosphate is hydrolysed more slowly. May be the same as EC 3.1.7.3

monoterpenyl-diphosphatase.

**References:** [1225, 3260]

[EC 3.1.7.11 created 2012]

EC 3.1.7.12

**Accepted name:** (+)-kolavelool synthase

**Reaction:** (+)-kolavenyl diphosphate +  $H_2O = (+)$ -kolavelool + diphosphate

Other name(s): Haur\_2146

**Systematic name:** kolavenyl-diphosphate diphosphohydrolase

**Comments:** Isolated from the bacterium *Herpetosiphon aurantiacus*.

**References:** [2050]

[EC 3.1.7.12 created 2017]

## EC 3.1.8 Phosphoric-triester hydrolases

EC 3.1.8.1

Accepted name: aryldialkylphosphatase

**Reaction:** an aryl dialkyl phosphate  $+ H_2O = dialkyl$  phosphate + an aryl alcohol

Other name(s): organophosphate hydrolase; paraoxonase; A-esterase; aryltriphosphatase; organophosphate es-

terase; esterase B1; esterase E4; paraoxon esterase; pirimiphos-methyloxon esterase; OPA anhydrase; organophosphorus hydrolase; phosphotriesterase; paraoxon hydrolase; OPH; organophosphorus acid

anhydrase

**Systematic name:** aryltriphosphate dialkylphosphohydrolase

**Comments:** Acts on organophosphorus compounds (such as paraoxon) including esters of phosphonic and phosphorus

phinic acids. Inhibited by chelating agents; requires divalent cations for activity. Previously regarded

as identical with EC 3.1.1.2 arylesterase.

**References:** [30, 267, 1768, 1777, 1]

[EC 3.1.8.1 created 1989]

EC 3.1.8.2

Accepted name: diisopropyl-fluorophosphatase

**Reaction:** diisopropyl fluorophosphate +  $H_2O$  = diisopropyl phosphate + fluoride

Other name(s): DFPase; tabunase; organophosphorus acid anhydrolase; organophosphate acid anhydrase;

OPA anhydrase; diisopropylphosphofluoridase; dialkylfluorophosphatase; diisopropyl phosphorofluo-

ridate hydrolase; isopropylphosphorofluoridase; diisopropylfluorophosphonate dehalogenase

**Systematic name:** diisopropyl-fluorophosphate fluorohydrolase

Comments: Acts on phosphorus anhydride bonds (such as phosphorus-halide and phosphorus-cyanide) in

organophosphorus compounds (including 'nerve gases'). Inhibited by chelating agents; requires di-

valent cations. Related to EC 3.1.8.1 aryldialkylphosphatase.

**References:** [99, 100, 101, 463, 1991, 1]

[EC 3.1.8.2 created 1961 as EC 3.8.2.1, transferred 1992 to EC 3.1.8.2]

# EC 3.1.11 Exodeoxyribonucleases producing 5'-phosphomonoesters

EC 3.1.11.1

Accepted name: exodeoxyribonuclease I

**Reaction:** Exonucleolytic cleavage in the 3'- to 5'-direction to yield nucleoside 5'-phosphates

Other name(s): Escherichia coli exonuclease I; E. coli exonuclease I; exonuclease I

**Comments:** Preference for single-stranded DNA. The *Escherichia coli* enzyme hydrolyses glucosylated DNA.

**References:** [236, 1415, 1640]

[EC 3.1.11.1 created 1972 as EC 3.1.4.25, transferred 1978 to EC 3.1.11.1]

EC 3.1.11.2

Accepted name: exodeoxyribonuclease III

**Reaction:** Exonucleolytic cleavage in the 3'- to 5'-direction to yield nucleoside 5'-phosphates **Other name(s):** Escherichia coli exonuclease III; E. coli exonuclease III; endoribonuclease III

**Comments:** Preference for double-stranded DNA. Has endonucleolytic activity near apurinic sites on DNA.

**References:** [1700, 2419, 2420]

[EC 3.1.11.2 created 1972 as EC 3.1.4.27, transferred 1978 to EC 3.1.11.2]

EC 3.1.11.3

**Accepted name:** exodeoxyribonuclease (lambda-induced)

**Reaction:** Exonucleolytic cleavage in the 5'- to 3'-direction to yield nucleoside 5'-phosphates

Other name(s): lambda exonuclease; phage lambda-induced exonuclease; Escherichia coli exonuclease IV; E. coli

exonuclease IV; exodeoxyribonuclease IV; exonuclease IV

**Comments:** Preference for double-stranded DNA. Does not attack single-strand breaks.

**References:** [1699, 1713]

### [EC 3.1.11.3 created 1972 as EC 3.1.4.28, transferred 1978 to EC 3.1.11.3]

EC 3.1.11.4

**Accepted name:** exodeoxyribonuclease (phage SP<sub>3</sub>-induced)

**Reaction:** Exonucleolytic cleavage in the 5'- to 3'-direction to yield nucleoside 5'-phosphates

Other name(s): phage SP<sub>3</sub> DNase; DNA 5'-dinucleotidohydrolase; deoxyribonucleate 5'-dinucleotidase; deoxyri-

bonucleic 5'-dinucleotidohydrolase; bacteriophage SP<sub>3</sub> deoxyribonuclease; deoxyribonucleate 5'-

dinucleotidase

**Comments:** Preference for single-stranded DNA.

**References:** [2964]

[EC 3.1.11.4 created 1972 as EC 3.1.4.31, transferred 1978 to EC 3.1.11.4]

EC 3.1.11.5

**Accepted name:** exodeoxyribonuclease V

**Reaction:** Exonucleolytic cleavage (in the presence of ATP) in either 5'- to 3'- or 3'- to 5'-direction to yield 5'-

phosphooligonucleotides

Other name(s): Escherichia coli exonuclease V; E. coli exonuclease V; gene recBC endoenzyme; RecBC deoxyri-

bonuclease; gene recBC DNase; exonuclease V; gene recBCD enzymes

Comments: Preference for double-stranded DNA. Possesses DNA-dependent ATPase activity. Acts endonucle-

olytically on single-stranded circular DNA.

**References:** [685, 951, 2177, 3216]

[EC 3.1.11.5 created 1978]

EC 3.1.11.6

Accepted name: exodeoxyribonuclease VII

**Reaction:** Exonucleolytic cleavage in either 5'- to 3'- or 3'- to 5'-direction to yield nucleoside 5'-phosphates **Other name(s):** Escherichia coli exonuclease VII; E. coli exonuclease VII; endodeoxyribonuclease VII; exonuclease

VII

**Comments:** Preference for single-stranded DNA.

**References:** [402, 401]

[EC 3.1.11.6 created 1978]

EC 3.1.11.7

**Accepted name:** adenosine-5'-diphospho-5'-[DNA] diphosphatase

**Reaction:** (1) adenosine-5'-diphospho-5'-[DNA] +  $H_2O = AMP + phospho-5'-[DNA]$ 

(2) adenosine-5'-diphospho-5'-(ribonucleotide)-[DNA] +  $H_2O = AMP + 5'$ -phospho-(ribonucleotide)-

[DNA]

Other name(s): aprataxin; 5'-App5'-DNA adenylate hydrolase; APTX (gene name); HNT3 (gene name)

**Systematic name:** adenosine-5'-diphospho-5'-[DNA] hydrolase (adenosine 5'-phosphate-forming)

**Comments:** Aprataxin is a DNA-binding protein involved in different types of DNA break repair. The enzyme

acts (among other activities) on abortive DNA ligation intermediates that contain an adenylate covalently linked to the 5'-phosphate DNA terminus. It also acts when the adenylate is covalently linked to the 5'-phosphate of a ribonucleotide linked to a DNA strand, which is the result of abortive ligase activity on products of EC 3.1.26.4, ribonuclease H, an enzyme that cleaves RNA-DNA hybrids on the 5' side of the ribonucleotide found in the 5'-RNA-DNA-3' junction. Aprataxin binds the adenylate group to a histidine residue within the active site, followed by its hydrolysis from the nucleic acid and eventual release, leaving a 5'-phosphate terminus that can be efficiently rejoined. The enzyme also possesses the activities of EC 3.1.11.8, guanosine-5'-diphospho-5'-[DNA] diphosphatase, and EC

3.1.12.2, DNA-3'-diphospho-5'-guanosine diphosphatase.

**References:** [22, 2996]

#### [EC 3.1.11.7 created 2017]

EC 3.1.11.8

**Accepted name:** guanosine-5'-diphospho-5'-[DNA] diphosphatase

**Reaction:** guanosine-5'-diphospho-5'-[DNA] +  $H_2O$  = phospho-5'-[DNA] + GMP

**Other name(s):** aprataxin; pp5'G5'DNA diphosphatase; pp5'G5'-DNA guanylate hydrolase; APTX (gene name);

HNT3 (gene name)

**Systematic name:** guanosine-5'-diphospho-5'-[DNA] hydrolase (guanosine 5'-phosphate-forming)

**Comments:** Aprataxin is a DNA-binding protein that catalyses (among other activities) the 5' decapping of Gpp-

DNA (formed by homologs of RtcB3 from the bacterium *Myxococcus xanthus*). The enzyme binds the guanylate group to a histidine residue at its active site, forming a covalent enzyme-nucleotide phosphate intermediate, followed by the hydrolysis of the guanylate from the nucleic acid and eventual release. The enzyme forms a 5'-phospho terminus that can be efficiently joined by "classical" ligases. The enzyme also possesses the activity of EC 3.1.11.7, adenosine-5'-diphospho-5'-[DNA]

diphosphatase and EC 3.1.12.2, DNA-3'-diphospho-5'-guanosine diphosphatase.

**References:** [1849]

[EC 3.1.11.8 created 2017]

# EC 3.1.12 Exodeoxyribonucleases producing 3'-phosphomonoesters

EC 3.1.12.1

**Accepted name:** 5' to 3' exodeoxyribonuclease (nucleoside 3'-phosphate-forming)

**Reaction:** exonucleolytic cleavage in the 5'- to 3'-direction to yield nucleoside 3'-phosphates

**Other name(s):** Cas4; 5' to 3' single stranded DNA exonuclease

**Comments:** Preference for single-stranded DNA. The enzyme from the archaeon *Sulfolobus solfataricus* contains

a [4Fe-4S] cluster and requires a divalent metal cation, such as Mg<sup>2+</sup> or Mn<sup>2+</sup>, for activity.

**References:** [3326, 1642]

[EC 3.1.12.1 created 2014]

EC 3.1.12.2

**Accepted name:** DNA-3'-diphospho-5'-guanosine diphosphatase

**Reaction:** [DNA]-3'-diphospho-5'-guanosine +  $H_2O = [DNA]-3'$ -phosphate + GMP

Other name(s): aprataxin; DNA-3'pp5'G guanylate hydrolase; APTX (gene name); HNT3 (gene name)

Systematic name: [DNA]-3'-diphospho-5'-guanosine hydrolase (guanosine 5'-phosphate-forming)

**Comments:** Aprataxin is a DNA-binding protein that catalyses (among other activities) the 3' decapping of DNA-

ppG (formed by EC 6.5.1.8, 3'-phosphate/5'-hydroxy nucleic acid ligase) [538]. The enzyme binds the guanylate group to a histidine residue at its active site, forming a covalent enzyme-nucleotide phosphate intermediate, followed by the hydrolysis of the guanylate from the nucleic acid and its eventual release. The enzyme also possesses the activity of EC 3.1.11.7, adenosine-5'-diphospho-5'-

[DNA] diphosphatase, and EC 3.1.11.8, guanosine-5'-diphospho-5'-[DNA] diphosphatase.

**References:** [538, 406]

[EC 3.1.12.2 created 2017]

# EC 3.1.13 Exoribonucleases producing 5'-phosphomonoesters

EC 3.1.13.1

Accepted name: exoribonuclease II

**Reaction:** Exonucleolytic cleavage in the 3'- to 5'-direction to yield nucleoside 5'-phosphates

Other name(s): ribonuclease II; ribonuclease Q; BN ribonuclease; Escherichia coli exo-RNase II; RNase II; exori-

bonuclease (misleading); 5'-exoribonuclease (misleading)

**Comments:** Preference for single-stranded RNA. The enzyme processes 3'-terminal extra-nucleotides of

monomeric tRNA precursors, following the action of EC 3.1.26.5 ribonuclease P.

**References:** [2117, 2573, 2635, 2742]

[EC 3.1.13.1 created 1972 as EC 3.1.4.20, transferred 1978 to EC 3.1.13.1]

#### EC 3.1.13.2

Accepted name: exoribonuclease H

**Reaction:** 3'-end directed exonucleolytic cleavage of viral RNA-DNA hybrid

**Comments:** This is a secondary reaction to the RNA 5'-end directed cleavage 13-19 nucleotides from the RNA

end performed by EC 3.1.26.13 (retroviral ribonuclease H).

**References:** [2555]

[EC 3.1.13.2 created 1978, modified 2010]

#### EC 3.1.13.3

Accepted name: oligonucleotidase

**Reaction:** Exonucleolytic cleavage of oligonucleotides to yield nucleoside 5'-phosphates

Other name(s): oligoribonuclease

**Comments:** Also hydrolyses NAD<sup>+</sup> to NMN and AMP.

**References:** [866]

[EC 3.1.13.3 created 1972 as EC 3.1.4.19, transferred 1978 to EC 3.1.13.3]

#### EC 3.1.13.4

Accepted name: poly(A)-specific ribonuclease

**Reaction:** Exonucleolytic cleavage of poly(A) to 5'-AMP **Other name(s):** 3'-exoribonuclease; 2',3'-exoribonuclease

**Comments:** Cleaves poly(A) in either the single- or double-stranded form.

**References:** [2586]

[EC 3.1.13.4 created 1984]

#### EC 3.1.13.5

**Accepted name:** ribonuclease D

**Reaction:** Exonucleolytic cleavage that removes extra residues from the 3'-terminus of tRNA to produce 5'-

mononucleotides

Other name(s): RNase D

**Comments:** Requires divalent cations for activity  $(Mg^{2+}, Mn^{2+} \text{ or } Co^{2+})$ . Alteration of the 3'-terminal base has

no effect on the rate of hydrolysis whereas modification of the 3'-terminal sugar has a major effect. tRNA terminating with a 3'-phosphate is completely inactive [512]. This enzyme can convert a tRNA

precursor into a mature tRNA [513].

**References:** [906, 513, 512, 3327]

[EC 3.1.13.5 created 2006]

#### EC 3.1.14 Exoribonucleases producing 3'-phosphomonoesters

EC 3.1.14.1

Accepted name: yeast ribonuclease

**Reaction:** Exonucleolytic cleavage to nucleoside 3'-phosphates

**Comments:** Similar enzyme: RNase U<sub>4</sub>.

**References:** [2212]

[EC 3.1.14.1 created 1978]

# EC 3.1.15 Exonucleases that are active with either ribo- or deoxyribonucleic acids and produce 5'-phosphomonoesters

EC 3.1.15.1

Accepted name: venom exonuclease

**Reaction:** Exonucleolytic cleavage in the 3'- to 5'- direction to yield nucleoside 5'-phosphates

Other name(s): venom phosphodiesterase

**Comments:** Preference for single-stranded substrate.

**References:** [1607]

[EC 3.1.15.1 created 1978]

# EC 3.1.16 Exonucleases that are active with either ribo- or deoxyribonucleic acids and produce 3'-phosphomonoesters

EC 3.1.16.1

Accepted name: spleen exonuclease

**Reaction:** Exonucleolytic cleavage in the 5'- to 3'-direction to yield nucleoside 3'-phosphates

Other name(s): 3'-exonuclease; spleen phosphodiesterase; 3'-nucleotide phosphodiesterase; phosphodiesterase II

**Comments:** Preference for single-stranded substrate.

**References:** [199]

[EC 3.1.16.1 created 1972 as EC 3.1.4.18, transferred 1978 to EC 3.1.16.1]

#### EC 3.1.21 Endodeoxyribonucleases producing 5'-phosphomonoesters

EC 3.1.21.1

Accepted name: deoxyribonuclease I

**Reaction:** Endonucleolytic cleavage to 5'-phosphodinucleotide and 5'-phosphooligonucleotide end-products **Other name(s):** pancreatic DNase; DNase; thymonuclease, dornase; dornava; dornavac; pancreatic deoxyribonuclease, dornase; dovayribonuclease, dornase; dovayribonuclease, d

ase; pancreatic dornase; deoxyribonuclease (pancreatic); pancreatic DNase; DNAase; deoxyribonucleic phosphatase; DNase I; alkaline deoxyribonuclease; alkaline DNase; endodeoxyribonuclease I; DNA depolymerase; *Escherichia coli* endonuclease I; deoxyribonuclease A; DNA endonuclease;

DNA nuclease

**Comments:** Preference for double-stranded DNA.

**References:** [557, 1555, 1608]

[EC 3.1.21.1 created 1961 as EC 3.1.4.5, transferred 1978 to EC 3.1.21.1, modified 1981]

EC 3.1.21.2

**Accepted name:** deoxyribonuclease IV

**Reaction:** Endonucleolytic cleavage of ssDNA at apurinic/apyrimidinic sites to 5'-phosphooligonucleotide end-

products

Other name(s): deoxyribonuclease IV (phage-T<sub>4</sub>-induced) (misleading); endodeoxyribonuclease IV (phage T<sub>4</sub>-

induced) (misleading);  $E.\ coli$  endonuclease IV; endodeoxyribonuclease (misleading); redoxyendonuclease; deoxriboendonuclease (misleading); endonuclease II; endonuclease IV; DNA-adenine-

transferase; nfo (gene name)

Comments: The enzyme is an apurinic/apyrimidinic (AP) site endonuclease that primes DNA repair synthesis at

AP sites. It specifically cleaves the DNA backbone at AP sites and also removes 3' DNA-blocking groups such as 3' phosphates, 3' phosphoglycolates, and 3'  $\alpha,\beta$ -unsaturated aldehydes that arise from oxidative base damage and the activity of combined glycosylase/lyase enzymes. It is also the only known repair enzyme that is able to cleave the DNA backbone 5' of the oxidative lesion  $\alpha$ -

deoxyadenosine. The enzyme has a strong preference for single-stranded DNA.

**References:** [819, 820, 1016, 520, 1221, 1183]

[EC 3.1.21.2 created 1972 as EC 3.1.4.30, transferred 1978 to EC 3.1.21.2, modified 2014]

#### EC 3.1.21.3

**Accepted name:** type I site-specific deoxyribonuclease

**Reaction:** Endonucleolytic cleavage of DNA to give random double-stranded fragments with terminal 5'-

phosphates; ATP is simultaneously hydrolysed

**Other name(s):** type I restriction enzyme; deoxyribonuclease (ATP- and S-adenosyl-L-methionine-dependent);

restriction-modification system; deoxyribonuclease (adenosine triphosphate-hydrolyzing); adenosine triphosphate-dependent deoxyribonuclease; ATP-dependent DNase; type 1 site-specific deoxyribonu-

clease

**Comments:** This is a large group of enzymes which, together with those now listed as EC 3.1.21.4 (type II site-

specific deoxyribonuclease) and EC 3.1.21.5 (type III site-specific deoxyribonuclease), were previously listed separately in sub-subclasses EC 3.1.23 and EC 3.1.24. They have an absolute requirement for ATP (or dATP) and S-adenosyl-L-methionine. They recognize specific short DNA sequences and cleave at sites remote from the recognition sequence. They are multifunctional proteins that also catalyse the reactions of EC 2.1.1.72 [site-specific DNA-methyltransferase (adenine-specific)] and EC

2.1.1.37

**References:** [2437]

[EC 3.1.21.3 created 1984 from EC 3.1.23 and EC 3.1.24]

### EC 3.1.21.4

**Accepted name:** type II site-specific deoxyribonuclease

**Reaction:** Endonucleolytic cleavage of DNA to give specific double-stranded fragments with terminal 5'-

phosphates

Other name(s): type II restriction enzyme

**Comments:** This is a large group of enzymes which, together with those now listed as EC 3.1.21.3 (type 1 site-

specific deoxyribonuclease) and EC 3.1.21.5.

**References:** [2437]

[EC 3.1.21.4 created 1984 from EC 3.1.23 and EC 3.1.24]

#### EC 3.1.21.5

**Accepted name:** type III site-specific deoxyribonuclease

**Reaction:** Endonucleolytic cleavage of DNA to give specific double-stranded fragments with terminal 5'-

phosphates

**Other name(s):** type III restriction enzyme; restriction-modification system

**Comments:** This is a large group of enzymes which, together with those now listed as EC 3.1.21.3 (type 1 site-

specific deoxyribonuclease) and EC 3.1.21.4 (type II site-specific deoxyribonuclease), were previously listed separately in sub-subclasses EC 3.1.23 and EC 3.1.24. They have an absolute requirement for ATP but do not hydrolyse it; *S*-adenosy-L-methionine stimulates the reaction, but is not absolutely required. They recognize specific, short DNA sequences and cleave a short distance away from the recognition sequence. These enzymes exist as complexes with enzymes of similar specificity listed under EC 2.1.1.72 [site-specific DNA-methyltransferase (adenine-specific)] or EC 2.1.1.73

**References:** [2437]

[EC 3.1.21.5 created 1984 from EC 3.1.23 and EC 3.1.24]

EC 3.1.21.6

Accepted name: CC-preferring endodeoxyribonuclease

**Reaction:** endonucleolytic cleavage to give 5'-phosphooligonucleotide end-products, with a preference for

cleavage within the sequence CC

**Other name(s):** Streptomyces glaucescens exocytoplasmic dodeoxyribonuclease

**Comments:** Prefers CC sites in double-stranded circular and linear DNA. Greater affinity for double-stranded

than single-stranded DNA. Produces nicks, generating double-stranded fragments with 5'- and/or 3'-protruding single-stranded tails. Requires magnesium ions for activity. The endonuclease from

Chlorella-like green algae infected with NYs-1 virus 4[3228] may be the same enzyme.

**References:** [3228, 62]

[EC 3.1.21.6 created 1999]

EC 3.1.21.7

Accepted name: deoxyribonuclease V

**Reaction:** Endonucleolytic cleavage at apurinic or apyrimidinic sites to products with a 5'-phosphate

Other name(s): endodeoxyribonuclease V; DNase V; Escherichia coli endodeoxyribonuclease V

**Comments:** Previously classified erroneously as EC 3.1.22.3.

**References:** [882]

[EC 3.1.21.7 created 1978 as EC 3.1.22.3, transferred 2001 to EC 3.1.21.7]

EC 3.1.21.8

**Accepted name:** T<sub>4</sub> deoxyribonuclease II

**Reaction:** Endonucleolytic nicking and cleavage of cytosine-containing double-stranded DNA.

**Other name(s):** T<sub>4</sub> endonuclease II; EndoII (ambiguous); *denA* (gene name)

**Comments:** Requires Mg<sup>2+</sup>. This phage T<sub>4</sub> enzyme is involved in degradation of host DNA. The enzyme primar-

ily catalyses nicking of the bottom strand of double stranded DNA between the first and second base pair to the right of a top-strand CCGC motif. Double-stranded breaks are produced 5- to 10-fold less frequently [370]. It does not cleave the T4 native DNA, which contains 5-hydroxymethylcytosine in-

stead of cytosine.

**References:** [371, 369, 370, 53]

[EC 3.1.21.8 created 2014]

EC 3.1.21.9

**Accepted name:** T<sub>4</sub> deoxyribonuclease IV

**Reaction:** Endonucleolytic cleavage of the 5' phosphodiester bond of deoxycytidine in single-stranded DNA.

**Other name(s):** T<sub>4</sub> endonuclease IV; EndoIV (ambiguous); *denB* (gene name)

**Comments:** This phage T<sub>4</sub> enzyme is involved in degradation of host DNA. The enzyme does not cleave double-

stranded DNA or native T4 DNA, which contains 5-hydroxymethylcytosine instead of cytosine.

**References:** [2489, 1705, 2488, 200, 1147, 2167]

# EC 3.1.22 Endodeoxyribonucleases producing 3'-phosphomonoesters

EC 3.1.22.1

Accepted name: deoxyribonuclease II

**Reaction:** Endonucleolytic cleavage to nucleoside 3'-phosphates and 3'-phosphooligonucleotide end-products **Other name(s):** DNase II; pancreatic DNase II; deoxyribonucleate 3'-nucleotidohydrolase; DNase II; pancreatic

DNase II; acid deoxyribonuclease; acid DNase

**Comments:** Preference for double-stranded DNA.

**References:** [201]

 $[EC\ 3.1.22.1\ created\ 1961\ as\ EC\ 3.1.4.6,\ transferred\ 1978\ to\ EC\ 3.1.22.1,\ modified\ 1981]$ 

EC 3.1.22.2

**Accepted name:** Aspergillus deoxyribonuclease K<sub>1</sub>

**Reaction:** Endonucleolytic cleavage to nucleoside 3'-phosphates and 3'-phosphooligonucleotide end-products

Other name(s): Aspergillus DNase  $K_1$ 

**Comments:** Preference for single-stranded DNA.

**References:** [1761, 2644]

[EC 3.1.22.2 created 1978, modified 1981]

[3.1.22.3 Transferred entry. deoxyribonuclease V. Now EC 3.1.21.7, deoxyribonuclease V]

[EC 3.1.22.3 created 1978, deleted 2001]

EC 3.1.22.4

**Accepted name:** crossover junction endodeoxyribonuclease

Reaction: Endonucleolytic cleavage at a junction such as a reciprocal single-stranded crossover between two

homologous DNA duplexes (Holliday junction)

Other name(s): Hie endonuclease; Holliday junction endonuclease CCE1; Holliday junction resolvase; Holliday

junction-cleaving endonuclease; Holliday junction-resolving endoribonuclease; RusA Holliday junction resolvase; RusA endonuclease; RuvC endonuclease; SpCCe<sub>1</sub> Holliday junction resolvase; crossover junction endoribonuclease; cruciform-cutting endonuclease; endo X3; endonuclease RuvC;

endonuclease VII; endonuclease X3; resolving enzyme CCE1

Comments: The enzyme from Saccharomyces cerevisiae has no endonuclease or exonuclease activity on single-

stranded or double-stranded DNA molecules that do not contain Holliday junctions.

**References:** [2822, 2630, 2616, 787, 1907]

[EC 3.1.22.4 created 1989, modified 2003]

EC 3.1.22.5

**Accepted name:** deoxyribonuclease X

**Reaction:** Endonucleolytic cleavage of supercoiled plasma DNA to linear DNA duplexes **Other name(s):** Escherichia coli endodeoxyribonuclease; Escherichia coli endodeoxyribonuclease X

Comments: Preference for supercoiled DNA; little activity on linear double-stranded DNA. Inhibited by single-

stranded DNA, ATP and AMP.

**References:** [907]

[EC 3.1.22.5 created 1992]

# EC 3.1.23 Site-specific endodeoxyribonucleases: cleavage is sequence specific (deleted sub-subclass)

[3.1.23.1	Transferred entry. endodeoxyribonuclease AluI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.1 created 1978, deleted 1984]
[3.1.23.2	Transferred entry. endodeoxyribonuclease AsuI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.2 created 1978, deleted 1984]
[3.1.23.3	Transferred entry. endodeoxyribonuclease AvaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.3 created 1978, deleted 1984]
[3.1.23.4	Transferred entry. endodeoxyribonuclease AvaII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.4 created 1978, deleted 1984]
[3.1.23.5	Transferred entry. endodeoxyribonuclease Ball. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.5 created 1978, deleted 1984]
[3.1.23.6	Transferred entry. endodeoxyribonuclease BamHI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.6 created 1978, deleted 1984]
[3.1.23.7	Transferred entry. endodeoxyribonuclease BbvI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.7 created 1978, deleted 1984]
[3.1.23.8	Transferred entry. endodeoxyribonuclease BclI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.8 created 1978, deleted 1984]
[3.1.23.9	Transferred entry. endodeoxyribonuclease BglI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.9 created 1978, deleted 1984]
[3.1.23.10	Transferred entry. endodeoxyribonuclease BglII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.10 created 1978, deleted 1984]
[3.1.23.11	Transferred entry. endodeoxyribonuclease BpuI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.11 created 1978, deleted 1984]
[3.1.23.12	Transferred entry. endodeoxyribonuclease DpnI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.12 created 1978, modified 1982, deleted 1984]
[3.1.23.13	Transferred entry. endodeoxyribonuclease EcoRI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.13 created 1978, deleted 1984]
[3.1.23.14	Transferred entry. endodeoxyribonuclease EcoRII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.14 created 1978, deleted 1984]
[3.1.23.15	Transferred entry. endodeoxyribonuclease HaeI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.15 created 1978, deleted 1984]
[3.1.23.16	Transferred entry. endodeoxyribonuclease HaeII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.16 created 1978, deleted 1984]
[3.1.23.17	Transferred entry. endodeoxyribonuclease HaeIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]

	[EC 3.1.23.17 created 1978, deleted 1984]
[3.1.23.18	Transferred entry. endodeoxyribonuclease HgaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.18 created 1978, deleted 1984]
[3.1.23.19	Transferred entry. endodeoxyribonuclease HhaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.19 created 1978, deleted 1984]
[3.1.23.20	Transferred entry. endodeoxyribonuclease HindII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.20 created 1978, deleted 1984]
[3.1.23.21	Transferred entry. endodeoxyribonuclease HindIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.21 created 1978, deleted 1984]
[3.1.23.22	Transferred entry. endodeoxyribonuclease Hinfl. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.22 created 1978, deleted 1984]
[3.1.23.23	Transferred entry. endodeoxyribonuclease HpaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.23 created 1978, deleted 1984]
[3.1.23.24	Transferred entry. endodeoxyribonuclease HpaII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.24 created 1978, deleted 1984]
[3.1.23.25	Transferred entry. endodeoxyribonuclease HphI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.25 created 1978, deleted 1984]
[3.1.23.26	Transferred entry. endodeoxyribonuclease KpnI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.26 created 1978, deleted 1984]
[3.1.23.27	Transferred entry. endodeoxyribonuclease MboI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.27 created 1978, deleted 1984]
[3.1.23.28	Transferred entry. endodeoxyribonuclease MboII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.28 created 1978, deleted 1984]
[3.1.23.29	Transferred entry. endodeoxyribonuclease Mnll. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.29 created 1978, deleted 1984]
[3.1.23.30	Transferred entry. endodeoxyribonuclease PfaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.30 created 1978, modified 1982, deleted 1984]
[3.1.23.31	Transferred entry. endodeoxyribonuclease PstI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.31 created 1978, deleted 1984]
[3.1.23.32	Transferred entry. endodeoxyribonuclease PvuI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.32 created 1978, modified 1982, deleted 1984]
[3.1.23.33	Transferred entry. endodeoxyribonuclease PvuII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.33 created 1978, deleted 1984]
[3.1.23.34	Transferred entry. endodeoxyribonuclease SacI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.34 created 1978, deleted 1984]

[3.1.23.35	Transferred entry. endodeoxyribonuclease SacII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.35 created 1978, deleted 1984]
[3.1.23.36	Transferred entry. endodeoxyribonuclease SacIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.36 created 1978, deleted 1984]
[3.1.23.37	Transferred entry. endodeoxyribonuclease Sall. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.37 created 1978, deleted 1984]
[3.1.23.38	Transferred entry. endodeoxyribonuclease SgrI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.38 created 1978, deleted 1984]
[3.1.23.39	Transferred entry. endodeoxyribonuclease TaqI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.39 created 1978, deleted 1984]
[3.1.23.40	Transferred entry. endodeoxyribonuclease TaqII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.40 created 1978, deleted 1984]
[3.1.23.41	Transferred entry. endodeoxyribonuclease XbaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.41 created 1978, deleted 1984]
[3.1.23.42	Transferred entry. endodeoxyribonuclease XhoI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.42 created 1978, deleted 1984]
[3.1.23.43	Transferred entry. endodeoxyribonuclease XhoII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.43 created 1978, modified 1982, deleted 1984]
[3.1.23.44	Transferred entry. endodeoxyribonuclease XmaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.44 created 1978, deleted 1984]
[3.1.23.45	Transferred entry. endodeoxyribonuclease XniI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.45 created 1978, modified 1982, deleted 1984]
[3.1.23.46	Transferred entry. endodeoxyribonuclease AimI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.46 created 1982, deleted 1984]
[3.1.23.47	Transferred entry. endodeoxyribonuclease AccI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.47 created 1982, deleted 1984]
[3.1.23.48	Transferred entry. endodeoxyribonuclease AccII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.48 created 1982, deleted 1984]
[3.1.23.49	Transferred entry. endodeoxyribonuclease AtuAI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.49 created 1982, deleted 1984]
[3.1.23.50	Transferred entry. endodeoxyribonuclease AtuBVI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease
	[EC 3.1.23.50 created 1982, deleted 1984]
[3.1.23.51	Transferred entry. endodeoxyribonuclease AcaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.51 created 1982, deleted 1984]
[3.1.23.52	Transferred entry, endodeoxyribonuclease Acyl. Now EC 3.1.21.4, type II site-specific deoxyribonuclease l

	[EC 3.1.23.52 created 1982, deleted 1984]
[3.1.23.53	Transferred entry. endodeoxyribonuclease AosI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.53 created 1982, deleted 1984]
[3.1.23.54	Transferred entry. endodeoxyribonuclease AsuII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.54 created 1982, deleted 1984]
[3.1.23.55	Transferred entry. endodeoxyribonuclease AvaIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.55 created 1982, deleted 1984]
[3.1.23.56	Transferred entry. endodeoxyribonuclease AvrII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.56 created 1982, deleted 1984]
[3.1.23.57 Assumed to be	Transferred entry. endodeoxyribonuclease BceI4579. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. the same as endodeoxyribonuclease Bce4579I (see http://rebase.neb.com/rebase/rebase.html)]
	[EC 3.1.23.57 created 1982, deleted 1984]
[3.1.23.58 Assumed to be	Transferred entry. endodeoxyribonuclease Bce1229. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. the same as endodeoxyribonuclease Bce1229I (see http://rebase.neb.com/rebase/rebase.html)]
	[EC 3.1.23.58 created 1982, deleted 1984]
[3.1.23.59 Assumed to be	Transferred entry. endodeoxyribonuclease Bme899. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. the same as endodeoxyribonuclease Bme899I (see http://rebase.neb.com/rebase/rebase.html)]
	[EC 3.1.23.59 created 1982, deleted 1984]
[3.1.23.60 Assumed to be	Transferred entry. endodeoxyribonuclease Bme205. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. the same as endodeoxyribonuclease Bme205I (see http://rebase.neb.com/rebase/rebase.html)]
	[EC 3.1.23.60 created 1982, deleted 1984]
[3.1.23.61	Transferred entry. endodeoxyribonuclease BmeI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.61 created 1982, deleted 1984]
[3.1.23.62 Assumed to be	Transferred entry. endodeoxyribonuclease Bsp1286. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. the same as endodeoxyribonuclease Bsp1286I (see http://rebase.neb.com/rebase/rebase.html)]
	[EC 3.1.23.62 created 1982, deleted 1984]
[3.1.23.63	Transferred entry. endodeoxyribonuclease BstAI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.63 created 1982, deleted 1984]
[3.1.23.64	Transferred entry. endodeoxyribonuclease BstEI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.64 created 1982, deleted 1984]
[3.1.23.65	Transferred entry. endodeoxyribonuclease BstEIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.65 created 1982, deleted 1984]
[3.1.23.66	Transferred entry. endodeoxyribonuclease BstPI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.66 created 1982, deleted 1984]

[EC 3.1.23.67 created 1982, deleted 1984]

sumed to be the same as endodeoxyribonuclease BsuMI (see http://rebase.neb.com/rebase/rebase.html)]

[3.1.23.67

Transferred entry. endodeoxyribonuclease BsuM. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. As-

[3.1.23.68 Transferred entry. endodeoxyribonuclease Bsu6633. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. The name was misprinted in supplement 3 of the 1978 edition. Assumed to be the same as endodeoxyribonuclease Bsu6633I (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.23.68 created 1982, deleted 1984]

[3.1.23.69 Transferred entry. endodeoxyribonuclease Bsu1145. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease Bsu1145I (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.23.69 created 1982, deleted 1984]

[3.1.23.70 Transferred entry. endodeoxyribonuclease Bsu1192. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease Bsu1192I or see Bsu1192II (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.23.70 created 1982, deleted 1984]

[3.1.23.71 Transferred entry. endodeoxyribonuclease Bsu1193. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease Bsu1193I (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.23.71 created 1982, deleted 1984]

[3.1.23.72 Transferred entry. endodeoxyribonuclease Bsu1231. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. Not found in http://rebase.neb.com/rebase/rebase.html]

[EC 3.1.23.72 created 1982, deleted 1984]

[3.1.23.73 Transferred entry. endodeoxyribonuclease Bsu1259. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease Bsu1259I (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.23.73 created 1982, deleted 1984]

- [3.1.23.74 Transferred entry. endodeoxyribonuclease ClaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
  [EC 3.1.23.74 created 1982, deleted 1984]
- [3.1.23.75 Transferred entry. endodeoxyribonuclease CauII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
  [EC 3.1.23.75 created 1982, deleted 1984]
- [3.1.23.76 Transferred entry. endodeoxyribonuclease CviI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
  [EC 3.1.23.76 created 1982, deleted 1984]
- [3.1.23.77 Transferred entry. endodeoxyribonuclease DdeI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
  [EC 3.1.23.77 created 1982, deleted 1984]
- [3.1.23.78 Transferred entry. endodeoxyribonuclease EclI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
  [EC 3.1.23.78 created 1982, deleted 1984]
- [3.1.23.79 Transferred entry. endodeoxyribonuclease Ecal. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
  [EC 3.1.23.79 created 1982, deleted 1984]
- [3.1.23.80 Transferred entry. endodeoxyribonuclease EcoRI'. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease EcoRI' (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.23.80 created 1982, deleted 1984]

- [3.1.23.81 Transferred entry. endodeoxyribonuclease Fnu48I. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
  [EC 3.1.23.81 created 1982, deleted 1984]
- [3.1.23.82 Transferred entry. endodeoxyribonuclease Fnu4H. Now EC 3.1.21.4, type II site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease Fnu4HI (see http://rebase.neb.com/rebase/rebase.html)]

	[EC 3.1.23.82 created 1982, deleted 1984]
[3.1.23.83	Transferred entry. endodeoxyribonuclease HapI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.83 created 1982, deleted 1984]
[3.1.23.84	Transferred entry. endodeoxyribonuclease Hin1056II. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.84 created 1982, deleted 1984]
[3.1.23.85	Transferred entry. endodeoxyribonuclease HinfIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.85 created 1982, deleted 1984]
[3.1.23.86	Transferred entry. endodeoxyribonuclease HgiAI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.86 created 1982, deleted 1984]
[3.1.23.87	Transferred entry. endodeoxyribonuclease HgiCI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.87 created 1982, deleted 1984]
[3.1.23.88	Transferred entry. endodeoxyribonuclease HgiDI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.88 created 1982, deleted 1984]
[3.1.23.89	Transferred entry. endodeoxyribonuclease HgiEII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.89 created 1982, deleted 1984]
[3.1.23.90	Transferred entry. endodeoxyribonuclease MstI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.90 created 1982, deleted 1984]
[3.1.23.91	Transferred entry. endodeoxyribonuclease MstII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.91 created 1982, deleted 1984]
[3.1.23.92	Transferred entry. endodeoxyribonuclease MglI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.92 created 1982, deleted 1984]
[3.1.23.93	Transferred entry. endodeoxyribonuclease MglII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.93 created 1982, deleted 1984]
[3.1.23.94	Transferred entry. endodeoxyribonuclease MnoII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.94 created 1982, deleted 1984]
[3.1.23.95	Transferred entry. endodeoxyribonuclease MnnIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.95 created 1982, deleted 1984]
[3.1.23.96	Transferred entry. endodeoxyribonuclease MviI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.96 created 1982, deleted 1984]
[3.1.23.97	Transferred entry. endodeoxyribonuclease MviII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.97 created 1982, deleted 1984]
[3.1.23.98	Transferred entry. endodeoxyribonuclease OxaII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.98 created 1982, deleted 1984]

Assumed to be the same as endodeoxyribonuclease PaeR7I (see http://rebase.neb.com/rebase/rebase.html)]

Transferred entry. endodeoxyribonuclease PaeR7. Now EC 3.1.21.4, type II site-specific deoxyribonuclease.

[3.1.23.99

	[EC 3.1.23.99 created 1982, deleted 1984]
[3.1.23.100	Transferred entry. endodeoxyribonuclease RspI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.100 created 1982, deleted 1984]
[3.1.23.101	Transferred entry. endodeoxyribonuclease RsaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.101 created 1982, deleted 1984]
[3.1.23.102	Transferred entry. endodeoxyribonuclease Smal. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.102 created 1982, deleted 1984]
[3.1.23.103	Transferred entry. endodeoxyribonuclease SspI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.103 created 1982, deleted 1984]
[3.1.23.104	Transferred entry. endodeoxyribonuclease SnaI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.104 created 1982, deleted 1984]
[3.1.23.105	Transferred entry. endodeoxyribonuclease SfaNI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.105 created 1982, deleted 1984]
[3.1.23.106	Transferred entry. endodeoxyribonuclease SalII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.106 created 1982, deleted 1984]
[3.1.23.107	Transferred entry. endodeoxyribonuclease Saul. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.107 created 1982, deleted 1984]
[3.1.23.108	Transferred entry. endodeoxyribonuclease SphI. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.108 created 1982, deleted 1984]
[3.1.23.109	Transferred entry. endodeoxyribonuclease XmaIII. Now EC 3.1.21.4, type II site-specific deoxyribonuclease]
	[EC 3.1.23.109 created 1982, deleted 1984]

# EC 3.1.24 Site specific endodeoxyribonucleases: cleavage is not sequence specific (deleted sub-subclass)

[3.1.24.1 Transferred entry. endodeoxyribonuclease EcoB. Now EC 3.1.21.3, type I site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease EcoBI (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.24.1 created 1978, modified 1982, deleted 1984]

[3.1.24.2 Transferred entry. endodeoxyribonuclease EcoK. Now EC 3.1.21.3, type I site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease EcoKI (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.24.2 created 1978, modified 1982, deleted 1984]

[3.1.24.3 Transferred entry. endodeoxyribonuclease EcoPI. Now EC 3.1.21.5, type III site-specific deoxyribonuclease. The name is misprinted in supplement 3 of the 1978 edition]

[EC 3.1.24.3 created 1978, modified 1982, deleted 1984]

[3.1.24.4 Transferred entry. endodeoxyribonuclease EcoP15. Now EC 3.1.21.5, type III site-specific deoxyribonuclease. Assumed to be the same as endodeoxyribonuclease EcoP15I (see http://rebase.neb.com/rebase/rebase.html)]

[EC 3.1.24.4 created 1978, modified 1982, deleted 1984]

## EC 3.1.25 Site-specific endodeoxyribonucleases that are specific for altered bases

#### EC 3.1.25.1

**Accepted name:** deoxyribonuclease (pyrimidine dimer)

**Reaction:** Endonucleolytic cleavage near pyrimidine dimers to products with 5'-phosphate

Other name(s): endodeoxyribonuclease (pyrimidine dimer); endodeoxyribonuclease (pyrimidine dimer); bacterio-

phage T<sub>4</sub> endodeoxyribonuclease V; T4 endonuclease V

**Comments:** Acts on a damaged strand, 5' from the damaged site.

**References:** [279, 2416]

[EC 3.1.25.1 created 1978]

[3.1.25.2 Transferred entry. endodeoxyribonuclease (apurinic or apyrimidinic). Now EC 4.2.99.18, DNA-(apurinic or apyrimidinic site) lyase]

[EC 3.1.25.2 created 1978, deleted 1992]

# EC 3.1.26 Endoribonucleases producing 5'-phosphomonoesters

#### EC 3.1.26.1

Accepted name: Physarum polycephalum ribonuclease

**Reaction:** Endonucleolytic cleavage to 5'-phosphomonoester

**References:** [1145]

[EC 3.1.26.1 created 1978]

#### EC 3.1.26.2

**Accepted name:** ribonuclease α

**Reaction:** Endonucleolytic cleavage to 5'-phosphomonoester

**Other name(s):** 2'-O-methyl RNase

**Comments:** Specific for *O*-methylated RNA.

**References:** [2116]

[EC 3.1.26.2 created 1978]

#### EC 3.1.26.3

Accepted name: ribonuclease III

**Reaction:** Endonucleolytic cleavage to a 5'-phosphomonoester

Other name(s): RNase III; ribonuclease 3

Comments: This is an endoribonuclease that cleaves double-stranded RNA molecules [994]. The cleavage can be

either a single-stranded nick or double-stranded break in the RNA, depending in part upon the degree of base-pairing in the region of the cleavage site [497]. Specificity is conferred by negative determinants, i.e., the presence of certain Watson-Crick base-pairs at specific positions that strongly inhibit cleavage [3328]. RNase III is involved in both rRNA processing and mRNA processing and decay.

**References:** [511, 2391, 2440, 994, 497, 3328]

[EC 3.1.26.3 created 1978, modified 2006]

#### EC 3.1.26.4

Accepted name: ribonuclease H

**Reaction:** Endonucleolytic cleavage to 5'-phosphomonoester

Other name(s): endoribonuclease H (calf thymus); RNase H; RNA\*DNA hybrid ribonucleotidohydrolase; hybrid ri-

bonuclease; hybridase; hybridase (ribonuclease H); ribonuclease H; hybrid nuclease; calf thymus ri-

bonuclease H

**Comments:** Acts on RNA-DNA hybrids.

**References:** [1014, 2757]

[EC 3.1.26.4 created 1978, modified 2010]

EC 3.1.26.5

Accepted name: ribonuclease P

> Endonucleolytic cleavage of RNA, removing 5'-extranucleotides from tRNA precursor Reaction:

Other name(s): RNase P

> **Comments:** An RNA-containing enzyme, essential for tRNA processing; generates 5'-termini or mature tRNA

> > molecules.

[215, 216, 2439] **References:** 

[EC 3.1.26.5 created 1978, modified 1982]

EC 3.1.26.6

Accepted name: ribonuclease IV

> Endonucleolytic cleavage of poly(A) to fragments terminated by 3'-hydroxy and 5'-phosphate groups **Reaction:**

Other name(s): endoribonuclease IV; poly(A)-specific ribonuclease

Forms oligonucleotides with an average chain length of 10. **Comments:** 

**References:** [1995, 1996]

[EC 3.1.26.6 created 1984]

EC 3.1.26.7

Accepted name: ribonuclease P4

> Endonucleolytic cleavage of RNA, removing 3'-extranucleotides from tRNA precursor Reaction:

**References:** [2610]

[EC 3.1.26.7 created 1984]

EC 3.1.26.8

Accepted name: ribonuclease M5

> Endonucleolytic cleavage of RNA, removing 21 and 42 nucleotides, respectively, from the 5'- and Reaction:

3'-termini of a 5S-rRNA precursor

Other name(s): RNase M5; 5S ribosomal maturation nuclease; 5S ribosomal RNA maturation endonuclease **Comments:** 

Converts the 5S-rRNA precursor from Bacillus subtilis into 5S-rRNA, with 5'-phosphate and 3'-

hydroxy groups.

**References:** [2708]

[EC 3.1.26.8 created 1986]

EC 3.1.26.9

Accepted name: ribonuclease [poly-(U)-specific]

> Reaction: Endonucleolytic cleavage of poly(U) to fragments terminated by 3'-hydroxy and 5'-phosphate groups

Other name(s): ribonuclease (uracil-specific); uracil-specific endoribonuclease; uracil-specific RNase

Forms oligonucleotides with chain lengths of 6 to 12. **Comments:** 

**References:** [115]

[EC 3.1.26.9 created 1986]

EC 3.1.26.10

**Accepted name:** ribonuclease IX

**Reaction:** Endonucleolytic cleavage of poly(U) or poly(C) to fragments terminated by 3'-hydroxy and 5'-

phosphate groups

**Other name(s):** poly(U)- and poly(C)-specific endoribonuclease

**Comments:** Acts on poly(U) and poly(C), with a higher affinity for poly(C), but does not act on poly(A) or

poly(G).

**References:** [2652]

[EC 3.1.26.10 created 1992]

EC 3.1.26.11

**Accepted name:** tRNase Z

**Reaction:** endonucleolytic cleavage of RNA, removing extra 3' nucleotides from tRNA precursor, generating 3'

termini of tRNAs. A 3'-hydroxy group is left at the tRNA terminus and a 5'-phosphoryl group is left

at the trailer molecule

Other name(s): 3 tRNase; tRNA 3 endonuclease; RNase Z; 3' tRNase

**Comments:** No cofactor requirements. An homologous enzyme to that found in *Arabidopsis thaliana* has been

found in Methanococcus janaschii.

**References:** [2565, 1853, 2564, 1558, 1979, 1921, 2851]

[EC 3.1.26.11 created 2002]

EC 3.1.26.12

Accepted name: ribonuclease E

**Reaction:** Endonucleolytic cleavage of single-stranded RNA in A- and U-rich regions

Other name(s): endoribonuclease E; RNase E; Rne protein

Comments: RNase E is a bacterial ribonuclease that plays a role in the processing of ribosomal RNA (9S to 5S

rRNA), the chemical degradation of bulk cellular RNA, the decay of specific regulatory, messenger and structural RNAs and the control of plasmid DNA replication [754]. The enzyme binds to monophosphorylated 5' ends of substrates but exhibits sequential cleavages in the 3' to 5' direction [754]. 2'-O-Methyl nucleotide substitutions at RNase E binding sites do not prevent binding but do prevent cleavage of non-modified target sequences 5' to that locus [754]. In *Escherichia coli*, the enzyme is found in the RNA degradosome. The C-terminal half of the protein contains binding sites for the three other major degradosomal components, the DEAD-box RNA helicase Rh1B, enolase (EC

4.1.1.11) and polynucleotide phosphorylase (EC 2.7.7.8).

**References:** [754, 683, 488, 3051, 2758, 356]

[EC 3.1.26.12 created 2008]

EC 3.1.26.13

**Accepted name:** retroviral ribonuclease H

**Reaction:** Endohydrolysis of RNA in RNA/DNA hybrids. Three different cleavage modes: 1. sequence-

specific internal cleavage of RNA [1-4]. Human immunodeficiency virus type 1 and Moloney murine leukemia virus enzymes prefer to cleave the RNA strand one nucleotide away from the RNA-DNA junction [5]. 2. RNA 5'-end directed cleavage 13-19 nucleotides from the RNA end [6,7]. 3. DNA

3'-end directed cleavage 15-20 nucleotides away from the primer terminus [8-10].

Other name(s): RT/RNase H; retroviral reverse transcriptase RNaseH; HIV RNase H

**Comments:** Comments: Retroviral reverse transcriptase is a multifunctional enzyme responsible for viral replica-

tion. To perform this task the enzyme combines two distinct activities. The polymerase domain (EC 2.7.7.49, RNA-directed DNA polymerase) occupies the N-terminal two-thirds of the reverse transcriptase whereas the ribonuclease H domain comprises the C-terminal remaining one-third [390, 2589]. The RNase H domain of Moloney murine leukemia virus and Human immunodeficiency virus display

two metal binding sites [937, 548, 2233]

**References:** [2590, 2527, 2378, 286, 2591, 591, 1396, 2225, 828, 185, 1203, 1540, 390, 2589, 937, 548, 2233]

### EC 3.1.27 Endoribonucleases producing 3'-phosphomonoesters

[3.1.27.1 Transferred entry. ribonuclease  $T_2$ . Now EC 4.6.1.19, ribonuclease  $T_2$ , since the primary reaction is that of a lyase]

[EC 3.1.27.1 created 1972 as EC 3.1.4.23, transferred 1978 to EC 3.1.27.1, modified 1981, deleted 2018]

[3.1.27.2 Transferred entry. Bacillus subtilis ribonuclease. Now EC 4.6.1.22, Bacillus subtilis ribonuclease, since the reaction catalysed is that of a lyase]

[EC 3.1.27.2 created 1978, deleted 2018]

#### EC 3.1.27.3

**Accepted name:** ribonuclease T<sub>1</sub>

**Reaction:** Two-stage endonucleolytic cleavage to nucleoside 3'-phosphotes and 3'-phosphooligonucleotides end-

ing in Gp with 2',3'-cyclic phosphate intermediates

Other name(s): guanyloribonuclease; Aspergillus oryzae ribonuclease; RNase N<sub>1</sub>; RNase N<sub>2</sub>; ribonuclease N<sub>3</sub>; ri-

bonuclease  $U_1$ ; ribonuclease F1; ribonuclease Ch; ribonuclease PP1; ribonuclease SA; RNase F1; ribonuclease C2; binase; RNase Sa; guanyl-specific RNase; RNase G; RNase  $T_1$ ; ribonuclease

guaninenucleotido-2'-transferase (cyclizing); ribonuclease N<sub>3</sub>; ribonuclease N<sub>1</sub>

**Comments:** Formerly EC 2.7.7.26 and EC 3.1.4.8.

**References:** [1392, 2837]

[EC 3.1.27.3 created 1961 as EC 3.1.4.8, transferred 1965 to EC 2.7.7.26, reinstated 1972 as EC 3.1.4.8, transferred 1978 to EC 3.1.27.3]

[3.1.27.4 Transferred entry. ribonuclease  $U_2$ . Now EC 4.6.1.20, ribonuclease  $U_2$ , since the primary reaction is that of a lyase]

[EC 3.1.27.4 created 1978, modified 1981, deleted 2018]

[3.1.27.5 Transferred entry. pancreatic ribonuclease. Now EC 4.6.1.18, pancreatic ribonuclease. This reaction is now known to involve an internal-transfer (lyase) process to produce the cyclic derivative, followed by a reversal of that step with water in the "hydrolytic step"]

[EC 3.1.27.5 created 1972 as EC 3.1.4.22, transferred 1978 to EC 3.1.27.5, modified 1981, deleted 2018]

[3.1.27.6 Transferred entry. Enterobacter ribonuclease. Now EC 4.6.1.21, Enterobacter ribonuclease, since the primary reaction is that of a lyase]

[EC 3.1.27.6 created 1978, modified 1981, deleted 2018]

#### EC 3.1.27.7

**Accepted name:** ribonuclease F

**Reaction:** Endonucleolytic cleavage of RNA precursor into two, leaving 5'-hydroxy and 3'-phosphate groups

**Other name(s):** ribonuclease F (*E. coli*)

**References:** [1004, 3138]

[EC 3.1.27.7 created 1984]

#### EC 3.1.27.8

**Accepted name:** ribonuclease V

**Reaction:** Hydrolysis of poly(A), forming oligoribonucleotides and ultimately 3'-AMP

**References:** [2585]

#### [EC 3.1.27.8 created 1984]

#### [3.1.27.9 Transferred entry. tRNA-intron endonuclease. Now EC 4.6.1.16, tRNA-intron lyase]

[EC 3.1.27.9 created 1992, deleted 2014]

#### EC 3.1.27.10

Accepted name: rRNA endonuclease

**Reaction:** Hydrolysis of the phosphodiester linkage between guanosine and adenosine residues at one specific

position in 28S rRNA from rat ribosomes

Other name(s):  $\alpha$ -sarcin

**Comments:** Also acts on bacterial rRNA.

**References:** [704]

[EC 3.1.27.10 created 1992]

# EC 3.1.30 Endoribonucleases that are active with either ribo- or deoxyribonucleic acids and produce 5'-phosphomonoesters

#### EC 3.1.30.1

**Accepted name:** Aspergillus nuclease S<sub>1</sub>

**Reaction:** Endonucleolytic cleavage to 5'-phosphomononucleotide and 5'-phosphooligonucleotide end-products **Other name(s):** endonuclease S<sub>1</sub> (*Aspergillus*); single-stranded-nucleate endonuclease; deoxyribonuclease S<sub>1</sub>; de-

oxyribonuclease S<sub>1</sub>; nuclease S<sub>1</sub>; *Neurospora crassa* single-strand specific endonuclease; S1 nuclease; single-strand endodeoxyribonuclease; single-stranded DNA specific endonuclease; single-strand-specific endodeoxyribonuclease; single strand-specific DNase; *Aspergillus oryzae* S1 nuclease

**References:** [55, 2809, 3076]

[EC 3.1.30.1 created 1972 as EC 3.1.4.21, transferred 1978 to EC 3.1.30.1, modified 1981]

#### EC 3.1.30.2

Accepted name: Serratia marcescens nuclease

**Reaction:** Endonucleolytic cleavage to 5'-phosphomononucleotide and 5'-phosphooligonucleotide end-products

**Other name(s):** endonuclease (*Serratia marcescens*); barley nuclease; plant nuclease I; nucleate endonuclease

**Comments:** Hydrolyses double- or single-stranded substrate.

**References:** [1909, 2767, 2768, 3145]

 $[EC\ 3.1.30.2\ created\ 1965\ as\ EC\ 3.1.4.9,\ transferred\ 1978\ to\ EC\ 3.1.30.2,\ modified\ 1981]$ 

# EC 3.1.31 Endoribonucleases that are active with either ribo- or deoxyribonucleic acids and produce 3'-phosphomonoesters

#### EC 3.1.31.1

Accepted name: micrococcal nuclease

**Reaction:** Endonucleolytic cleavage to nucleoside 3'-phosphates and 3'-phosphooligonucleotide end-products **Other name(s):** spleen endonuclease; thermonuclease; nuclease T; micrococcal endonuclease; nuclease T'; staphylo-

coccal nuclease; spleen phosphodiesterase; Staphylococcus aureus nuclease; Staphylococcus aureus

nuclease B; ribonucleate (deoxynucleate) 3'-nucleotidohydrolase

**Comments:** Hydrolyses double- or single-stranded substrate.

**References:** [31, 57, 2393, 2798]

# EC 3.2 Glycosylases

This subclass contains the glycosylases, which are classified as hydrolases, although some of them can also transfer glycosyl residues to oligosaccharides, polysaccharides and other alcoholic acceptors. The glycosylases are subdivided into glycosidases, i.e., enzymes that hydrolyse *O*- and *S*-glycosyl compounds (EC 3.2.1) and those that hydrolyse *N*-glycosyl compounds (EC 3.2.2). Common names for enzymes acting on D-sugars or their derivatives do not normally contain 'D', unless ambiguity would result from the common existence of the corresponding L-sugar. Enzymes that hydrolyse a terminal, non-reducing-end glycose (or a well-defined di-, tri- or oligosaccharide) from a glycan, i.e. exoenzymes, are given systematic names based on 'glycohydrolase'; enzymes that hydrolyse internal glycosidic bonds, i.e. endoenzymes, are given systematic names based on 'glycanohydrolase'. The same structure is often used when providing accepted names for these enzymes.

### EC 3.2.1 Glycosidases, i.e. enzymes that hydrolyse O- and S-glycosyl compounds

EC 3.2.1.1

**Accepted name:** α-amylase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkages in polysaccharides containing three or more

 $(1\rightarrow 4)$ - $\alpha$ -linked D-glucose units

**Other name(s):** glycogenase;  $\alpha$  amylase,  $\alpha$ -amylase; endoamylase; Taka-amylase A; 1,4- $\alpha$ -D-glucan glucanohydro-

lase

**Systematic name:** 4-α-D-glucan glucanohydrolase

Comments: Acts on starch, glycogen and related polysaccharides and oligosaccharides in a random manner; re-

ducing groups are liberated in the  $\alpha$ -configuration. The term " $\alpha$ " relates to the initial anomeric config-

uration of the free sugar group released and not to the configuration of the linkage hydrolysed.

**References:** [771, 1800, 2598]

[EC 3.2.1.1 created 1961]

EC 3.2.1.2

**Accepted name:** β-amylase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkages in polysaccharides so as to remove successive maltose

units from the non-reducing ends of the chains

**Other name(s):** saccharogen amylase; glycogenase;  $\beta$  amylase,  $\beta$ -amylase; 1,4- $\alpha$ -D-glucan maltohydrolase

**Systematic name:** 4-α-D-glucan maltohydrolase

**Comments:** Acts on starch, glycogen and related polysaccharides and oligosaccharides producing  $\beta$ -maltose by an

inversion. The term 'β" relates to the initial anomeric configuration of the free sugar group released

and not to the configuration of the linkage hydrolysed.

**References:** [129, 807, 1800]

[EC 3.2.1.2 created 1961]

EC 3.2.1.3

**Accepted name:** glucan 1,4-α-glucosidase

**Reaction:** Hydrolysis of terminal  $(1\rightarrow 4)$ -linked  $\alpha$ -D-glucose residues successively from non-reducing ends of

the chains with release of β-D-glucose

Other name(s): glucoamylase; amyloglucosidase;  $\gamma$ -amylase; lysosomal  $\alpha$ -glucosidase; acid maltase; exo-1,4- $\alpha$ -

glucosidase; glucose amylase; γ-1,4-glucan glucohydrolase; acid maltase; 1,4-α-D-glucan glucohy-

drolase

**Systematic name:** 4-α-D-glucan glucohydrolase

**Comments:** Most forms of the enzyme can rapidly hydrolyse 1,6-α-D-glucosidic bonds when the next bond in the

sequence is 1,4, and some preparations of this enzyme hydrolyse 1,6- and 1,3- $\alpha$ -D-glucosidic bonds in other polysaccharides. This entry covers all such enzymes acting on polysaccharides more rapidly than on oligosaccharides. EC 3.2.1.20  $\alpha$ -glucosidase, from mammalian intestine, can catalyse similar

reactions.

**References:** [808, 307, 1314, 1417, 1916, 2980]

[EC 3.2.1.3 created 1961]

EC 3.2.1.4

**Accepted name:** cellulase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-glucosidic linkages in cellulose, lichenin and cereal  $\beta$ -D-glucans **Other name(s):** endo-1,4- $\beta$ -D-glucanase;  $\beta$ -1,4-endoglucan hydrolase; cellulose A; cellulosin AP;

endoglucanase D; alkali cellulase; cellulase A 3; celludextrinase; 9.5 cellulase; avicelase; pancellase

SS;  $1,4-(1,3;1,4)-\beta$ -D-glucan 4-glucanohydrolase

**Systematic name:** 4-β-D-glucan 4-glucanohydrolase

**Comments:** Will also hydrolyse 1,4-linkages in β-D-glucans also containing 1,3-linkages.

**References:** [540, 1603, 2015, 2096, 3169, 1077, 1078, 1252]

[EC 3.2.1.4 created 1961, modified 2001]

[3.2.1.5 Deleted entry. licheninase]

[EC 3.2.1.5 created 1961, deleted 1964]

EC 3.2.1.6

**Accepted name:** endo-1,3(4)-β-glucanase

**Reaction:** Endohydrolysis of  $(1\rightarrow 3)$ - or  $(1\rightarrow 4)$ -linkages in  $\beta$ -D-glucans when the glucose residue whose reduc-

ing group is involved in the linkage to be hydrolysed is itself substituted at C-3

Other name(s): endo-1,3-β-D-glucanase; laminarinase; laminaranase; β-1,3-glucanase; β-1,3-1,4-glucanase; endo-1,3-

 $\beta$ -glucanase; endo- $\beta$ -1,3(4)-glucanase; endo- $\beta$ -1,3-1,4-glucanase; endo- $\beta$ -(1 $\rightarrow$ 3)-D-glucanase; endo-1,3-1,4- $\beta$ -D-glucanase; endo- $\beta$ -1,3-glucanase; endo- $\beta$ -1,3-gluca

 $1,3-(1,3;1,4)-\beta$ -D-glucan 3(4)-glucanohydrolase

**Systematic name:** 3(or 4)-β-D-glucan 3(4)-glucanohydrolase

Comments: Substrates include laminarin, lichenin and cereal D-glucans; different from EC 3.2.1.52 β-N-

acetylhexosaminidase.

**References:** [152, 153, 519, 2397, 2734]

[EC 3.2.1.6 created 1961, modified 1976]

EC 3.2.1.7

Accepted name: inulinase

**Reaction:** Endohydrolysis of  $(2\rightarrow 1)$ - $\beta$ -D-fructosidic linkages in inulin

Other name(s): inulase; indoinulinase; endo-inulinase; exoinulinase; 2,1-β-D-fructan fructanohydrolase

**Systematic name:** 1-β-D-fructan fructanohydrolase

**References:** [16]

[EC 3.2.1.7 created 1961]

EC 3.2.1.8

**Accepted name:** endo-1,4-β-xylanase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-xylosidic linkages in xylans

Other name(s): endo- $(1\rightarrow 4)$ - $\beta$ -xylan 4-xylanohydrolase; endo-1,4-xylanase; xylanase;  $\beta$ -1,4-xylanase; endo-1,4-

xylanase; endo- $\beta$ -1,4-xylanase; endo-1,4- $\beta$ -D-xylanase; 1,4- $\beta$ -xylan xylanohydrolase;  $\beta$ -xylanase;

 $\beta$ -1,4-xylan xylanohydrolase; endo-1,4- $\beta$ -xylanase;  $\beta$ -D-xylanase

**Systematic name:**  $4-\beta$ -D-xylan xylanohydrolase

**References:** [1192, 3168]

[EC 3.2.1.8 created 1961]

[3.2.1.9 Deleted entry. amylopectin-1,6-glucosidase]

[EC 3.2.1.9 created 1961, deleted 1972]

EC 3.2.1.10

Accepted name: oligo-1,6-glucosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic linkages in some oligosaccharides produced from starch and

glycogen by EC 3.2.1.1 (α-amylase), and in isomaltose

Other name(s): limit dextrinase (erroneous); isomaltase; sucrase-isomaltase; exo-oligo-1,6-glucosidase; dextrin

 $6\alpha$ -glucanohydrolase;  $\alpha$ -limit dextrinase; dextrin 6-glucanohydrolase; oligosaccharide  $\alpha$ -1,6-

glucohydrolase;  $\alpha$ -methylglucosidase

**Systematic name:** oligosaccharide  $6-\alpha$ -glucohydrolase

**Comments:** This enzyme, like EC 3.2.1.33 (amylo- $\alpha$ -1,6-glucosidase), can release an  $\alpha$ -1 $\rightarrow$ 6-linked glucose,

whereas the shortest chain that can be released by EC 3.2.1.41 (pullulanase), EC 3.2.1.142 (limit dextrinase), and EC 3.2.1.68 (isoamylase) is maltose. It also hydrolyses isomaltulose (palatinose), isomaltotriose and panose, but has no action on glycogen or phosphorylase limit dextrin. The enzyme from intestinal mucosa is a single polypeptide chain that also catalyses the reaction of EC 3.2.1.48 (sucrose  $\alpha$ -glucosidase). Differs from EC 3.2.1.33 (amylo- $\alpha$ -1,6-glucosidase) in its preference for short-chain substrates and in its not requiring the 6-glucosylated residue to be at a branch point, i.e. linked at both

C-1 and C-4.

**References:** [1080, 2678, 2449, 1433, 3248]

[EC 3.2.1.10 created 1961, modified 2000, modified 2013]

EC 3.2.1.11

Accepted name: dextranase

**Reaction:** Endohydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic linkages in dextran

Other name(s): dextran hydrolase; endodextranase; dextranase DL 2; DL 2; endo-dextranase; α-D-1,6-glucan-6-

glucanohydrolase; 1,6-α-D-glucan 6-glucanohydrolase

**Systematic name:** 6-α-D-glucan 6-glucanohydrolase

**References:** [121, 593, 772, 2474]

[EC 3.2.1.11 created 1961]

[3.2.1.12 Deleted entry. cycloheptaglucanase. Now included with EC 3.2.1.54 cyclomaltodextrinase]

[EC 3.2.1.12 created 1961, deleted 1976]

[3.2.1.13 Deleted entry. cyclohexaglucanase. Now included with EC 3.2.1.54 cyclomaltodextrinase]

[EC 3.2.1.13 created 1961, deleted 1976]

EC 3.2.1.14

Accepted name: chitinase

**Reaction:** Random endo-hydrolysis of *N*-acetyl- $\beta$ -D-glucosaminide (1 $\rightarrow$ 4)- $\beta$ -linkages in chitin and chitodex-

trins

Other name(s): ChiC; chitodextrinase (ambiguous); 1,4-β-poly-N-acetylglucosaminidase; poly-β-glucosaminidase;

β-1,4-poly-*N*-acetyl glucosamidinase; poly[1,4-(*N*-acetyl-β-D-glucosaminide)] glycanohydrolase

**Systematic name:**  $(1\rightarrow 4)$ -2-acetamido-2-deoxy- $\beta$ -D-glucan glycanohydrolase

**Comments:** The enzyme binds to chitin and randomly cleaves glycosidic linkages in chitin and chitodextrins in

a non-processive mode, generating chitooligosaccharides and free ends on which exo-chitinases and exo-chitodextrinases can act. Activity is greatly stimulated in the presence of EC 1.14.99.53, lytic chitin monoxygenase, which attacks the crystalline structure of chitin and makes the polymer more

accesible to the chitinase. cf. EC 3.2.1.202, endo-chitodextrinase.

**References:** [3319, 2958, 772, 477, 802, 3352, 2465]

[EC 3.2.1.14 created 1961, modified 2017]

EC 3.2.1.15

Accepted name: polygalacturonase

**Reaction:** Random hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-galactosiduronic linkages in pectate and other galacturonans pectin depolymerase; pectinase; endopolygalacturonase; pectin hydrolase; pectin poly-

 $galacturonase;\ endo-polygalacturonase;\ poly-\alpha-1,4-galacturonide\ glycanohydrolase;\ endogalacturonase$ 

onase; endo-D-galacturonase; poly(1,4- $\alpha$ -D-galacturonide) glycanohydrolase

**Systematic name:**  $(1\rightarrow 4)$ - $\alpha$ -D-galacturonan glycanohydrolase

**References:** [593, 1704, 1864, 1911, 2272]

[EC 3.2.1.15 created 1961]

[3.2.1.16 Deleted entry. alginase]

[EC 3.2.1.16 created 1961, deleted 1972]

EC 3.2.1.17

Accepted name: lysozyme

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -linkages between *N*-acetylmuramic acid and *N*-acetyl-D-glucosamine residues

in a peptidoglycan and between N-acetyl-D-glucosamine residues in chitodextrins

Other name(s): muramidase; globulin G; mucopeptide glucohydrolase; globulin G1; N,O-diacetylmuramidase;

lysozyme g; L-7001; 1,4-N-acetylmuramidase; mucopeptide N-acetylmuramoylhydrolase; PR1-

lysozyme

**Systematic name:** peptidoglycan *N*-acetylmuramoylhydrolase

**Comments:** *cf.* also EC 3.2.1.14 chitinase.

**References:** [232, 234, 1334]

[EC 3.2.1.17 created 1961]

EC 3.2.1.18

**Accepted name:** exo-α-sialidase

**Reaction:** Hydrolysis of  $\alpha$ -(2 $\rightarrow$ 3)-,  $\alpha$ -(2 $\rightarrow$ 6)-,  $\alpha$ -(2 $\rightarrow$ 8)- glycosidic linkages of terminal sialic acid residues in

oligosaccharides, glycoproteins, glycolipids, colominic acid and synthetic substrates

Other name(s): neuraminidase; sialidase;  $\alpha$ -neuraminidase; acetylneuraminidase

Systematic name: acetylneuraminyl hydrolase

Comments: The enzyme does not act on 4-O-acetylated sialic acids. endo- $\alpha$ -Sialidase activity is listed as EC

3.2.1.129, endo- $\alpha$ -sialidase. See also EC 4.2.2.15 anhydrosialidase.

**References:** [2556, 347]

[EC 3.2.1.18 created 1961, modified 1999]

[3.2.1.19 Deleted entry. heparinase]

[EC 3.2.1.19 created 1961, deleted 1978]

EC 3.2.1.20

**Accepted name:** α-glucosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $(1\rightarrow 4)$ -linked  $\alpha$ -D-glucose residues with release of D-glucose **Other name(s):** maltase; glucosidosucrase; maltase-glucoamylase;  $\alpha$ -glucopyranosidase; glucosidoin-

vertase; α-D-glucosidase; α-glucoside hydrolase; α-1,4-glucosidase

**Systematic name:**  $\alpha$ -D-glucoside glucohydrolase

**Comments:** This single entry covers a group of enzymes whose specificity is directed mainly towards the exo-

hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -glucosidic linkages, and that hydrolyse oligosaccharides rapidly, relative to polysaccharide, which are hydrolysed relatively slowly, or not at all. The intestinal enzyme also hydrolyses polysaccharides, catalysing the reactions of EC 3.2.1.3 glucan 1,4- $\alpha$ -glucosidase and, more

slowly, hydrolyses  $(1\rightarrow 6)$ - $\alpha$ -D-glucose links.

**References:** [319, 781, 1603, 2675, 2726]

[EC 3.2.1.20 created 1961]

EC 3.2.1.21

**Accepted name:** β-glucosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $\beta$ -D-glucosyl residues with release of  $\beta$ -D-glucose

Other name(s): gentiobiase; cellobiase; emulsin; elaterase; aryl- $\beta$ -glucosidase;  $\beta$ -D-glucosidase;  $\beta$ -glucoside glucohy-

drolase; arbutinase; amygdalinase; *p*-nitrophenyl β-glucosidase; primeverosidase; amygdalase; lina-

marase; salicilinase; β-1,6-glucosidase

**Systematic name:**  $\beta$ -D-glucoside glucohydrolase

**Comments:** Wide specificity for  $\beta$ -D-glucosides. Some examples also hydrolyse one or more of the following:

β-D-galactosides, α-L-arabinosides, β-D-xylosides, β-D-fucosides.

**References:** [435, 471, 529, 1135, 1603, 2520]

[EC 3.2.1.21 created 1961]

EC 3.2.1.22

**Accepted name:** α-galactosidase

Reaction: Hydrolysis of terminal, non-reducing  $\alpha$ -D-galactose residues in  $\alpha$ -D-galactosides, including galactose

oligosaccharides, galactomannans and galactolipids

**Other name(s):** melibiase;  $\alpha$ -D-galactosidase;  $\alpha$ -galactosidase A;  $\alpha$ -galactoside galactohydrolase

**Systematic name:**  $\alpha$ -D-galactoside galactohydrolase **Comments:** Also hydrolyses  $\alpha$ -D-fucosides.

**References:** [2811, 3181]

[EC 3.2.1.22 created 1961]

EC 3.2.1.23

**Accepted name:**  $\beta$ -galactosidase

**Reaction:** Hydrolysis of terminal non-reducing  $\beta$ -D-galactose residues in  $\beta$ -D-galactosides

Other name(s): lactase (ambiguous); β-lactosidase; maxilact; hydrolact; β-D-lactosidase; S 2107; lactozym; trilactase;

 $\beta$ -D-galactanase; oryzatym; sumiklat

**Systematic name:**  $\beta$ -D-galactoside galactohydrolase

**Comments:** Some enzymes in this group hydrolyse  $\alpha$ -L-arabinosides; some animal enzymes also hydrolyse  $\beta$ -D-

fucosides and  $\beta$ -D-glucosides; *cf.* EC 3.2.1.108 lactase.

**References:** [235, 1542, 1559, 1596, 1729, 1953, 3099, 89]

[EC 3.2.1.23 created 1961, modified 1980]

EC 3.2.1.24

**Accepted name:** α-mannosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $\alpha$ -D-mannose residues in  $\alpha$ -D-mannosides

Other name(s):  $\alpha$ -D-mannosidase; p-nitrophenyl- $\alpha$ -mannosidase;  $\alpha$ -D-mannopyranosidase; 1,2- $\alpha$ -mannosidase; 1,2- $\alpha$ -mannosidase

α-D-mannosidase; exo-α-mannosidase

**Systematic name:** α-D-mannoside mannohydrolase

Comments: Also hydrolyses  $\alpha$ -D-lyxosides and heptopyranosides with the same configuration at C-2, C-3 and C-4

as mannose.

**References:** [1675, 3194]

[EC 3.2.1.24 created 1961]

EC 3.2.1.25

**Accepted name:** β-mannosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $\beta$ -D-mannose residues in  $\beta$ -D-mannosides

**Other name(s):** mannanase; mannase; β-D-mannosidase; β-mannoside mannohydrolase; exo-β-D-mannanase

**Systematic name:** β-D-mannoside mannohydrolase

**References:** [16, 165, 592, 1215]

[EC 3.2.1.25 created 1961]

EC 3.2.1.26

**Accepted name:** β-fructofuranosidase

**Reaction:** Hydrolysis of terminal non-reducing  $\beta$ -D-fructofuranoside residues in  $\beta$ -D-fructofuranosides **Other name(s):** invertase; saccharase; glucosucrase;  $\beta$ -h-fructosidase;  $\beta$ -fructosidase; invertin; sucrase; maxinvert L

1000; fructosylinvertase; alkaline invertase; acid invertase

**Systematic name:** β-D-fructofuranoside fructohydrolase

**Comments:** Substrates include sucrose; also catalyses fructotransferase reactions.

**References:** [2017, 2071]

[EC 3.2.1.26 created 1961]

[3.2.1.27 Deleted entry.  $\alpha$ -1,3-glucosidase]

[EC 3.2.1.27 created 1961, deleted 1972]

EC 3.2.1.28

**Accepted name:**  $\alpha, \alpha$ -trehalase

**Reaction:**  $\alpha, \alpha$ -trehalose + H<sub>2</sub>O =  $\beta$ -D-glucose +  $\alpha$ -D-glucose

**Other name(s):** trehalase

**Systematic name:**  $\alpha, \alpha$ -trehalose glucohydrolase

Comments: The enzyme is an anomer-inverting glucosidase that catalyses the hydrolysis of the  $\alpha$ -glucosidic O-

linkage of  $\alpha$ , $\alpha$ -trehalose, releasing initially equimolar amounts of  $\alpha$ - and  $\beta$ -D-glucose. It is widely

distributed in microorganisms, plants, invertebrates and vertebrates.

**References:** [2018, 1363, 1099, 1965]

[EC 3.2.1.28 created 1961, modified 2012]

[3.2.1.29 Deleted entry. chitobiase. Now included with EC 3.2.1.52,  $\beta$ -N-acetylhexosaminidase]

[EC 3.2.1.29 created 1961, deleted 1972]

[3.2.1.30 Deleted entry. \( \beta\)-D-acetylglucosaminidase. Now included with EC 3.2.1.52, \( \beta\)-N-acetylhexosaminidase]

[EC 3.2.1.30 created 1961, deleted 1992]

EC 3.2.1.31

**Accepted name:** β-glucuronidase

**Reaction:** a  $\beta$ -D-glucuronoside + H<sub>2</sub>O = D-glucuronate + an alcohol

Other name(s): β-glucuronide glucuronohydrolase glucuronidase; exo-β-D-glucuronidase; ketodase

**Systematic name:**  $\beta$ -D-glucuronoside glucuronosohydrolase

**References:** [612, 644, 777, 1657, 3085]

[EC 3.2.1.31 created 1961]

EC 3.2.1.32

**Accepted name:** *endo-*1,3-β-xylanase

**Reaction:** Random endohydrolysis of  $(1\rightarrow 3)$ - $\beta$ -D-glycosidic linkages in  $(1\rightarrow 3)$ - $\beta$ -D-xylans

Other name(s): xylanase (ambiguous); endo-1,3-β-xylosidase (misleading); 1,3-β-xylanase; 1,3-xylanase; β-1,3-

xylanase; *endo*-β-1,3-xylanase; 1,3-β-D-xylan xylanohydrolase; xylan *endo*-1,3-β-xylosidase

**Systematic name:**  $3-\beta$ -D-xylan xylanohydrolase

Comments: This enzyme is found mostly in marine bacteria, which break down the  $\beta(1,3)$ -xylan found in the cell

wall of some green and red algae. The enzyme produces mainly xylobiose, xylotriose and xylote-

traose.

**References:** [417, 61, 68, 66, 2186]

[EC 3.2.1.32 created 1965, modified 2011]

EC 3.2.1.33

**Accepted name:** amylo-α-1,6-glucosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic branch linkages in glycogen phosphorylase limit dextrin **Other name(s):** amylo-1,6-glucosidase; dextrin  $6-\alpha$ -D-glucosidase; amylopectin 1,6-glucosidase; dextrin-1,6-

glucosidase; glycogen phosphorylase-limit dextrin α-1,6-glucohydrolase

**Systematic name:** glycogen phosphorylase-limit dextrin 6-α-glucohydrolase

**Comments:** This enzyme hydrolyses an unsubstituted glucose unit linked by an  $\alpha(1\rightarrow 6)$  bond to an  $\alpha(1\rightarrow 4)$  glu-

cose chain. The enzyme activity found in mammals and yeast is in a polypeptide chain containing two active centres. The other activity is similar to that of EC 2.4.1.25 (4- $\alpha$ -glucanotransferase), which acts on the glycogen phosphorylase limit dextrin chains to expose the single glucose residues, which the 6- $\alpha$ -glucosidase activity can then hydrolyse. Together, these two activities constitute the glycogen

debranching system.

**References:** [308, 1621, 2068]

[EC 3.2.1.33 created 1965, modified 2000]

[3.2.1.34 Deleted entry. chondroitinase. Now included with EC 3.2.1.35 hyalurononglucosaminidase]

[EC 3.2.1.34 created 1965, deleted 1972]

EC 3.2.1.35

Accepted name: hyaluronoglucosaminidase

**Reaction:** Random hydrolysis of  $(1\rightarrow 4)$ -linkages between *N*-acetyl- $\beta$ -D-glucosamine and D-glucuronate

residues in hyaluronate

**Other name(s):** hyaluronidase; hyaluronoglucosidase; chondroitinase; chondroitinase I

**Systematic name:** hyaluronate 4-glycanohydrolase

**Comments:** Also hydrolyses 1,4-β-D-glycosidic linkages between *N*-acetyl-galactosamine or *N*-

acetylgalactosamine sulfate and glucuronic acid in chondroitin, chondroitin 4- and 6-sulfates, and

dermatan.

**References:** [1896, 2375, 3155]

[EC 3.2.1.35 created 1965, modified 1976, modified 2001 (EC 3.2.1.34 created 1965, incorporated 1972)]

EC 3.2.1.36

Accepted name: hyaluronoglucuronidase

**Reaction:** Random hydrolysis of  $(1\rightarrow 3)$ -linkages between  $\beta$ -D-glucuronate and N-acetyl-D-glucosamine

residues in hyaluronate

**Other name(s):** hyaluronidase; glucuronoglucosaminoglycan hyaluronate lyase; orgelase

**Systematic name:** hyaluronate 3-glycanohydrolase

**References:** [1707, 1896]

[EC 3.2.1.36 created 1965, modified 1980]

#### EC 3.2.1.37

**Accepted name:** xylan 1,4-β-xylosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-xylans, to remove successive D-xylose residues from the non-reducing ter-

mini

**Other name(s):** xylobiase;  $\beta$ -xylosidase; exo-1,4- $\beta$ -xylosidase;  $\beta$ -D-xylopyranosidase;  $\beta$ -xylosidase;

exo-1,4-xylosidase; exo-1,4-β-D-xylosidase; 1,4-β-D-xylan xylohydrolase

**Systematic name:** 4-β-D-xylan xylohydrolase

Comments: Also hydrolyses xylobiose. Some other exoglycosidase activities have been found associated with this

enzyme in sheep liver.

**References:** [435, 1192]

[EC 3.2.1.37 created 1965]

#### EC 3.2.1.38

**Accepted name:** β-D-fucosidase

**Reaction:** Hydrolysis of terminal non-reducing  $\beta$ -D-fucose residues in  $\beta$ -D-fucosides

Other name(s):  $\beta$ -fucosidase

**Systematic name:** β-D-fucoside fucohydrolase

**Comments:** Enzymes from some sources also hydrolyse  $\beta$ -D-galactosides and/or  $\beta$ -D-glucosides and/or  $\alpha$ -L-

arabinosides. The activity of EC 3.2.1.37 xylan 1,4- $\beta$ -xylosidase, is an associated activity found in

some sources (e.g. liver).

**References:** [434, 435, 2450, 3182, 3183]

[EC 3.2.1.38 created 1965, deleted 1972, reinstated 1978]

#### EC 3.2.1.39

**Accepted name:** glucan endo-1,3-β-D-glucosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 3)$ - $\beta$ -D-glucosidic linkages in  $(1\rightarrow 3)$ - $\beta$ -D-glucans

 $\textbf{Other name(s):} \quad \text{endo-1,3-$\beta$-glucanase; laminarinase; oligo-1,3-glucosidase; endo-1,3-$\beta$-glucanase;}$ 

callase;  $\beta$ -1,3-glucanase; kitalase; 1,3- $\beta$ -D-glucan 3-glucanohydrolase; endo-(1,3)- $\beta$ -D-glucanase; (1 $\rightarrow$ 3)- $\beta$ -glucan 3-glucanohydrolase; endo-1,3- $\beta$ -D-glucanase; endo-1,3- $\beta$ -glucosidase; 1,3- $\beta$ -D-glucanase; 1,

glucan glucanohydrolase

**Systematic name:**  $3-\beta$ -D-glucan glucanohydrolase

**Comments:** Different from EC 3.2.1.6 endo-1,3(4)- $\beta$ -glucanase. Very limited action on mixed-link (1 $\rightarrow$ 3,1 $\rightarrow$ 4)- $\beta$ -

D-glucans. Hydrolyses laminarin, paramylon and pachyman.

**References:** [428, 2397]

[EC 3.2.1.39 created 1965]

#### EC 3.2.1.40

**Accepted name:** α-L-rhamnosidase

**Reaction:** Hydrolysis of terminal non-reducing  $\alpha$ -L-rhamnose residues in  $\alpha$ -L-rhamnosides

Other name(s): α-L-rhamnosidase T; α-L-rhamnosidase N

**Systematic name:** α-L-rhamnoside rhamnohydrolase

Comments: The enzyme, found in animal tissues, plants, yeasts, fungi and bacteria, utilizes an inverting mecha-

nism of hydrolysis, releasing  $\beta$ -L-rhamnose. Substrates include naringin, rutin, quercitrin, hesperidin, dioscin, terpenyl glycosides and many other natural glycosides containing terminal  $\alpha$ -L-rhamnose.

**References:** [2461, 1575, 3353, 3255, 514, 2346]

[EC 3.2.1.40 created 1972]

#### EC 3.2.1.41

Accepted name: pullulanase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic linkages in pullulan, amylopectin and glycogen, and in the  $\alpha$ -

and β-limit dextrins of amylopectin and glycogen

Other name(s): limit dextrinase (erroneous); amylopectin 6-glucanohydrolase; bacterial debranching enzyme; de-

branching enzyme;  $\alpha$ -dextrin endo-1,6- $\alpha$ -glucosidase; R-enzyme; pullulan  $\alpha$ -1,6-glucanohydrolase

**Systematic name:** pullulan 6-α-glucanohydrolase

**Comments:** Different from EC 3.2.1.142 (limit dextrinase) in its action on glycogen, and its rate of hydrolysis of

limit dextrins. Its action on amylopectin is complete. Maltose is the smallest sugar that it can release

from an  $\alpha$ -(1 $\rightarrow$ 6)-linkage.

**References:** [1622, 192, 1801]

[EC 3.2.1.41 created 1972, modified 1976, modified 2000 (EC 3.2.1.69 created 1972, incorporated 1976)]

#### EC 3.2.1.42

Accepted name: GDP-glucosidase

**Reaction:** GDP-glucose +  $H_2O$  = D-glucose + GDP

Other name(s): guanosine diphosphoglucosidase; guanosine diphosphate D-glucose glucohydrolase

Systematic name: GDP-glucose glucohydrolase

**References:** [2723]

[EC 3.2.1.42 created 1972]

#### EC 3.2.1.43

**Accepted name:**  $\beta$ -L-rhamnosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $\beta$ -L-rhamnose residues in  $\beta$ -L-rhamnosides

**Systematic name:** β-L-rhamnoside rhamnohydrolase

**References:** [142]

[EC 3.2.1.43 created 1972]

#### EC 3.2.1.44

Accepted name: fucoidanase

**Reaction:** Endohydrolysis of  $(1\rightarrow 2)$ - $\alpha$ -L-fucoside linkages in fucoidan without release of sulfate

Other name(s):  $\alpha$ -L-fucosidase; poly(1,2- $\alpha$ -L-fucoside-4-sulfate) glycanohydrolase

**Systematic name:** poly[ $(1\rightarrow 2)$ - $\alpha$ -L-fucoside-4-sulfate] glycanohydrolase

**References:** [2906]

[EC 3.2.1.44 created 1972]

# EC 3.2.1.45

Accepted name: glucosylceramidase

**Reaction:** a D-glucosyl-N-acylsphingosine +  $H_2O$  = D-glucose + a ceramide

Other name(s): psychosine hydrolase; glucosphingosine glucosylhydrolase; GlcCer-β-glucosidase; β-D-

glucocerebrosidase; glucosylcerebrosidase;  $\beta$ -glucosylceramidase; ceramide glucosidase; glucocere-

brosidase; glucosylsphingosine β-glucosidase; glucosylsphingosine β-D-glucosidase

**Systematic name:** D-glucosyl-*N*-acylsphingosine glucohydrolase

**Comments:** Also acts on glucosylsphingosine (*cf.* EC 3.2.1.62 glycosylceramidase).

**References:** [275, 3025]

[EC 3.2.1.45 created 1972]

EC 3.2.1.46

Accepted name: galactosylceramidase

**Reaction:** a D-galactosyl-N-acylsphingosine +  $H_2O$  = D-galactose + a ceramide

Other name(s): cerebroside galactosidase; galactocerebrosides, galactosidase; ga

cerebrosidase; ceramide galactosidase; galactoserebroside galactosidase; galactosylceramide.  $\beta$ -galactosidase; cerebroside  $\beta$ -galactosidase; galactosylceramidase I;  $\beta$ -galactosylceramidase; galactocerebroside- $\beta$ -D-galactosidase; lactosylceramidase I;  $\beta$ -galactocerebrosidase; lactosylceramidase

dase

**Systematic name:** D-galactosyl-*N*-acylsphingosine galactohydrolase

**Comments:** *cf.* EC 3.2.1.62 glycosylceramidase.

**References:** [274]

[EC 3.2.1.46 created 1972]

EC 3.2.1.47

Accepted name: galactosylgalactosylglucosylceramidase

**Reaction:**  $\alpha$ -D-galactosyl- $(1\rightarrow 4)$ - $\beta$ -D-galactosyl- $(1\rightarrow 4)$ - $\beta$ -D-glucosyl- $(1\leftrightarrow 1)$ -ceramide +  $H_2O$  = D-galactose +

 $\beta$ -D-galactosyl-(1 $\rightarrow$ 4)-β-D-glucosyl-(1 $\leftrightarrow$ 1)-ceramide

Other name(s): trihexosyl ceramide galactosidase; ceramide trihexosidase; ceramidetrihexoside  $\alpha$ -galactosidase; tri-

hexosylceramide  $\alpha$ -galactosidase; ceramidetrihexosidase

**Systematic name:**  $\alpha$ -D-galactosyl- $(1 \rightarrow 4)$ - $\beta$ -D-galactosyl- $(1 \rightarrow 4)$ - $\beta$ -D-glucosyl- $(1 \leftrightarrow 1)$ -ceramide galactohydrolase

**References:** [273, 1804]

[EC 3.2.1.47 created 1972, modified 2011]

EC 3.2.1.48

**Accepted name:** sucrose α-glucosidase

**Reaction:** Hydrolysis of sucrose and maltose by an  $\alpha$ -D-glucosidase-type action

Other name(s): sucrose  $\alpha$ -glucohydrolase; sucrase; sucrase; sucrose. $\alpha$ -glucohydrolase; intestinal sucrase;

sucrase(invertase)

**Systematic name:** sucrose-α-D-glucohydrolase

**Comments:** This enzyme is isolated from intestinal mucosa as a single polypeptide chain that also displays activ-

ity towards isomaltose (EC 3.2.1.10 oligo-1,6-glucosidase).

**References:** [473, 1080, 1499, 2657, 2678, 2860]

[EC 3.2.1.48 created 1972]

EC 3.2.1.49

**Accepted name:** α-N-acetylgalactosaminidase

**Reaction:** Cleavage of non-reducing  $\alpha$ - $(1\rightarrow 3)$ -N-acetylgalactosamine residues from human blood group A and

AB mucin glycoproteins, Forssman hapten and blood group A lacto series glycolipids

**Other name(s):**  $\alpha$ -acetylgalactosaminidase; N-acetyl- $\alpha$ -D-galactosaminidase; N-acetyl- $\alpha$ -galactosaminidase;  $\alpha$ -

NAGAL; α-NAGA; α-GalNAcase

**Systematic name:**  $\alpha$ -N-acetyl-D-galactosaminide N-acetylgalactosaminohydrolase

**Comments:** The human lysosomal enzyme is involved in the degradation of blood type A epitope.

**References:** [84, 3339, 458, 1185, 1064, 3151, 87]

#### [EC 3.2.1.49 created 1972, modified 2011]

EC 3.2.1.50

**Accepted name:** α-N-acetylglucosaminidase

**Reaction:** Hydrolysis of terminal non-reducing N-acetyl-D-glucosamine residues in N-acetyl- $\alpha$ -D-

glucosaminides

Other name(s):  $\alpha$ -acetylglucosaminidase; N-acetyl- $\alpha$ -D-glucosaminidase; N-acetyl- $\alpha$ -glucosaminidase;  $\alpha$ -D-2-

acetamido-2-deoxyglucosidase

**Systematic name:**  $\alpha$ -N-acetyl-D-glucosaminide N-acetylglucosaminohydrolase

**Comments:** Hydrolyses UDP-*N*-acetylglucosamine.

**References:** [3078, 3079, 3157, 3163]

[EC 3.2.1.50 created 1972]

EC 3.2.1.51

**Accepted name:** α-L-fucosidase

**Reaction:** an  $\alpha$ -L-fucoside + H<sub>2</sub>O = L-fucose + an alcohol

Other name(s):  $\alpha$ -fucosidase

**Systematic name:**  $\alpha$ -L-fucoside fucohydrolase

**References:** [1658, 2400, 2870]

[EC 3.2.1.51 created 1972]

EC 3.2.1.52

**Accepted name:**  $\beta$ -*N*-acetylhexosaminidase

**Reaction:** Hydrolysis of terminal non-reducing N-acetyl-D-hexosamine residues in N-acetyl- $\beta$ -D-hexosaminides

Other name(s): hexosaminidase; β-acetylaminodeoxyhexosidase; N-acetyl-β-D-hexosaminidase; N-acetyl-β-

hexosaminidase;  $\beta$ -hexosaminidase;  $\beta$ -acetylhexosaminidase;  $\beta$ -D-N-acetylhexosaminidase;  $\beta$ -N-acetyl-D-hexosaminidase;  $\beta$ -N-acetylglucosaminidase; hexosaminidase A; N-acetylhexosaminidase;

 $\beta$ -D-hexosaminidase

**Systematic name:**  $\beta$ -*N*-acetyl-D-hexosaminide *N*-acetylhexosaminohydrolase

**Comments:** Acts on *N*-acetylglucosides and *N*-acetylgalactosides.

**References:** [346, 357, 823, 1667]

[EC 3.2.1.52 created 1972 (EC 3.2.1.30 created 1961, incorporated 1992 [EC 3.2.1.29 created 1961, incorporated 1972])]

EC 3.2.1.53

**Accepted name:** β-*N*-acetylgalactosaminidase

**Reaction:** Hydrolysis of terminal non-reducing N-acetyl-D-galactosamine residues in N-acetyl- $\beta$ -D-

galactosaminides

Other name(s): N-acetyl- $\beta$ -galactosaminidase; N-acetyl- $\beta$ -D-galactosaminidase;  $\beta$ -acetylgalactosaminidase;  $\beta$ -D-N-

 $acetyl galactosaminidase; {\it N-} acetyl galactosaminidase$ 

**Systematic name:**  $\beta$ -*N*-acetyl-D-galactosaminide *N*-acetylgalactosaminohydrolase

**References:** [823, 1170]

[EC 3.2.1.53 created 1972]

EC 3.2.1.54

Accepted name: cyclomaltodextrinase

**Reaction:** cyclomaltodextrin +  $H_2O$  = linear maltodextrin

Other name(s): cycloheptaglucanase; cyclohexaglucanase; cyclodextrinase; cyclomaltodextrin dextrin-hydrolase (de-

cyclizing)

**Systematic name:** cyclomaltodextrin dextrin-hydrolase (ring-opening)

**Comments:** Also hydrolyses linear maltodextrin.

**References:** [585]

[EC 3.2.1.54 created 1972 (EC 3.2.1.12 and EC 3.2.1.13 both created 1961 and incorporated 1976)]

#### EC 3.2.1.55

**Accepted name:** non-reducing end α-L-arabinofuranosidase

**Reaction:** Hydrolysis of terminal non-reducing  $\alpha$ -L-arabinofuranoside residues in  $\alpha$ -L-arabinosides.

Other name(s): arabinosidase (ambiguous);  $\alpha$ -arabinosidase;  $\alpha$ -L-arabinosidase;  $\alpha$ -arabinofuranosidase; polysaccha-

ride α-L-arabinofuranosidase; α-L-arabinofuranoside hydrolase; L-arabinosidase (ambiguous); α-L-

arabinanase

**Systematic name:**  $\alpha$ -L-arabinofuranoside non-reducing end  $\alpha$ -L-arabinofuranosidase

**Comments:** The enzyme acts on  $\alpha$ -L-arabinofuranosides,  $\alpha$ -L-arabinans containing (1,3)- and/or (1,5)-

linkages, arabinoxylans and arabinogalactans. Some  $\beta$ -galactosidases (EC 3.2.1.23) and  $\beta$ -D-fucosidases (EC 3.2.1.38) also hydrolyse  $\alpha$ -L-arabinosides. *cf.* EC 3.2.1.185, non-reducing end  $\beta$ -

L-arabinofuranosidase.

**References:** [2829, 1355, 1356, 1810, 1244]

[EC 3.2.1.55 created 1972, modified 1976 (EC 3.2.1.79 created 1972, incorporated 1976), modified 2013]

#### EC 3.2.1.56

Accepted name: glucuronosyl-disulfoglucosamine glucuronidase

**Reaction:** 3-D-glucuronosyl- $N^2$ ,6-disulfo- $\beta$ -D-glucosamine +  $H_2O$  = D-glucuronate +  $N^2$ ,6-disulfo-D-

glucosamine

Other name(s): glycuronidase; 3-D-glucuronsyl-2-*N*,6-disulfo-β-D-glucosamine glucuronohydrolase

**Systematic name:** 3-D-glucuronsyl- $N^2$ ,6-disulfo- $\beta$ -D-glucosamine glucuronohydrolase

**References:** [608]

[EC 3.2.1.56 created 1972]

## EC 3.2.1.57

Accepted name: isopullulanase

**Reaction:** Hydrolysis of pullulan to isopanose (6-α-maltosylglucose)

**Systematic name:** pullulan 4-glucanohydrolase (isopanose-forming)

**Comments:** The enzyme has practically no action on starch. Panose  $(4-\alpha$ -isomaltosylglucose) is hydrolysed to

isomaltose and glucose. cf. EC 3.2.1.41 (pullulanase) and EC 3.2.1.135 (neopullulanase).

**References:** [2504]

[EC 3.2.1.57 created 1972]

#### EC 3.2.1.58

**Accepted name:** glucan 1,3-β-glucosidase

**Reaction:** Successive hydrolysis of  $\beta$ -D-glucose units from the non-reducing ends of  $(1 \rightarrow 3)$ - $\beta$ -D-glucans, releas-

ing α-glucose

Other name(s): exo-1,3- $\beta$ -glucosidase;  $\beta$ -1,3-glucan exo-hydrolase; exo (1 $\rightarrow$ 3)-glucanohydrolase; 1,3- $\beta$ -glucan glu-

cohydrolase

**Systematic name:** 3-β-D-glucan glucohydrolase

Comments: Acts on oligosaccharides, but very slowly on laminaribiose.

**References:** [152, 153]

[EC 3.2.1.58 created 1972]

#### EC 3.2.1.59

**Accepted name:** glucan endo-1,3-α-glucosidase

**Reaction:** Endohydrolysis of  $(1\rightarrow 3)$ - $\alpha$ -D-glucosidic linkages in isolichenin, pseudonigeran and nigeran

Other name(s): endo-1,3- $\alpha$ -glucanase; mutanase; endo-(1 $\rightarrow$ 3)- $\alpha$ -glucanase; cariogenase; cariogenanase; endo-1,3- $\alpha$ -

D-glucanase; 1,3(1,3;1,4)-α-D-glucan 3-glucanohydrolase

**Systematic name:** 3-α-D-glucan 3-glucanohydrolase

**Comments:** Products from pseudonigeran (1,3- $\alpha$ -D-glucan) are nigerose and  $\alpha$ -D-glucose.

**References:** [1068]

[EC 3.2.1.59 created 1972]

#### EC 3.2.1.60

**Accepted name:** glucan 1,4-α-maltotetraohydrolase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkages in amylaceous polysaccharides, to remove successive

maltotetraose residues from the non-reducing chain ends

**Other name(s):** exo-maltotetraohydrolase; 1,4-α-D-glucan maltotetraohydrolase

**Systematic name:** 4-α-D-glucan maltotetraohydrolase

Comments: Compare EC 3.2.1.2 β-amylase, which removes successive maltose residues, and EC 3.2.1.98 (glucan

1,4-α-maltohexaosidase) and EC 3.2.1.116 (glucan 1,4-α-maltotriohydrolase).

**References:** [2043, 2443]

[EC 3.2.1.60 created 1972]

#### EC 3.2.1.61

Accepted name: mycodextranase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkages in  $\alpha$ -D-glucans containing both  $(1\rightarrow 3)$ - and

 $(1\rightarrow 4)$ -bonds

Other name(s): 1,3-1,4- $\alpha$ -D-glucan 4-glucanohydrolase Systematic name:  $(1\rightarrow 3)$ - $(1\rightarrow 4)$ - $\alpha$ -D-glucan 4-glucanohydrolase

Comments: Products are nigerose and 4- $\alpha$ -D-nigerosylglucose. No hydrolysis of  $\alpha$ -D-glucans containing only 1,3-

or 1,4-bonds.

**References:** [2997]

[EC 3.2.1.61 created 1972]

#### EC 3.2.1.62

Accepted name: glycosylceramidase

**Reaction:** a glycosyl-*N*-acylsphingosine +  $H_2O$  = a ceramide + a sugar

Other name(s): phlorizin hydrolase; phloretin-glucosidase; glycosyl ceramide glycosylhydrolase; cerebrosidase; phloretin-glucosidase; phl

ridzin β-glucosidase; lactase-phlorizin hydrolase; phloridzin glucosidase

**Systematic name:** glycosyl-*N*-acylsphingosine glycohydrolase

**Comments:** Broad specificity [cf. EC 3.2.1.45 (glucosylceramidase) and EC 3.2.1.46 (galactosylceramidase)].

Also hydrolyses phlorizin to phloretin and glucose. The intestinal enzyme is a complex that also

catalyses the reaction of EC 3.2.1.108 lactase.

**References:** [1639, 1736, 1786]

[EC 3.2.1.62 created 1972, modified 1976]

#### EC 3.2.1.63

**Accepted name:** 1,2-α-L-fucosidase

**Reaction:** methyl-2- $\alpha$ -L-fucopyranosyl- $\beta$ -D-galactoside + H<sub>2</sub>O = L-fucose + methyl  $\beta$ -D-galactoside

Other name(s): almond emulsin fucosidase;  $\alpha$ - $(1\rightarrow 2)$ -L-fucosidase Systematic name:  $2-\alpha$ -L-fucopyranosyl- $\beta$ -D-galactoside fucohydrolase Comments: Highly specific for non-reducing terminal L-fucose residues linked to D-galactose residues by a 1,2-α-

linkage. Not identical with EC 3.2.1.111 1,3-α-L-fucosidase.

**References:** [118, 2144, 2400]

[EC 3.2.1.63 created 1972]

EC 3.2.1.64

**Accepted name:** 2,6-β-fructan 6-levanbiohydrolase

**Reaction:** Hydrolysis of  $(2\rightarrow 6)$ - $\beta$ -D-fructofuranan, to remove successive disaccharide residues as levanbiose,

i.e. 6-(β-D-fructofuranosyl)-D-fructose, from the end of the chain

Other name(s): β-2,6-fructan-6-levanbiohydrolase; 2,6-β-D-fructan 6-levanbiohydrolase; levanbiose-producing lev-

anase; 2,6-β-D-fructan 6-β-D-fructofuranosylfructohydrolase

**Systematic name:**  $(2\rightarrow 6)$ - $\beta$ -D-fructofuranan 6- $(\beta$ -D-fructosyl)-D-fructose-hydrolase

**References:** [105, 2498, 2499, 2715, 1385]

[EC 3.2.1.64 created 1972, modified 2004]

EC 3.2.1.65

Accepted name: levanase

**Reaction:** Random hydrolysis of  $(2\rightarrow 6)$ - $\beta$ -D-fructofuranosidic linkages in  $(2\rightarrow 6)$ - $\beta$ -D-fructans (levans) contain-

ing more than 3 fructose units

**Other name(s):** levan hydrolase; 2,6-β-D-fructan fructanohydrolase

**Systematic name:**  $(2\rightarrow 6)$ - $\beta$ -D-fructan fructanohydrolase

**References:** [104]

[EC 3.2.1.65 created 1972]

EC 3.2.1.66

Accepted name: quercitrinase

**Reaction:** quercitrin +  $H_2O$  = L-rhamnose + quercetin

**Systematic name:** quercitrin 3-L-rhamnohydrolase **Comments:** Quercitrin is quercetin 3-L-rhamnoside.

**References:** [3164]

[EC 3.2.1.66 created 1972]

EC 3.2.1.67

**Accepted name:** galacturan 1,4-α-galacturonidase

**Reaction:**  $[(1 \rightarrow 4) - \alpha - D$ -galacturonide]<sub>n</sub> + H<sub>2</sub>O =  $[(1 \rightarrow 4) - \alpha - D$ -galacturonide]<sub>n-1</sub> + D-galacturonate

Other name(s): exopolygalacturonase; poly(galacturonate) hydrolase; exo-D-galacturonase; exo-D-galacturonase;

exopoly-D-galacturonase; poly(1,4- $\alpha$ -D-galacturonide) galacturonohydrolase

**Systematic name:** poly $[(1\rightarrow 4)-\alpha$ -D-galacturonide] galacturonohydrolase

References: [1067]

[EC 3.2.1.67 created 1972]

EC 3.2.1.68

Accepted name: isoamylase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic branch linkages in glycogen, amylopectin and their  $\beta$ -limit dex-

trins

**Other name(s):** debranching enzyme; glycogen α-1,6-glucanohydrolase

**Systematic name:** glycogen 6-α-D-glucanohydrolase

Comments: Also readily hydrolyses amylopectin. Differs from EC 3.2.1.41 (pullulanase) and EC 3.2.1.142 (limit

dextrinase) by its inability to hydrolyse pullulan, and by limited action on α-limit dextrins. Maltose is

the smallest sugar it can release from an  $\alpha$ - $(1\rightarrow 6)$ -linkage.

**References:** [3286]

[EC 3.2.1.68 created 1972, modified 1976, modified 2000]

[3.2.1.69 Deleted entry, amylopectin 6-glucanohydrolase. Now included with EC 3.2.1.41 pullulanase]

[EC 3.2.1.69 created 1972, deleted 1976]

EC 3.2.1.70

**Accepted name:** glucan 1,6-α-glucosidase

**Reaction:** Hydrolysis of  $(1 \rightarrow 6)$ - $\alpha$ -D-glucosidic linkages in  $(1 \rightarrow 6)$ - $\alpha$ -D-glucans and derived oligosaccharides

Other name(s): exo-1,6- $\beta$ -glucosidase; glucodextrinase; glucan  $\alpha$ -1,6-D-glucohydrolase

**Systematic name:** glucan 6-α-D-glucohydrolase

**Comments:** Hydrolysis is accompanied by inversion at C-1, so that new reducing ends are released in the β-

configuration. Dextrans and isomaltosaccharides are hydrolysed, as is isomaltose, but very slowly. The enzyme from some sources also possesses the activity of EC 3.2.1.59 (glucan endo-1,3- $\alpha$ -

glucosidase).

**References:** [2176, 2547, 3095]

[EC 3.2.1.70 created 1972, modified 2001]

EC 3.2.1.71

**Accepted name:** glucan endo-1,2-β-glucosidase

**Reaction:** Random hydrolysis of  $(1\rightarrow 2)$ -glucosidic linkages in  $(1\rightarrow 2)$ - $\beta$ -D-glucans

**Other name(s):** endo-1,2- $\beta$ -glucanase;  $\beta$ -D-1,2-glucanase; endo- $(1\rightarrow 2)$ - $\beta$ -D-glucanase; 1,2- $\beta$ -D-glucan glucanohydro-

lase

**Systematic name:** 2-β-D-glucan glucanohydrolase

**References:** [2398]

[EC 3.2.1.71 created 1972]

EC 3.2.1.72

**Accepted name:** xylan 1,3-β-xylosidase

**Reaction:** Hydrolysis of successive xylose residues from the non-reducing termini of  $(1 \rightarrow 3)$ -β-D-xylans **Other name(s):** 1,3-β-D-xylosidase, exo-1,3-β-xylosidase; β-1,3'-xylanase; exo-β-1,3'-xylanase; 1,3-β-D-xylan xylo-

hvdrolase

**Systematic name:** 3-β-D-xylan xylohydrolase

**References:** [859]

[EC 3.2.1.72 created 1972]

EC 3.2.1.73

Accepted name: licheninase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ -β-D-glucosidic linkages in β-D-glucans containing  $(1\rightarrow 3)$ - and  $(1\rightarrow 4)$ -bonds **Other name(s):** lichenase; β- $(1\rightarrow 4)$ -D-glucan 4-glucanohydrolase; 1,3;1,4-β-glucan endohydrolase; 1,3;1,4-β-glucan

4-glucanohydrolase; 1,3-1,4-β-D-glucan 4-glucanohydrolase

**Systematic name:**  $(1\rightarrow 3)$ - $(1\rightarrow 4)$ - $\beta$ -D-glucan 4-glucanohydrolase

**Comments:** Acts on lichenin and cereal  $\beta$ -D-glucans, but not on  $\beta$ -D-glucans containing only 1,3- or 1,4-bonds.

**References:** [151]

#### [EC 3.2.1.73 created 1972]

EC 3.2.1.74

**Accepted name:** glucan 1,4-β-glucosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ -linkages in  $(1\rightarrow 4)$ - $\beta$ -D-glucans, to remove successive glucose units

Other name(s):  $\exp(-1,4-\beta)$ -glucosidase;  $\exp(-\beta-1,4-\beta)$ -glucosidase;  $\exp(-\beta-1,4-\beta)$ -glucanase;  $\exp($ 

β-glucosidase; exo-1,4-β-glucanase; 1,4-β-D-glucan glucohydrolase

**Systematic name:** 4-β-D-glucan glucohydrolase

Comments: Acts on 1,4- $\beta$ -D-glucans and related oligosaccharides. Cellobiose is hydrolysed, but very slowly.

**References:** [151]

[EC 3.2.1.74 created 1972]

EC 3.2.1.75

Accepted name: glucan endo-1,6-β-glucosidase

**Reaction:** Random hydrolysis of  $(1\rightarrow 6)$ -linkages in  $(1\rightarrow 6)$ - $\beta$ -D-glucans

**Other name(s):** endo-1,6- $\beta$ -glucanase;  $\beta$ -1 $\rightarrow$ 6)- $\beta$ -D-glucanase;  $\beta$ -1,6-glucanase-pustulanase;  $\beta$ -1,6-glucan hydrolase;

β-1,6-glucan 6-glucanohydrolase; 1,6-β-D-glucan glucanohydrolase

**Systematic name:** 6-β-D-glucan glucanohydrolase

**Comments:** Acts on lutean, pustulan and 1,6-oligo-β-D-glucosides.

References: [2399]

[EC 3.2.1.75 created 1972]

EC 3.2.1.76

Accepted name: L-iduronidase

**Reaction:** Hydrolysis of unsulfated  $\alpha$ -L-iduronosidic linkages in dermatan sulfate

**Other name**(s):  $\alpha$ -L-iduronidase

**Systematic name:** glycosaminoglycan α-L-iduronohydrolase

**References:** [1834, 2454, 2750]

[EC 3.2.1.76 created 1972]

EC 3.2.1.77

**Accepted name:** mannan  $1,2-(1,3)-\alpha$ -mannosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 2)$ - and  $(1\rightarrow 3)$ -linkages in yeast mannan, releasing mannose

Other name(s):  $exo-1,2-1,3-\alpha$ -mannosidase;  $1,2-1,3-\alpha$ -D-mannan mannohydrolase

**Systematic name:**  $(1\rightarrow 2)$ - $(1\rightarrow 3)$ - $\alpha$ -D-mannan mannohydrolase

Comments: A 1,6- $\alpha$ -D-mannan backbone remains after action on yeast mannan. This is further attacked, but

slowly.

**References:** [1336, 1337]

[EC 3.2.1.77 created 1972]

EC 3.2.1.78

**Accepted name:** mannan endo-1,4-β-mannosidase

**Reaction:** Random hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-mannosidic linkages in mannans, galactomannans and glucoman-

nans

Other name(s): endo-1,4- $\beta$ -mannanase; endo- $\beta$ -1,4-mannase;  $\beta$ -mannanase B;  $\beta$ -1, 4-mannan 4-mannanohydrolase;

endo- $\beta$ -mannanase;  $\beta$ -D-mannanase; 1,4- $\beta$ -D-mannan mannanohydrolase

**Systematic name:**  $4-\beta$ -D-mannan mannanohydrolase

**References:** [712, 2396]

#### [EC 3.2.1.78 created 1972]

#### [3.2.1.79 Deleted entry. \( \alpha \)-L-arabinofuranoside hydrolase. Now included with EC 3.2.1.55 \( \alpha \)-N-arabinofuranosidase]

#### [EC 3.2.1.79 created 1972, deleted 1976]

#### EC 3.2.1.80

**Accepted name:** fructan  $\beta$ -fructosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $(2\rightarrow 1)$ - and  $(2\rightarrow 6)$ -linked  $\beta$ -D-fructofuranose residues in fruc-

tans

Other name(s):  $\exp-\beta$ -D-fructosidase;  $\exp-\beta$ -fructosidase; polysaccharide  $\beta$ -fructofuranosidase; fructan exohydrolase

**Systematic name:** β-D-fructan fructohydrolase

**Comments:** Hydrolyses inulin and levan, and also sucrose.

**References:** [526, 1292]

[EC 3.2.1.80 created 1972]

#### EC 3.2.1.81

**Accepted name:** β-agarase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-galactosidic linkages in agarose, giving the tetramer as the predominant

product

Other name(s): agarase (ambiguous); AgaA; AgaB; endo-β-agarase; agarose 3-glycanohydrolase (incorrect)

**Systematic name:** agarose 4-glycanohydrolase

**Comments:** Also acts on porphyran, but more slowly [659]. This enzyme cleaves the  $\beta$ -(1 $\rightarrow$ 4) linkages of agarose

in a random manner with retention of the anomeric-bond configuration, producing  $\beta$ -anomers that give rise progressively to  $\alpha$ -anomers when mutarotation takes place [1303]. The end products of hydrolysis are neoagarotetraose and neoagarohexaose in the case of AgaA from the marine bacterium Zobellia galactanivorans, and neoagarotetraose and neoagarobiose in the case of AgaB [1303].

**References:** [659, 37, 2172, 2171, 2792, 1303]

[EC 3.2.1.81 created 1972, modified 2006]

#### EC 3.2.1.82

**Accepted name:** exo-poly-α-galacturonosidase

**Reaction:** Hydrolysis of pectic acid from the non-reducing end, releasing digalacturonate

Other name(s): exopolygalacturonosidase; exopolygalacturanosidase; poly(1,4-α-D-galactosiduronate) digalacturono-

hydrolase

**Systematic name:**  $poly[(1\rightarrow 4)-\alpha-D-galactosiduronate]$  digalacturonohydrolase

**References:** [1067, 1075, 1076]

[EC 3.2.1.82 created 1972]

#### EC 3.2.1.83

**Accepted name:** κ-carrageenase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-linkages between D-galactose 4-sulfate and 3,6-anhydro-D-galactose in

 $\kappa\text{-}carrageen ans$ 

Other name(s):  $\kappa$ -carrageenan 4- $\beta$ -D-glycanohydrolase

**Systematic name:** κ-carrageenan 4-β-D-glycanohydrolase (configuration-retaining)

Comments: The main products of hydrolysis are neocarrabiose-sulfate and neocarratetraose-sulfate [1903]. Un-

like EC 3.2.1.157 (ι-carrageenase), but similar to EC 3.2.1.81 (β-agarase), this enzyme proceeds with

retention of the anomeric configuration.

**References:** [3150, 2318, 2316, 1902, 1903]

[EC 3.2.1.83 created 1972, modified 2006]

#### EC 3.2.1.84

**Accepted name:** glucan 1,3-α-glucosidase

**Reaction:** Hydrolysis of terminal  $(1\rightarrow 3)$ - $\alpha$ -D-glucosidic links in  $(1\rightarrow 3)$ - $\alpha$ -D-glucans **Other name(s):** exo-1,3- $\alpha$ -glucanase; glucosidase II; 1,3- $\alpha$ -D-glucan 3-glucohydrolase

**Systematic name:** 3-α-D-glucan 3-glucohydrolase **Comments:** Does not act on nigeran.

**References:** [3348]

[EC 3.2.1.84 created 1972]

#### EC 3.2.1.85

**Accepted name:** 6-phospho-β-galactosidase

**Reaction:** a 6-phospho- $\beta$ -D-galactoside + H<sub>2</sub>O = 6-phospho-D-galactose + an alcohol

Other name(s): phospho-β-galactosidase; β-D-phosphogalactoside galactohydrolase; phospho-β-D-galactosidase; 6-

phospho-β-D-galactosidase

**Systematic name:** 6-phospho-β-D-galactoside 6-phosphogalactohydrolase

**References:** [1111]

[EC 3.2.1.85 created 1976]

#### EC 3.2.1.86

**Accepted name:** 6-phospho-β-glucosidase

**Reaction:** 6-phospho- $\beta$ -D-glucosyl-(1 $\rightarrow$ 4)-D-glucose + H<sub>2</sub>O = D-glucose + D-glucose 6-phosphate

Other name(s): phospho-β-glucosidase A; phospho-β-glucosidase; phosphocellobiase; 6-phospho-β-D-glucosyl-(1,4)-

D-glucose glucohydrolase

**Systematic name:** 6-phospho- $\beta$ -D-glucosyl- $(1\rightarrow 4)$ -D-glucose glucohydrolase

Comments: Also hydrolyses several other phospho- $\beta$ -D-glucosides, but not their non-phosphorylated forms.

**References:** [2230]

[EC 3.2.1.86 created 1976]

### EC 3.2.1.87

Accepted name: capsular-polysaccharide endo-1,3-α-galactosidase

**Reaction:** Random hydrolysis of  $(1\rightarrow 3)$ - $\alpha$ -D-galactosidic linkages in *Aerobacter aerogenes* capsular polysac-

charide

**Other name(s):** polysaccharide depolymerase; capsular polysaccharide galactohydrolase

Systematic name: Aerobacter-capsular-polysaccharide galactohydrolase

 $\textbf{Comments:} \quad \text{Hydrolyses the galactosyl-} \alpha\text{-1,3-D-galactose linkages only in the complex substrate, bringing about}$ 

depolymerization.

**References:** [3313, 3314]

[EC 3.2.1.87 created 1976]

#### EC 3.2.1.88

**Accepted name:** non-reducing end  $\beta$ -L-arabinopyranosidase

 $\begin{tabular}{lll} \textbf{Removal of a terminal $\beta$-L-arabinopyranose residue from the non-reducing end of its substrate.} \\ \textbf{Other name(s):} & vicianosidase; $\beta$-L-arabinosidase (ambiguous); $\beta$-L-arabinoside arabinohydrolase (ambiguous).} \\ \end{tabular}$ 

**Systematic name:**  $\beta$ -L-arabinopyranoside non-reducing end  $\beta$ -L-arabinopyranosidase

**Comments:** The enzyme, which was characterized from dormant seeds of the plant *Cajanus cajan* (pigeon pea),

has been shown to remove the terminal non-reducing  $\beta$ -L-arabinopyranoside residue from the artificial substrate p-nitrophenyl- $\beta$ -L-arabinopyranose [599]. In the presence of methanol the enzyme demonstrates transglycosylase activity, transferring the arabinose moiety to methanol while retaining the

anomeric configuration, generating 1-O-methyl-β-L-arabinopyranose [598].

**References:** [599, 598]

# [EC 3.2.1.88 created 1976, modified 2013]

EC 3.2.1.89

**Accepted name:** arabinogalactan endo-β-1,4-galactanase

**Reaction:** The enzyme specifically hydrolyses  $(1\rightarrow 4)$ - $\beta$ -D-galactosidic linkages in type I arabinogalactans. **Other name(s):** endo-1,4- $\beta$ -galactanase; galactanase (ambiguous); arabinogalactanase; ganB (gene name)

**Systematic name:** arabinogalactan 4-β-D-galactanohydrolase

**Comments:** This enzyme, isolated from the bacterium *Bacillus subtilis*, hydrolyses the  $\beta(1\rightarrow 4)$  bonds found in

type I plant arabinogalactans, which are a component of the primary cell walls of dicots. The predom-

inant product is a tetrasaccharide. cf. EC 3.2.1.181, galactan endo-β-1,3-galactanase.

**References:** [699, 1583, 2642]

[EC 3.2.1.89 created 1976, modified 2012]

[3.2.1.90 Deleted entry. arabinogalactan endo-1,3- $\beta$ -galactosidase. The enzyme was not sufficiently characterized to warrant an EC number]

[EC 3.2.1.90 created 1976, deleted 2001]

EC 3.2.1.91

**Accepted name:** cellulose 1,4-β-cellobiosidase (non-reducing end)

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-glucosidic linkages in cellulose and cellotetraose, releasing cellobiose from

the non-reducing ends of the chains

Other name(s): exo-cellobiohydrolase; β-1,4-glucan cellobiohydrolase; β-1,4-glucan cellobiosylhydrolase; 1,4-β-

glucan cellobiosidase; exoglucanase; avicelase; CBH 1;  $C_1$  cellulase; cellobiohydrolase I; cellobiohydrolase; exo- $\beta$ -1,4-glucan cellobiohydrolase; 1,4- $\beta$ -D-glucan cellobiohydrolase; cellobiosidase

Systematic name: 4-B-D-glucan cellobiohydrolase (non-reducing end)

**References:** [195, 713, 1030]

[EC 3.2.1.91 created 1976, modified 2011]

EC 3.2.1.92

**Accepted name:** peptidoglycan β-*N*-acetylmuramidase

**Reaction:** Hydrolysis of terminal, non-reducing *N*-acetylmuramic residues

Other name(s):  $\exp{-\beta-N}$ -acetylmuramidase;  $\exp{-\beta}$ -acetylmuramidase;  $\exp{-\beta}$ -acetamido-3-O-(D-1-carboxyethyl)-2-

deoxy-D-glucoside acetamidodeoxyglucohydrolase

**Systematic name:** peptidoglycan β-*N*-acetylmuramoylexohydrolase

**References:** [2428]

[EC 3.2.1.92 created 1976]

EC 3.2.1.93

**Accepted name:** α,α-phosphotrehalase

**Reaction:**  $\alpha, \alpha$ -trehalose 6-phosphate + H<sub>2</sub>O = D-glucose + D-glucose 6-phosphate

**Other name(s):** phosphotrehalase

**Systematic name:** α,α-trehalose-6-phosphate phosphoglucohydrolase

**References:** [212]

[EC 3.2.1.93 created 1976]

EC 3.2.1.94

**Accepted name:** glucan 1,6-α-isomaltosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic linkages in polysaccharides, to remove successive isomaltose

units from the non-reducing ends of the chains

Other name(s): exo-isomaltohydrolase; isomalto-dextranase; isomaltodextranase; G2-dextranase; 1,6-α-D-glucan iso-

maltohydrolase

**Systematic name:** 6-α-D-glucan isomaltohydrolase

Comments: Optimum activity is on those 1,6- $\alpha$ -D-glucans containing 6, 7 and 8 glucose units; those containing 3,

4 and 5 glucose units are hydrolysed at slower rates.

**References:** [2546, 2545]

[EC 3.2.1.94 created 1976]

EC 3.2.1.95

**Accepted name:** dextran 1,6-α-isomaltotriosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic linkages in dextrans, to remove successive isomaltotriose units

from the non-reducing ends of the chains

**Other name(s):** exo-isomaltotriohydrolase; 1,6-α-D-glucan isomaltotriohydrolase

**Systematic name:** 6-α-D-glucan isomaltotriohydrolase

**References:** [2795]

[EC 3.2.1.95 created 1978]

EC 3.2.1.96

**Accepted name:** mannosyl-glycoprotein endo- $\beta$ -N-acetylglucosaminidase

**Reaction:** Endohydrolysis of the N,N'-diacetylchitobiosyl unit in high-mannose glycopeptides and glycopro-

teins containing the -[Man(GlcNAc)<sub>2</sub>]Asn- structure. One N-acetyl-D-glucosamine residue remains

attached to the protein; the rest of the oligosaccharide is released intact

Other name(s): N,N'-diacetylchitobiosyl  $\beta$ -N-acetylglucosaminidase; endo- $\beta$ -N-acetylglucosaminidase; mannosyl-

glycoprotein endo- $\beta$ -N-acetylglucosamidase; di-N-acetylchitobiosyl  $\beta$ -N-acetylglucosaminidase; endo- $\beta$ -acetylglucosaminidase; endo- $\beta$ -(1 $\rightarrow$ 4)-N-acetylglucosaminidase; mannosyl-glycoprotein

1,4-N-acetamidodeoxy-β-D-glycohydrolase; endoglycosidase S; endo-N-acetyl-β-D-

glucosaminidase; endo-N-acetyl- $\beta$ -glucosaminidase; endo- $\beta$ -N-acetylglucosaminidase D; endo- $\beta$ -N-acetylglucosaminidase H; endo- $\beta$ -N-acetylglucosaminidase

 $L; glycopeptide-D-mannosyl-4-\textit{N-}(\textit{N-}acetyl-D-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-\beta-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-b-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-b-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-b-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-b-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-b-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-b-glucosaminyl})_2-a sparagine~1,4-\textit{N-}acetyl-b-glucosaminyl})_2-acetyl-b-glucosaminyl)_2-acetyl-b-glucosaminyl_2-acetyl-b-glucosaminyl_2-acetyl-b-glucosaminyl_2-acetyl-b-glucosaminyl_2-acetyl-b-glucosaminyl_2-acetyl-b-glucosaminyl_2-acetyl-b-glucosaminyl_2$ 

glucosaminohydrolase; endoglycosidase H

**Systematic name:** glycopeptide-D-mannosyl- $N^4$ -(N-acetyl-D-glucosaminyl)<sub>2</sub>-asparagine 1,4-N-acetyl- $\beta$ -

glucosaminohydrolase

**Comments:** A group of related enzymes.

**References:** [431, 1498, 2277, 2278, 2831, 2888]

[EC 3.2.1.96 created 1978]

EC 3.2.1.97

**Accepted name:** endo-α-*N*-acetylgalactosaminidase

**Reaction:**  $\beta$ -D-galactosyl- $(1 \rightarrow 3)$ -N-acetyl- $\alpha$ -D-galactosaminyl-[glycoprotein]-L-serine/L-threonine + H<sub>2</sub>O =

 $\beta$ -D-galactosyl-(1 $\rightarrow$ 3)-N-acetyl-D-galactosamine + [glycoprotein]-L-serine/L-threonine

Other name(s): endo- $\alpha$ -acetylgalactosaminidase; endo- $\alpha$ -N-acetyl-D-galactosaminidase; mucinaminylserine muci-

naminidase; D-galactosyl-3-(N-acetyl- $\alpha$ -D-galactosaminyl)-L-serine mucinaminohydrolase; endo- $\alpha$ -GalNAc-ase; glycopeptide  $\alpha$ -N-acetylgalactosaminidase; D-galactosyl-N-acetyl- $\alpha$ -D-galactosamine

D-galactosyl-N-acetyl-galactosaminohydrolase

**Systematic name:** glycopeptide-D-galactosyl-*N*-acetyl-α-D-galactosamine D-galactosyl-*N*-acetyl-

galactosaminohydrolase

**Comments:** The enzyme catalyses the liberation of Gal- $(1\rightarrow 3)$ - $\beta$ -GalNAc  $\alpha$ -linked to serine or threonine residues

of mucin-type glycoproteins. EngBF from the bacterium *Bifidobacterium longum* specifically acts on core 1-type O-glycan to release the disaccharide Gal- $(1\rightarrow 3)$ - $\beta$ -GalNAc. The enzymes from the bacteria *Clostridium perfringens*, *Enterococcus faecalis*, Propionibacterium acnes and *Alcaligenes faecalis* show broader specificity (e.g. they can also release the core 2 trisaccharide Gal- $(1\rightarrow 3)$ - $\beta$ -(GlcNAc- $(1\rightarrow 6)$ - $\beta$ )-GalNAc or the core 3 disaccharide GlcNAc- $(1\rightarrow 3)$ - $\beta$ -GalNAc) [85, 1523]. The enzyme may play an important role in the degradation and utilization of mucins having core 1 O-glycan.

**References:** [85, 1523, 850, 2814, 976, 88, 936]

[EC 3.2.1.97 created 1978 (EC 3.2.1.110 created 1984, incorporated 2008), modified 2008, modified 2011]

# EC 3.2.1.98

**Accepted name:** glucan 1,4-α-maltohexaosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkages in amylaceous polysaccharides, to remove successive

maltohexaose residues from the non-reducing chain ends

**Other name(s):** exo-maltohexaohydrolase; 1,4- $\alpha$ -D-glucan maltohexaohydrolase

**Systematic name:** 4-α-D-glucan maltohexaohydrolase

Comments: cf. EC 3.2.1.3 glucan 1,4-α-glucosidase, which removes successive glucose residues; EC 3.2.1.2 β-

amylase, which removes successive maltose residues; EC 3.2.1.116 glucan 1,4- $\alpha$ -maltotriohydrolase, which removes successive maltotriose units and EC 3.2.1.60 glucan 1,4- $\alpha$ -maltotetraohydrolase,

which removes successive maltotetraose residues. The products have the  $\alpha$ -configuration.

**References:** [1353, 2043]

[EC 3.2.1.98 created 1978]

# EC 3.2.1.99

**Accepted name:** arabinan endo-1,5-α-L-arabinanase

**Reaction:** Endohydrolysis of  $(1 \rightarrow 5)$ - $\alpha$ -arabinofuranosidic linkages in  $(1 \rightarrow 5)$ -arabinans

**Other name(s):** endo-1,5- $\alpha$ -L-arabinanase; endo- $\alpha$ -1,5-arabanase; endo-arabanase; 1,5- $\alpha$ -L-arabinan 1,5- $\alpha$ -L-

arabinanohydrolase; arabinan endo-1,5- $\alpha$ -L-arabinosidase (misleading)

**Systematic name:**  $5-\alpha$ -L-arabinan  $5-\alpha$ -L-arabinanohydrolase

**Comments:** Acts best on linear 1,5- $\alpha$ -L-arabinan. Also acts on branched arabinan, but more slowly.

**References:** [1354, 3154, 785, 1612]

[EC 3.2.1.99 created 1981, modified 2011]

# EC 3.2.1.100

**Accepted name:** mannan 1,4-mannobiosidase

 $\textbf{Reaction:} \quad \text{Hydrolysis of } (1 \rightarrow 4) \text{-}\beta \text{-}D \text{-}mannosidic linkages in } (1 \rightarrow 4) \text{-}\beta \text{-}D \text{-}mannans, to remove successive mannosidic linkages}$ 

biose residues from the non-reducing chain ends

**Other name(s):** 1,4- $\beta$ -D-mannan mannobiohydrolase; exo- $\beta$ -mannanase; exo-1,4- $\beta$ -mannobiohydrolase

**Systematic name:** 4-β-D-mannan mannobiohydrolase

**References:** [67]

[EC 3.2.1.100 created 1983]

# EC 3.2.1.101

**Accepted name:** mannan endo-1,6-α-mannosidase

**Reaction:** Random hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-mannosidic linkages in unbranched  $(1\rightarrow 6)$ -mannans

Other name(s): endo- $\alpha$ -1 $\rightarrow$ 6-D-mannanase; endo-1,6- $\beta$ -mannanase; mannan endo-1,6- $\beta$ -mannosidase; 1,6- $\alpha$ -D-

mannan mannanohydrolase

**Systematic name:** 6-α-D-mannan mannanohydrolase

**References:** [2041, 289, 2040]

# [EC 3.2.1.101 created 1984, modified 2001]

EC 3.2.1.102

**Accepted name:** blood-group-substance endo-1,4-β-galactosidase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ -β-D-galactosidic linkages in blood group A and B substances Other name(s): endo-β-galactosidase (ambiguous); blood-group-substance 1,4-β-D-galactanohydrolase

**Systematic name:** blood-group-substance 4-β-D-galactanohydrolase

**Comments:** Hydrolyses the 1,4- $\beta$ -D-galactosyl linkages adjacent to a 1,3- $\alpha$ -D-galactosyl or N-

acetylgalactosaminyl residues and a 1,2-α-D-fucosyl residue.

**References:** [858, 2054, 2852]

[EC 3.2.1.102 created 1984]

EC 3.2.1.103

**Accepted name:** keratan-sulfate endo-1,4-β-galactosidase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-galactosidic linkages in keratan sulfate

Other name(s): endo-β-galactosidase (ambiguous); keratan sulfate endogalactosidase; keratanase; keratan-sulfate 1,4-

 $\beta$ -D-galactanohydrolase

**Systematic name:** keratan-sulfate 4-β-D-galactanohydrolase

**Comments:** Hydrolyses the 1,4- $\beta$ -D-galactosyl linkages adjacent to 1,3-*N*-acetyl- $\alpha$ -D-glucosaminyl residues.

Also acts on some non-sulfated oligosaccharides, but only acts on blood group substances when the 1,2-linked fucosyl residues have been removed ( $\it cf$ . EC 3.2.1.102 blood-group-substance endo-1,4- $\beta$ -

galactosidase).

**References:** [858]

[EC 3.2.1.103 created 1984]

EC 3.2.1.104

**Accepted name:** steryl-β-glucosidase

**Reaction:** cholesteryl- $\beta$ -D-glucoside + H<sub>2</sub>O = D-glucose + cholesterol

Systematic name: cholesteryl- $\beta$ -D-glucoside glucohydrolase

**Comments:** Acts on glucosides of cholesterol and sitosterol, but not on some related sterols such as coprostanol.

**References:** [1364]

[EC 3.2.1.104 created 1984]

EC 3.2.1.105

**Accepted name:**  $3\alpha(S)$ -strictosidine  $\beta$ -glucosidase

**Reaction:** strictosidine +  $H_2O = D$ -glucose + strictosidine aglycone

**Systematic name:** strictosidine  $\beta$ -D-glucohydrolase

Comments: Does not act on a number of closely related glycosides. Strictosidine is a precursor of indole alka-

loids.

**References:** [1109, 144]

[EC 3.2.1.105 created 1984]

EC 3.2.1.106

**Accepted name:** mannosyl-oligosaccharide glucosidase

**Reaction:**  $Glc_3Man_9GlcNAc_2$ -[protein] +  $H_2O = Glc_2Man_9GlcNAc_2$ -[protein] +  $\beta$ -D-glucopyranose

Other name(s): Glc3Man9NAc2 oligosaccharide glucosidase; trimming glucosidase I; CWH41 (gene name); MOGS

(gene name); mannosyl-oligosaccharide glucohydrolase

**Systematic name:** Glc<sub>3</sub>Man<sub>9</sub>GlcNAc<sub>2</sub>-[protein] glucohydrolase (configuration-inverting)

**Comments:** This enzyme catalyses the first step in the processing of the N-glycan tetradecasaccharide precursor

Glc<sub>3</sub>Man<sub>9</sub>GlcNAc<sub>2</sub>, which takes place in the endoplasmic reticulum, by removing the distal  $\alpha$ -1,2-linked glucose residue. This and subsequent processing steps are required before complex *N*-glycans

can be synthesized.

**References:** [696, 981, 1441, 982, 1812]

[EC 3.2.1.106 created 1984, modified 2018]

#### EC 3.2.1.107

Accepted name: protein-glucosylgalactosylhydroxylysine glucosidase

**Reaction:** [collagen]-(5R)-5-O-[ $\alpha$ -D-glucosyl- $(1\rightarrow 2)$ - $\beta$ -D-galactosyl]-5-hydroxy-L-lysine + H<sub>2</sub>O = D-glucose +

[collagen]-(5R)-5-O- $(\beta$ -D-galactosyl)-5-hydroxy-L-lysine

**Other name(s):** PGGHG (gene name); 2-*O*-α-D-glucopyranosyl-5-*O*-α-D-galactopyranosylhydroxy-L-lysine gluco-

hydrolase; protein-α-D-glucosyl-1,2-β-D-galactosyl-L-hydroxylysine glucohydrolase; protein-α-D-

glucosyl- $(1\rightarrow 2)$ - $\beta$ -D-galactosyl-L-hydroxylysine glucohydrolase

**Systematic name:** [collagen]-(5R)-5-O-[ $\alpha$ -D-glucosyl- $(1\rightarrow 2)$ - $\beta$ -D-galactosyl]-5-hydroxy-L-lysine glucohydrolase

**Comments:** The enzyme specifically hydrolyses glucose from  $\alpha$ -D-glucosyl- $(1\rightarrow 2)$ - $\beta$ -D-galactosyl disaccharide

units that are linked to hydroxylysine residues of collagen and collagen-like proteins. Acetylation of

the ε-amino group of the glycosylated hydroxylysine abolishes activity.

**References:** [1035, 1036, 2766, 1034]

[EC 3.2.1.107 created 1984]

#### EC 3.2.1.108

Accepted name: lactase

**Reaction:** lactose +  $H_2O$  = D-galactose + D-glucose

Other name(s): lactase-phlorizin hydrolase Systematic name: lactose galactohydrolase

Comments: The enzyme from intestinal mucosa is isolated as a complex that also catalyses the reaction of EC

3.2.1.62 glycosylceramidase. cf. EC 3.2.1.33 amylo-α-1,6-glucosidase.

**References:** [1736, 2365, 2567, 2685, 2686, 89]

[EC 3.2.1.108 created 1984]

# EC 3.2.1.109

Accepted name: endogalactosaminidase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-galactosaminidic linkages in poly(D-galactosamine)

**Systematic name:** galactosaminoglycan glycanohydrolase

**References:** [2403, 2867]

[EC 3.2.1.109 created 1984]

[3.2.1.110 Deleted entry. mucinaminylserine mucinaminidase. The enzyme is identical to EC 3.2.1.97, glycopeptide  $\alpha$ -N-acetylgalactosaminidase]

[EC 3.2.1.110 created 1984, deleted 2008]

### EC 3.2.1.111

**Accepted name:**  $1,3-\alpha$ -L-fucosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 3)$ -linkages between  $\alpha$ -L-fucose and N-acetylglucosamine residues in glycopro-

teins

Other name(s): almond emulsin fucosidase I

**Systematic name:** 3-α-L-fucosyl-*N*-acetylglucosaminyl-glycoprotein fucohydrolase

**Comments:** Not identical with EC 3.2.1.63 1,2- $\alpha$ -L-fucosidase.

**References:** [1243, 2144, 3303]

#### [EC 3.2.1.111 created 1986]

EC 3.2.1.112

Accepted name: 2-deoxyglucosidase

**Reaction:** a 2-deoxy- $\alpha$ -D-glucoside + H<sub>2</sub>O = 2-deoxy-D-glucose + an alcohol

Other name(s): 2-deoxy-α-glucosidase; 2-deoxy-α-D-glucosidase Systematic name: 2-deoxy-α-D-glucoside deoxyglucohydrolase

**References:** [366]

[EC 3.2.1.112 created 1986]

EC 3.2.1.113

**Accepted name:** mannosyl-oligosaccharide 1,2-α-mannosidase

**Reaction:** Hydrolysis of the terminal  $(1\rightarrow 2)$ -linked  $\alpha$ -D-mannose residues in the oligo-mannose oligosaccharide

Man<sub>9</sub>(GlcNAc)<sub>2</sub>

Other name(s): mannosidase 1A; mannosidase 1B; 1,2-α-mannosidase; exo-α-1,2-mannanase; mannose-9 processing

α-mannosidase; glycoprotein processing mannosidase I; mannosidase I; Man<sub>9</sub>-mannosidase; ManI;

1,2-α-mannosyl-oligosaccharide α-D-mannohydrolase

**Systematic name:**  $2-\alpha$ -mannosyl-oligosaccharide  $\alpha$ -D-mannohydrolase

**Comments:** Involved in the synthesis of glycoproteins.

**References:** [2826, 2994]

[EC 3.2.1.113 created 1986]

EC 3.2.1.114

**Accepted name:** mannosyl-oligosaccharide 1,3-1,6-α-mannosidase

**Reaction:** Man<sub>5</sub>GlcNAc<sub>3</sub>-[protein] + 2 H<sub>2</sub>O = Man<sub>3</sub>GlcNAc<sub>3</sub>-[protein] + 2  $\alpha$ -D-mannopyranose

Other name(s): MAN2A1 (gene name); MAN2A2 (gene name); mannosidase II; exo-1,3-1,6-α-mannosidase;

 $\alpha$ -D-mannosidase II;  $\alpha$ -mannosidase II;  $\alpha$ 1-3,6-mannosidase; GlcNAc transferase I-dependent  $\alpha$ 1,3[ $\alpha$ 1,6]mannosidase; Golgi  $\alpha$ -mannosidase II; ManII; 1,3(1,6)- $\alpha$ -D-mannosidase; 1,3-(1,6-)mannosyl-oligosaccharide  $\alpha$ -D-mannohydrolase; (1 $\rightarrow$ 3)-(1 $\rightarrow$ 6)-mannosyl-oligosaccharide  $\alpha$ -D-mannohydrolase; (1 $\rightarrow$ 3)-(1 $\rightarrow$ 6)-mannosyl-oligosaccharide  $\alpha$ -D-mannohydrolase; (1 $\rightarrow$ 3)-(1 $\rightarrow$ 6)-mannohydrolase; (1 $\rightarrow$ 8)-mannohydrolase; (1 $\rightarrow$ 8)-mannohydrolase

mannohydrolase

**Systematic name:**  $(1\rightarrow 3)$ - $(1\rightarrow 6)$ -mannosyl-oligosaccharide  $\alpha$ -D-mannohydrolase (configuration-retaining)

**Comments:** The enzyme, found in plants and animals, participates in the processing of N-glycans in the Golgi ap-

paratus. It removes two mannosyl residues, one linked by  $\alpha$ 1,3 linkage, and the other linked by  $\alpha$ 1,6 linkage, both of which are removed by the same catalytic site. The enzyme is sensitive to swainso-

nine.

**References:** [2995, 2825, 1057, 2994, 1964, 1926, 3036, 92, 2615, 2458]

[EC 3.2.1.114 created 1986, modified 2018]

EC 3.2.1.115

**Accepted name:** branched-dextran exo-1,2-α-glucosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 2)$ - $\alpha$ -D-glucosidic linkages at the branch points of dextrans and related polysaccha-

rides, producing free D-glucose

Other name(s): dextran 1,2- $\alpha$ -glucosidase; dextran  $\alpha$ -1,2 debranching enzymel 1,2- $\alpha$ -D-glucosyl-branched-dextran

2-glucohydrolase

**Systematic name:**  $(1\rightarrow 2)$ - $\alpha$ -D-glucosyl-branched-dextran 2-glucohydrolase

**Comments:** Does not hydrolyse disaccharides or oligosaccharides containing linear 1,2-α-glucosidic linkages.

**References:** [1933, 1934]

[EC 3.2.1.115 created 1989]

# EC 3.2.1.116

**Accepted name:** glucan 1,4-α-maltotriohydrolase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkages in amylaceous polysaccharides, to remove successive

maltotriose residues from the non-reducing chain ends

Other name(s): exo-maltotriohydrolase; maltotriohydrolase; 1,4-α-D-glucan maltotriohydrolase

**Systematic name:** 4-α-D-glucan maltotriohydrolase

**Comments:** cf. EC 3.2.1.2 (β-amylase), EC 3.2.1.60 (glucan 1,4-α-maltotetraohydrolase) and EC 3.2.1.98 (glucan

1,4- $\alpha$ -maltohexaosidase). The products have the  $\alpha$ -configuration.

**References:** [2043]

[EC 3.2.1.116 created 1989]

# EC 3.2.1.117

**Accepted name:** amygdalin β-glucosidase

**Reaction:** (*R*)-amygdalin +  $H_2O = (R)$ -prunasin + D-glucose

Other name(s): amygdalase; amygdalinase; amygdalin hydrolase; amygdalin glucosidase

**Systematic name:** amygdalin β-D-glucohydrolase

**Comments:** Highly specific; does not act on prunasin, linamarin, gentiobiose or cellobiose (cf. EC 3.2.1.21 β-

glucosidase).

**References:** [1572]

[EC 3.2.1.117 created 1989]

### EC 3.2.1.118

**Accepted name:** prunasin β-glucosidase

**Reaction:** (*R*)-prunasin +  $H_2O$  = D-glucose + mandelonitrile

**Other name(s):** prunasin hydrolase

**Systematic name:** prunasin  $\beta$ -D-glucohydrolase

**Comments:** Highly specific; does not act on amygdalin, linamarin or gentiobiose. (*cf.* EC 3.2.1.21 β-glucosidase).

**References:** [1572]

[EC 3.2.1.118 created 1989]

### EC 3.2.1.119

**Accepted name:** vicianin β-glucosidase

**Reaction:** (*R*)-vicianin +  $H_2O$  = mandelonitrile + vicianose

**Other name(s):** vicianin hydrolase

**Systematic name:** (R)-vicianin  $\beta$ -D-glucohydrolase

**Comments:** Also hydrolyses, more slowly, (*R*)-amygdalin and (*R*)-prunasin, but not gentiobiose, linamarin or cel-

lobiose.

**References:** [1572]

[EC 3.2.1.119 created 1989]

# EC 3.2.1.120

**Accepted name:** oligoxyloglucan  $\beta$ -glycosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-glucosidic links in oligoxyloglucans so as to remove successive iso-

primeverose [i.e.  $\alpha$ -xylo- $(1\rightarrow 6)$ - $\beta$ -D-glucosyl-] residues from the non-reducing chain ends

Other name(s): isoprimeverose-producing oligoxyloglucan hydrolase; oligoxyloglucan hydrolase

**Systematic name:** oligoxyloglucan xyloglucohydrolase

References: [1402]

[EC 3.2.1.120 created 1989]

# EC 3.2.1.121

Accepted name: polymannuronate hydrolase

**Reaction:** Endohydrolysis of the D-mannuronide linkages of polymannuronate

Other name(s): polymannuronic acid polymerase Systematic name: poly(mannuronide) mannuronohydrolase

**Comments:** Does not act on alginic acid, which is a copolymer of polymannuronate.

**References:** [668]

[EC 3.2.1.121 created 1989]

#### EC 3.2.1.122

**Accepted name:** maltose-6'-phosphate glucosidase

**Reaction:**  $\alpha$ -maltose 6'-phosphate +  $H_2O$  = D-glucose + D-glucose 6-phosphate **Other name(s):** phospho- $\alpha$ -glucosidase; maltose-6'-phosphate 6-phosphoglucohydrolase

**Systematic name:** α-maltose-6'-phosphate 6-phosphoglucohydrolase

**Comments:** Hydrolyses a variety of 6-phospho- $\alpha$ -D-glucosides, including  $\alpha$ -maltose 6'-phosphate,  $\alpha$ , $\alpha$ -trehalose

6-phosphate, sucrose 6-phosphate and p-nitrophenyl- $\alpha$ -D-glucopyranoside 6-phosphate (as a chromogenic substrate). The enzyme is activated by  $Fe^{II}$ ,  $Mn^{II}$ ,  $Co^{II}$  and  $Ni^{II}$ . It is rapidly inactivated in

air.

**References:** [2916]

[EC 3.2.1.122 created 1989, modified 1999]

#### EC 3.2.1.123

Accepted name: endoglycosylceramidase

**Reaction:** oligoglycosylglucosyl- $(1 \leftrightarrow 1)$ -ceramide +  $H_2O$  = ceramide + oligoglycosylglucose

Other name(s): endoglycoceramidase; EGCase; glycosyl-*N*-acetyl-sphingosine 1,1-β-D-glucanohydrolase, oligogly-

cosylglucosylceramide glycohydrolase; oligoglycosylglucosyl(1↔1)ceramide glycohydrolase

**Systematic name:** oligoglycosylglucosyl- $(1 \leftrightarrow 1)$ -ceramide glycohydrolase

Comments: An enzyme from *Rhodococcus* sp. that degrades various acidic and neutral glycosphingolipids to

oligosaccharides and ceramides, by cleaving a glucosyl bond. Does not act on monoglycosylce-

ramides. cf. EC 3.2.1.62 glycosylceramidase.

**References:** [1266]

[EC 3.2.1.123 created 1989]

# EC 3.2.1.124

**Accepted name:** 3-deoxy-2-octulosonidase

**Reaction:** Endohydrolysis of the  $\beta$ -ketopyranosidic linkages of 3-deoxy-D-manno-2-octulosonate in capsular

polysaccharides

Other name(s): 2-keto-3-deoxyoctonate hydrolase; octulosylono hydrolase; octulofuranosylono hydrolase; octulopy-

ranosylonohydrolase

**Systematic name:** capsular-polysaccharide 3-deoxy-D-manno-2-octulosonohydrolase

Comments: The enzyme from a bacteriophage catalyses the depolymerization of capsular polysaccharides con-

taining 3-deoxy-2-octulosonide in the cell wall of Escherichia coli.

**References:** [42]

[EC 3.2.1.124 created 1989]

# EC 3.2.1.125

**Accepted name:** raucaffricine β-glucosidase

**Reaction:** raucaffricine +  $H_2O$  = D-glucose + vomilenine

Other name(s): raucaffricine  $\beta$ -D-glucosidase; raucaffricine glucosidase

**Systematic name:** raucaffricine  $\beta$ -D-glucohydrolase

**Comments:** Highly specific; some other ajmalan glucoside alkaloids are hydrolysed, but more slowly.

**References:** [2587]

[EC 3.2.1.125 created 1989]

EC 3.2.1.126

**Accepted name:** coniferin β-glucosidase

**Systematic name:** coniferin  $\beta$ -D-glucosidase

**Comments:** Also hydrolyses syringin, 4-cinnamyl alcohol β-glucoside and, more slowly, some other aryl β-

glycosides. A plant cell-wall enzyme involved in the biosynthesis of lignin.

**References:** [1182, 1807]

[EC 3.2.1.126 created 1989]

EC 3.2.1.127

**Accepted name:** 1,6-α-L-fucosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ -linkages between  $\alpha$ -L-fucose and N-acetyl-D-glucosamine in glycopeptides

such as immunoglobulin G glycopeptide and fucosyl-asialo-agalacto-fetuin

**Other name(s):** α-L-fucosidase; 1,6-L-fucosyl-*N*-acetyl-D-glucosaminylglycopeptide fucohydrolase

**Systematic name:** 6-L-fucosyl-*N*-acetyl-D-glucosaminylglycopeptide fucohydrolase

**Comments:** The enzyme from *Aspergillus niger* does not act on 1,2-, 1,3-, or 1,4-L-fucosyl linkages.

References: [3280]

[EC 3.2.1.127 created 1989]

EC 3.2.1.128

**Accepted name:** glycyrrhizin hydrolase

**Reaction:** glycyrrhizin + H<sub>2</sub>O =  $\beta$ -D-glucuronosyl- $(1\rightarrow 2)$ -D-glucuronate + glycyrrhetinate

Other name(s): glycyrrhizinate  $\beta$ -glucuronidase; glycyrrhizin  $\beta$ -hydrolase; glycyrrhizinic acid hydrolase

**Systematic name:** glycyrrhizinate glucuronosylhydrolase

**Comments:** The enzyme from Aspergillus niger is specific for the hydrolysis of the triterpenoid glycoside gly-

cyrrhizin from roots of Glycyrrhiza sp.

**References:** [2008]

[EC 3.2.1.128 created 1989]

EC 3.2.1.129

**Accepted name:** endo-α-sialidase

**Reaction:** Endohydrolysis of  $(2\rightarrow 8)$ - $\alpha$ -sialosyl linkages in oligo- or poly(sialic) acids

**Other name(s):** endo-*N*-acylneuraminidase; endoneuraminidase; endo-*N*-acetylneuraminidase; poly( $\alpha$ -2,8-sialosyl)

 $endo-\textit{N}-acetylneuraminidase; poly(\alpha-2,8-sialoside) ~\alpha-2,8-sialosylhydrolase; endosialidase; endo-Nacetylneuraminidase; endo-N$ 

**Systematic name:** polysialoside  $(2\rightarrow 8)$ - $\alpha$ -sialosylhydrolase

**Comments:** Although the name endo-*N*-acetylneuraminidase has also been used for this enzyme, this is mislead-

ing since its activity is not restricted to acetylated substrates. An exo- $\alpha$ -sialidase activity is listed as

EC 3.2.1.18 exo-α-sialidase. See also EC 4.2.2.15 anhydrosialidase.

**References:** [769, 1029, 1470, 1580, 2253, 2932, 347]

[EC 3.2.1.129 created 1990, modified 1999]

# EC 3.2.1.130

**Accepted name:** glycoprotein endo-α-1,2-mannosidase

**Reaction:** GlcMan<sub>9</sub>GlcNAc<sub>2</sub>-[protein] +  $H_2O = Man_8GlcNAc_2$ -[protein] (isomer  $8A_{1,2,3}B_{1,2}$ ) +  $\alpha$ -D-glucosyl-

 $(1\rightarrow 3)$ - $\alpha$ -D-mannopyranose

Other name(s): glucosylmannosidase; endo-α-mannosidase; endo-α-mannosidase; endomannosidase; glucosyl

mannosidase; MANEA (gene name); glycoprotein glucosylmannohydrolase

**Systematic name:** glycoprotein glucosylmannohydrolase (configuration-retaining)

**Comments:** The enzyme catalyses the hydrolysis of the terminal  $\alpha$ -D-glucosyl- $(1\rightarrow 3)$ -D-mannosyl unit from the

 $GlcMan_9(GlcNAc)_2$  oligosaccharide component of N-glucosylated proteins during their processing in the Golgi apparatus. The name for the isomer is based on a nomenclature proposed by Prien et al

[2325].

**References:** [1749, 2993, 1144, 2741, 1037, 1052, 2325]

[EC 3.2.1.130 created 1990, modified 2017]

#### EC 3.2.1.131

**Accepted name:** xylan  $\alpha$ -1,2-glucuronosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 2)$ - $\alpha$ -D-(4-O-methyl)glucuronosyl links in the main chain of hardwood xylans **Other name(s):** 1,2- $\alpha$ -glucuronidase;  $\alpha$ - $(1\rightarrow 2)$ -glucuronidase; xylan  $\alpha$ -D-1,2-(4-O-methyl)glucuronohydrolase

**Systematic name:** xylan  $2-\alpha$ -D-(4-O-methyl)glucuronohydrolase

**References:** [1261]

[EC 3.2.1.131 created 1990]

# EC 3.2.1.132

Accepted name: chitosanase

**Reaction:** Endohydrolysis of  $\beta$ -(1 $\rightarrow$ 4)-linkages between D-glucosamine residues in a partly acetylated chitosan

**Systematic name:** chitosan *N*-acetylglucosaminohydrolase

**Comments:** A whole spectrum of chitosanases are now known (for more details, see

http://rbrzezinski.recherche.usherbrooke.ca/). They can hydrolyse various types of links in chitosan. The only constant property is the endohydrolysis of GlcN-GlcN links, which is common to all known chitosanases. One known chitosanase is limited to this link recognition [1808], while the majority can also recognize GlcN-GlcNAc links or GlcNAc-GlcN links but not both. They also do not

recognize GlcNAc-GlcNAc links in partly acetylated chitosan.

**References:** [755, 2496, 1286, 1808]

[EC 3.2.1.132 created 1990, modified 2004]

# EC 3.2.1.133

**Accepted name:** glucan 1,4-α-maltohydrolase

**Reaction:** hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkages in polysaccharides so as to remove successive  $\alpha$ -

maltose residues from the non-reducing ends of the chains

**Other name(s):** maltogenic  $\alpha$ -amylase; 1,4- $\alpha$ -D-glucan  $\alpha$ -maltohydrolase

**Systematic name:**  $4-\alpha$ -D-glucan  $\alpha$ -maltohydrolase

**Comments:** Acts on starch and related polysaccharides and oligosaccharides. The product is  $\alpha$ -maltose; cf. EC

3.2.1.2  $\beta$ -amylase.

**References:** [607, 2218]

[EC 3.2.1.133 created 1992, modified 1999]

# EC 3.2.1.134

**Accepted name:** difructose-anhydride synthase

**Reaction:** bis-D-fructose 2',1:2,1'-dianhydride +  $H_2O$  = inulobiose

**Other name(s):** inulobiose hydrolase

**Systematic name:** bis-D-fructose 2',1:2,1'-dianhydride fructohydrolase

**Comments:** Produces diffructose anhydride by the reverse reaction of partial hydrolysis, forming an  $\alpha$ -fructosidic

linkage.

**References:** [1841, 1842]

[EC 3.2.1.134 created 1992]

EC 3.2.1.135

Accepted name: neopullulanase

**Reaction:** Hydrolysis of pullulan to panose (6-α-D-glucosylmaltose)

Other name(s): pullulanase II

**Systematic name:** pullulan 4-D-glucanohydrolase (panose-forming)

**Comments:** *cf.* EC 3.2.1.41 (pullulanase ) and EC 3.2.1.57 (isopullulanase).

**References:** [1242]

[EC 3.2.1.135 created 1992]

EC 3.2.1.136

**Accepted name:** glucuronoarabinoxylan endo-1,4-β-xylanase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-xylosyl links in some glucuronoarabinoxylans

Other name(s): feraxan endoxylanase; feraxanase; endoarabinoxylanase; glucuronoxylan xylohydrolase; glu-

curonoxylanase; glucuronoxylan xylanohydrolase; glucuronoarabinoxylan 1,4-β-D-xylanohydrolase

**Systematic name:** glucuronoarabinoxylan 4-β-D-xylanohydrolase

Comments: High activity towards feruloylated arabinoxylans from cereal plant cell walls.

**References:** [2095]

[EC 3.2.1.136 created 1992]

EC 3.2.1.137

**Accepted name:** mannan exo-1,2-1,6-α-mannosidase

**Reaction:** Hydrolysis of  $(1\rightarrow 2)$ - $\alpha$ -D- and  $(1\rightarrow 6)$ - $\alpha$ -D- linkages in yeast mannan, releasing D-mannose

Other name(s): exo-1,2-1,6-α-mannosidase; 1,2-1,6-α-D-mannan D-mannohydrolase

**Systematic name:**  $(1\rightarrow 2)$ - $(1\rightarrow 6)$ - $\alpha$ -D-mannan D-mannohydrolase

**Comments:** Mannose residues linked  $\alpha$ -D-1,3- are also released, but very slowly.

**References:** [2856]

[EC 3.2.1.137 created 1992]

[3.2.1.138 Transferred entry. anhydrosialidase. Now EC 4.2.2.15, anhydrosialidase]

[EC 3.2.1.138 created 1992, deleted 2003]

EC 3.2.1.139

Accepted name: α-glucuronidase

**Reaction:** an  $\alpha$ -D-glucuronoside + H<sub>2</sub>O = an alcohol + D-glucuronate

Other name(s):  $\alpha$ -glucosiduronase

**Systematic name:** α-D-glucosiduronate glucuronohydrolase

**Comments:** Considerable differences in the specificities of the enzymes from different fungi for  $\alpha$ -D-

glucosiduronates have been reported. Activity is also found in the snail.

**References:** [2329, 3006]

[EC 3.2.1.139 created 1999]

# EC 3.2.1.140

Accepted name: lacto-N-biosidase

**Reaction:**  $\beta$ -D-Gal- $(1\rightarrow 3)$ - $\beta$ -D-GlcNAc- $(1\rightarrow 3)$ - $\beta$ -D-Gal- $(1\rightarrow 4)$ -D-Glc +  $H_2O = \beta$ -D-Gal- $(1\rightarrow 3)$ -D-GlcNAc +

 $\beta$ -D-Gal-(1 $\rightarrow$ 4)-D-Glc

**Systematic name:** oligosaccharide lacto-*N*-biosylhydrolase

**Comments:** The enzyme from *Streptomyces* specifically hydrolyses the terminal lacto-*N*-biosyl residue (β-D-Gal-

 $(1\rightarrow 3)$ -D-GlcNAc) from the non-reducing end of oligosaccharides with the structure  $\beta$ -D-Gal- $(1\rightarrow 3)$ - $\beta$ -D-GlcNAc- $(1\rightarrow 3)$ - $\beta$ -D-Gal- $(1\rightarrow R)$ . Lacto-N-hexaose  $(\beta$ -D-Gal- $(1\rightarrow 3)$ - $\beta$ -D-GlcNAc- $(1\rightarrow 3)$ - $\beta$ -D-Gal- $(1\rightarrow 4)$ -D-Glc) is hydrolysed to form first lacto-N-tetraose plus lacto-N-biose, with the subsequent formation of lactose. Oligosaccharides in which the non-reducing terminal Gal or the penultimate GlcNAc are replaced by fucose or sialic acid are not substrates. Asialo GM1 tetraose  $(\beta$ -D-Gal- $(1\rightarrow 3)$ - $\beta$ -D-GalNAc- $(1\rightarrow 3)$ - $\beta$ -D-Gal- $(1\rightarrow 4)$ -D-Glc) is hydrolysed very slowly, but lacto-N-neotetraose  $(\beta$ -D-Gal- $(1\rightarrow 4)$ - $\beta$ -D-GalNAc- $(1\rightarrow 3)$ - $\beta$ -D-Gal- $(1\rightarrow 4)$ -D-D-Gal- $(1\rightarrow 4)$ -D-Gal- $(1\rightarrow 4)$ -D-D-Gal- $(1\rightarrow 4)$ -D-D-Gal- $(1\rightarrow 4)$ -D-Gal- $(1\rightarrow 4)$ -D-Ga

Glc) is not a substrate

**References:** [2521, 2522]

[EC 3.2.1.140 created 1999]

#### EC 3.2.1.141

**Accepted name:**  $4-\alpha-D-(1\rightarrow 4)-\alpha-D$ -glucanotrehalose trehalohydrolase

**Reaction:** hydrolysis of  $(1\rightarrow 4)$ - $\alpha$ -D-glucosidic linkage in 4- $\alpha$ -D-[ $(1\rightarrow 4)$ - $\alpha$ -D-glucanosyl]<sub>n</sub> trehalose to yield

trehalose and  $(1\rightarrow 4)$ - $\alpha$ -D-glucan

**Other name(s):** malto-oligosyltrehalose trehalohydrolase

**Systematic name:**  $4-\alpha-D-[(1\rightarrow4)-\alpha-D-glucano]$  trehalose glucanohydrolase (trehalose-producing)

**References:** [1825, 2032, 2031]

[EC 3.2.1.141 created 1999]

# EC 3.2.1.142

Accepted name: limit dextrinase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic linkages in  $\alpha$ - and  $\beta$ -limit dextrins of amylopectin and glycogen,

and in amylopectin and pullulan

**Other name(s):** *R*-enzyme; amylopectin-1,6-glucosidase; dextrin  $\alpha$ -1,6-glucanohydrolase

**Systematic name:** dextrin 6-α-glucanohydrolase

**Comments:** Plant enzymes with little or no action on glycogen. Action on amylopectin is incomplete, but action

on  $\alpha$ -limit dextrins is complete. Maltose is the smallest sugar it can release from an  $\alpha$ -(1 $\rightarrow$ 6)-linkage.

**References:** [958, 1801]

[EC 3.2.1.142 created 2000]

# EC 3.2.1.143

**Accepted name:** poly(ADP-ribose) glycohydrolase

**Reaction:** hydrolyses poly(ADP-D-ribose) at glycosidic (1"-2') linkage of ribose-ribose bond to produce free

ADP-D-ribose

**Comments:** Specific to (1''-2') linkage of ribose-ribose bond of poly(ADP-D-ribose).

**References:** [1935, 1694]

[EC 3.2.1.143 created 2000]

#### EC 3.2.1.144

**Accepted name:** 3-deoxyoctulosonase

**Reaction:** 3-deoxyoctulosonyl-lipopolysaccharide + H<sub>2</sub>O = 3-deoxyoctulosonic acid + lipopolysaccharide

Other name(s):  $\alpha$ -Kdo-ase

**Systematic name:** 3-deoxyoctulosonyl-lipopolysaccharide hydrolase

**Comments:** Releases Kdo (α- and β-linked 3-deoxy-D-manno-octulosonic acid) from different lipopolysaccha-

rides, including Re-LPS from Escherichia coli and Salmonella, Rd-LPS from S. minnesota, and de-O-

acyl-re-LPS. 4-Methylumbelliferyl- $\alpha$ -Kdo ( $\alpha$ -Kdo-OMec) is also a substrate.

**References:** [1676]

[EC 3.2.1.144 created 2000]

EC 3.2.1.145

**Accepted name:** galactan 1,3-β-galactosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $\beta$ -D-galactose residues in  $(1\rightarrow 3)$ - $\beta$ -D-galactopyranans

Other name(s): galactan  $(1\rightarrow 3)$ - $\beta$ -D-galactosidase Systematic name: galactan 3- $\beta$ -D-galactosidase

**Comments:** This enzyme removes not only free galactose, but also 6-glycosylated residues, e.g.,  $(1\rightarrow 6)$ - $\beta$ -D-

galactobiose, and galactose bearing oligosaccharide chains on O-6. Hence, it releases branches from

[arabino-galacto- $(1\rightarrow 6)$ ]- $(1\rightarrow 3)$ - $\beta$ -D-galactans.

**References:** [2983, 2254]

[EC 3.2.1.145 created 2001]

EC 3.2.1.146

**Accepted name:** β-galactofuranosidase

**Reaction:** Hydrolysis of terminal non-reducing  $\beta$ -D-galactofuranosides, releasing galactose **Other name(s):** exo- $\beta$ -galactofuranosidase; exo- $\beta$ -D-galactofuranosidase

**Systematic name:**  $\beta$ -D-galactofuranoside hydrolase

Comments: The enzyme from *Helminthosporium sacchari* detoxifies helminthosporoside, a

bis(digalactosyl)terpene produced by this fungus, by releasing its four molecules of bound galactose.

**References:** [2425, 530, 498, 1910]

[EC 3.2.1.146 created 2001]

EC 3.2.1.147

Accepted name: thioglucosidase

**Reaction:** a thioglucoside  $+ H_2O = a$  sugar + a thiol **Other name(s):** myrosinase; sinigrinase; sinigrase **Systematic name:** thioglucoside glucohydrolase

**Comments:** Has a wide specificity for thioglycosides.

**References:** [956, 2281]

[EC 3.2.1.147 created 1972 as EC 3.2.3.1, transferred 2001 to EC 3.2.1.147]

[3.2.1.148 Deleted entry. ribosylhomocysteinase. This enzyme was transferred to EC 3.13.1.2, 5-deoxyribos-5-ylhomocysteinase, which has since been deleted. The activity is most probably attributable to EC 4.4.1.21, S-ribosylhomocysteine lyase]

[EC 3.2.1.148 created 1972 as EC 3.3.1.3, transferred 2001 to EC 3.2.1.148, deleted 2004]

EC 3.2.1.149

**Accepted name:** β-primeverosidase

**Reaction:** a 6-O-( $\beta$ -D-xylopyranosyl)- $\beta$ -D-glucopyranoside + H<sub>2</sub>O = 6-O-( $\beta$ -D-xylopyranosyl)- $\beta$ -D-

glucopyranose + an alcohol

**Systematic name:** 6-*O*-(β-D-xylopyranosyl)-β-D-glucopyranoside 6-*O*-(β-D-xylosyl)-β-D-glucohydrolase

**Comments:** The enzyme is responsible for the formation of the alcoholic aroma in oolong and black tea. In ad-

dition to  $\beta$ -primeverosides [i.e. 6-O-( $\beta$ -D-xylopyranosyl)- $\beta$ -D-glucopyranosides], it also hydrolyses 6-O-( $\beta$ -D-apiofuranosyl)- $\beta$ -D-glucopyranosides and, less rapidly,  $\beta$ -vicianosides and 6-O-( $\alpha$ -L-arabinofuranosyl)- $\beta$ -D-glucopyranosides, but not  $\beta$ -glucosides. Geranyl-, linaloyl-, benzyl- and p-

nitrophenol glycosides are all hydrolysed.

**References:** [1227, 2149]

[EC 3.2.1.149 created 2001]

EC 3.2.1.150

**Accepted name:** oligoxyloglucan reducing-end-specific cellobiohydrolase

**Reaction:** Hydrolysis of cellobiose from the reducing end of xyloglucans consisting of a  $(1\rightarrow 4)$ - $\beta$ -linked glu-

can carrying  $\alpha$ -D-xylosyl groups on O-6 of the glucose residues. To be a substrate, the first residue must be unsubstituted, the second residue may bear a xylosyl group, whether further glycosylated or not, and the third residue, which becomes the new terminus by the action of the enzyme, is preferably

xylosylated, but this xylose residue must not be further substituted.

Systematic name: oligoxyloglucan reducing-end cellobiohydrolase

**Comments:** The enzyme is found in the fungus *Geotrichum* sp. M128. The substrate is a hemicellulose found in

plant cell walls.

**References:** [3266]

[EC 3.2.1.150 created 2003]

EC 3.2.1.151

**Accepted name:** xyloglucan-specific endo-β-1,4-glucanase

**Reaction:**  $xyloglucan + H_2O = xyloglucan oligosaccharides$ 

Other name(s): XEG; xyloglucan endo-β-1,4-glucanase; xyloglucanese; xyloglucanendohydrolase; XH; 1,4-β-D-

glucan glucanohydrolase

**Systematic name:**  $[(1\rightarrow 6)-\alpha-D-xylo]-(1\rightarrow 4)-\beta-D-glucan glucanohydrolase$ 

**Comments:** The enzyme from *Aspergillus aculeatus* is specific for xyloglucan and does not hydrolyse other cell-

wall components. The reaction involves endohydrolysis of 1,4-β-D-glucosidic linkages in xyloglucan

with retention of the  $\beta$ -configuration of the glycosyl residues.

**References:** [2248, 985]

[EC 3.2.1.151 created 2003]

EC 3.2.1.152

**Accepted name:** mannosylglycoprotein endo-β-mannosidase

 $\textbf{Reaction:} \quad \text{Hydrolysis of the } \alpha\text{-D-mannosyl-}(1\rightarrow 6)\text{-}\beta\text{-D-mannosyl-}(1\rightarrow 4)\text{-}N\text{-acetyl-}\beta\text{-D-glucosaminyl-}(1\rightarrow 4)\text{-}N\text{-acetyl-}\beta\text{-}$ 

acetyl- $\beta$ -D-glucosaminyl sequence of glycoprotein to  $\alpha$ -D-mannosyl- $(1\rightarrow 6)$ -D-mannose and N-acetyl-

 $\beta\text{-D-glucosaminyl-}(1{\rightarrow}4)\text{--}\textit{N-acetyl-}\beta\text{-D-glucosaminyl sequences}$ 

Other name(s): endo- $\beta$ -mannosidase

**Comments:** The substrate group is a substituent on N-4 of an asparagine residue in the glycoprotein. The mannose

residue at the non-reducing end of the sequence may carry further  $\alpha$ -D-mannosyl groups on O-3 or O-6, but such a substituent on O-3 of the  $\beta$ -D-mannosyl group prevents the action of the enzyme. The

enzyme was obtained from the lily, Lilium longiflorum.

**References:** [1263, 2533]

[EC 3.2.1.152 created 2005]

EC 3.2.1.153

**Accepted name:** fructan  $\beta$ -(2,1)-fructosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $(2\rightarrow 1)$ -linked  $\beta$ -D-fructofuranose residues in fructans

Other name(s):  $\beta$ -(2-1)-D-fructan fructohydrolase;  $\beta$ -(2-1)fructan exohydrolase; inulinase; 1-FEH II; 1-fructan exo-

hydrolase; 1-FEH w1; 1-FEH w2; β-(2-1)-linkage-specific fructan-β-fructosidase; β-(2,1)-D-fructan

fructohydrolase

**Systematic name:**  $\beta$ -(2 $\rightarrow$ 1)-D-fructan fructohydrolase

**Comments:** Possesses one of the activities of EC 3.2.1.80, fructan  $\beta$ -fructosidase. While the best substrates are

the inulin-type fructans, such as 1-kestose [ $\beta$ -D-fructofuranosyl-( $2\rightarrow 1$ )- $\beta$ -D-fructofuranosyl  $\alpha$ -D-glucopyranoside], some (but not all) levan-type fructans can also be hydrolysed, but more slowly [see EC 3.2.1.154, fructan  $\beta$ -(2,6)-fructosidase]. Sucrose, while being a very poor

substrate, can substantially inhibit enzyme activity in some cases.

**References:** [2457, 579]

[EC 3.2.1.153 created 2005]

EC 3.2.1.154

**Accepted name:** fructan  $\beta$ -(2,6)-fructosidase

**Reaction:** Hydrolysis of terminal, non-reducing  $(2\rightarrow 6)$ -linked  $\beta$ -D-fructofuranose residues in fructans

Other name(s):  $\beta$ -(2-6)-fructan exohydrolase; levanase; 6-FEH;  $\beta$ -(2,6)-D-fructan fructohydrolase

**Systematic name:**  $(2\rightarrow 6)$ - $\beta$ -D-fructan fructohydrolase

Comments: Possesses one of the activities of EC 3.2.1.80, fructan  $\beta$ -fructosidase. While the best substrates are

the levan-type fructans such as 6-kestotriose [ $\beta$ -D-fructofuranosyl-( $2\rightarrow 6$ )- $\beta$ -D-fructofuranosyl  $\alpha$ -D-glucopyranoside], some (but not all) inulin-type fructans can also be hydrolysed, but more slowly [cf. EC 3.2.1.153, fructan  $\beta$ -(2,1)-fructosidase]. Sucrose, while being a very poor

substrate, can substantially inhibit enzyme activity in some cases.

**References:** [1829, 580, 1119]

[EC 3.2.1.154 created 2005]

EC 3.2.1.155

**Accepted name:** xyloglucan-specific exo-β-1,4-glucanase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ -D-glucosidic linkages in xyloglucans so as to successively remove oligosaccha-

rides from the chain end.

Other name(s): Cel74A

**Systematic name:**  $[(1\rightarrow 6)-\alpha-D-xylo]-(1\rightarrow 4)-\beta-D-glucan$  exo-glucohydrolase

Comments: The enzyme removes XXXG heptasaccharides, XXLG/XLXG octasaccharides and XLLG nonasac-

charides from the end of tamarind seed xyloglucan polymers in a processive manner. Hydrolysis occurs at the unsubstituted D-glucopyranose residue in the main backbone. It is not known whether the cleavage takes place at the reducing or non-reducing end of the polymer. Very low activity with  $\beta$ -D-glucans. The enzyme from *Chrysosporium lucknowense* shifts to an endoglucanase mode when acting on linear substrates without bulky substituents on the polymeric backbone such as barley  $\beta$ -glucan.

**References:** [985]

[EC 3.2.1.155 created 2005, withdrawn at public-review stage, modified and reinstated 2006, modified 2011]

EC 3.2.1.156

**Accepted name:** oligosaccharide reducing-end xylanase

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-xylose residues from the reducing end of oligosaccharides

**Other name(s):** Rex; reducing end xylose-releasing exo-oligoxylanase

**Systematic name:**  $\beta$ -D-xylopyranosyl- $(1\rightarrow 4)$ - $\beta$ -D-xylopyranose reducing-end xylanase

**Comments:** The enzyme, originally isolated from the bacterium *Bacillus halodurans* C-125, releases the xylose

unit at the reducing end of oligosaccharides ending with the structure  $\beta$ -D-xylopyranosyl- $(1\rightarrow 4)$ - $\beta$ -D-xylopyranosyl- $(1\rightarrow 4)$ - $\beta$ -D-xylopyranose, leaving the new reducing end in the  $\alpha$  configuration. It is specific for the  $\beta$  anomers of xylooligosaccharides whose degree of polymerization is equal to or greater than 3. The penultimate residue must be  $\beta$ -D-xylopyranose, but replacing either of the flanking

residues with glucose merely slows the rate greatly.

**References:** [1166, 865]

[EC 3.2.1.156 created 2005]

EC 3.2.1.157

Accepted name: 1-carrageenase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-linkages between D-galactose 4-sulfate and 3,6-anhydro-D-galactose-

2-sulfate in t-carrageenans

**Systematic name:** 1-carrageenan 4-β-D-glycanohydrolase (configuration-inverting)

Comments: The main products of hydrolysis are t-neocarratetraose sulfate and t-neocarrahexaose sulfate. t-

Neocarraoctaose is the shortest substrate oligomer that can be cleaved. Unlike EC 3.2.1.81,  $\beta$ -agarase and EC 3.2.1.83,  $\kappa$ -carrageenase, this enzyme proceeds with inversion of the anomeric configuration. 1-Carrageenan differs from  $\kappa$ -carrageenan by possessing a sulfo group on O-2 of the 3,6-anhydro-D-galactose residues, in addition to that present in the  $\kappa$ -compound on O-4 of the D-galactose residues.

**References:** [137, 1904, 1905]

[EC 3.2.1.157 created 2006]

EC 3.2.1.158

**Accepted name:** α-agarase

**Reaction:** Endohydrolysis of  $(1\rightarrow 3)-\alpha$ -L-galactosidic linkages in agarose, yielding agarotetraose as the major

product

Other name(s): agarase (ambiguous); agaraseA33

Systematic name: agarose 3-glycanohydrolase

**Comments:** Requires Ca<sup>2+</sup>. The enzyme from *Thalassomonas* sp. can use agarose, agarohexaose and neoa-

garohexaose as substrate. The products of agarohexaose hydrolysis are dimers and tetramers, with agarotetraose being the predominant product, whereas hydrolysis of neoagarohexaose gives rise to two types of trimer. While the enzyme can also hydrolyse the highly sulfated agarose porphyran very efficiently, it cannot hydrolyse the related compounds  $\kappa$ -carrageenan (see EC 3.2.1.83) and 1-

carrageenan (see EC 3.2.1.157) [2170]. See also EC 3.2.1.81, β-agarase.

**References:** [2317, 2170]

[EC 3.2.1.158 created 2006]

EC 3.2.1.159

**Accepted name:** α-neoagaro-oligosaccharide hydrolase

**Reaction:** Hydrolysis of the  $(1\rightarrow 3)$ - $\alpha$ -L-galactosidic linkages of neoagaro-oligosaccharides that are smaller than

a hexamer, yielding 3,6-anhydro-L-galactose and D-galactose

**Other name(s):** α-neoagarooligosaccharide hydrolase; α-NAOS hydrolase **Systematic name:** α-neoagaro-oligosaccharide 3-glycohydrolase

**Comments:** When neoagarohexaose is used as a substrate, the oligosaccharide is cleaved at the non-reducing end

to produce 3,6-anhydro-L-galactose and agaropentaose, which is further hydrolysed to agarobiose and agarotriose. With neoagarotetraose as substrate, the products are predominantly agarotriose and 3,6-anhydro-L-galactose. In *Vibrio* sp. the actions of EC 3.2.1.81,  $\beta$ -agarase and EC 3.2.1.159 can be used

to degrade agarose to 3,6-anhydro-L-galactose and D-galactose.

**References:** [2791]

[EC 3.2.1.159 created 2006]

[3.2.1.160 Deleted entry. xyloglucan-specific exo- $\beta$ -1,4-glucanase. The enzyme was shown to be identical to EC 3.2.1.155, xyloglucan-specific exo- $\beta$ -1,4-glucanase, during the public-review process so was withdrawn before being made official]

#### [EC 3.2.1.160 created 2006, deleted 2006]

# EC 3.2.1.161

**Accepted name:**  $\beta$ -apiosyl- $\beta$ -glucosidase

**Reaction:**  $7-[\beta-D-apiofuranosyl-(1\rightarrow 6)-\beta-D-glucopyranosyloxy]isoflavonoid + H<sub>2</sub>O = a 7-hydroxyisoflavonoid$ 

+  $\beta$ -D-apiofuranosyl- $(1\rightarrow 6)$ -D-glucose

Other name(s): isoflavonoid-7-O- $\beta$ [D-apiosyl-(1 $\rightarrow$ 6)- $\beta$ -D-glucoside] disaccharidase; isoflavonoid 7-O- $\beta$ -apiosyl-

glucoside β-glucosidase; furcatin hydrolase

**Systematic name:** 7-[ $\beta$ -D-apiofuranosyl-( $1\rightarrow 6$ )- $\beta$ -D-glucopyranosyloxy]isoflavonoid  $\beta$ -D-apiofuranosyl-( $1\rightarrow 6$ )-D-

glucohydrolase

**Comments:** The enzyme from the tropical tree *Dalbergia nigrescens* Kurz belongs in glycosyl hydrolase family

1. The enzyme removes disaccharides from the natural substrates dalpatein 7-O- $\beta$ -D-apiofuranosyl- $(1\rightarrow 6)$ - $\beta$ -D-glucopyranoside and 7-hydroxy-2',4',5',6-tetramethoxy-7-O- $\beta$ -D-apiofuranosyl- $(1\rightarrow 6)$ - $\beta$ -D-glucopyranoside (dalnigrein 7-O- $\beta$ -D-apiofuranosyl- $(1\rightarrow 6)$ - $\beta$ -D-glucopyranoside) although it can also remove a single glucose residue from isoflavonoid 7-O-glucosides [445]. Daidzin and genistin

are also substrates.

**References:** [1181, 445, 23]

[EC 3.2.1.161 created 2006]

# EC 3.2.1.162

**Accepted name:** λ-carrageenase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -linkages in the backbone of  $\lambda$ -carrageenan, resulting in the tetrasaccha-

ride  $\alpha$ -D-Galp2,6 $S_2$ -(1 $\rightarrow$ 3)- $\beta$ -D-Galp2S-(1 $\rightarrow$ 4)- $\alpha$ -D-Galp2,6 $S_2$ -(1 $\rightarrow$ 3)-D-Galp2S

Other name(s): endo- $\beta$ -1,4-carrageenose 2,6,2'-trisulfate-hydrolase Systematic name: endo- $(1\rightarrow 4)$ - $\beta$ -carrageenose 2,6,2'-trisulfate-hydrolase

Comments: The enzyme from *Pseudoalteromonas* sp. is specific for  $\lambda$ -carrageenan. 1-Carrageenan (see EC

3.2.1.157, t-carrageenase), κ-carrageenan (see EC 3.2.1.83, κ-carrageenase), agarose and porphyran

are not substrates.

**References:** [2169]

[EC 3.2.1.162 created 2007]

# EC 3.2.1.163

**Accepted name:** 1,6-α-D-mannosidase

**Reaction:** Hydrolysis of the  $(1\rightarrow 6)$ -linked  $\alpha$ -D-mannose residues in  $\alpha$ -D-Manp- $(1\rightarrow 6)$ -D-Manp

**Systematic name:**  $(1\rightarrow 6)$ - $\alpha$ -mannosyl  $\alpha$ -D-mannohydrolase

**Comments:** The enzyme is specific for  $(1 \rightarrow 6)$ -linked mannobiose and has no activity towards any other linkages,

or towards p-nitrophenyl- $\alpha$ -D-mannopyranoside or baker's yeast mannan. It is strongly inhibited by

Mn<sup>2+</sup> but does not require Ca<sup>2+</sup> or any other metal cofactor for activity.

**References:** [92]

[EC 3.2.1.163 created 2007]

#### EC 3.2.1.164

**Accepted name:** galactan endo-1,6-β-galactosidase

**Reaction:** Endohydrolysis of  $(1\rightarrow 6)$ - $\beta$ -D-galactosidic linkages in arabinogalactan proteins and  $(1\rightarrow 3)$ : $(1\rightarrow 6)$ - $\beta$ -

galactans to yield galactose and  $(1\rightarrow 6)$ - $\beta$ -galactobiose as the final products

Other name(s): endo-1,6- $\beta$ -galactanase Systematic name: endo- $\beta$ -(1 $\rightarrow$ 6)-galactanase **Comments:** The enzyme specifically hydrolyses 1,6-β-D-galactooligosaccharides with a degree of polymerization

(DP) higher than 3, and their acidic derivatives with 4-O-methylglucosyluronate or glucosyluronate groups at the non-reducing terminals [2188]. 1,3- $\beta$ -D- and 1,4- $\beta$ -D-galactosyl residues cannot act as substrates. The enzyme can also hydrolyse  $\alpha$ -L-arabinofuranosidase-treated arabinogalactan protein (AGP) extracted from radish roots [2188, 1519]. AGPs are thought to be involved in many physiologi-

cal events, such as cell division, cell expansion and cell death [1519].

**References:** [293, 2188, 1519]

[EC 3.2.1.164 created 2007]

EC 3.2.1.165

**Accepted name:** exo-1,4-β-D-glucosaminidase

Reaction: Hydrolysis of chitosan or chitosan oligosaccharides to remove successive D-glucosamine residues

from the non-reducing termini

Other name(s): CsxA; GlcNase; exochitosanase; GlmA; exo-β-D-glucosaminidase; chitosan exo-1,4-β-D-

glucosaminidase

**Systematic name:** chitosan exo- $(1\rightarrow 4)$ - $\beta$ -D-glucosaminidase

**Comments:** Chitosan is a partially or totally *N*-deacetylated chitin derivative that is found in the cell walls of some

phytopathogenic fungi and comprises D-glucosamine residues with a variable content of GlcNAc residues [496]. Acts specifically on chitooligosaccharides and chitosan, having maximal activity on chitotetraose, chitopentaose and their corresponding alcohols [2055]. The enzyme can degrade GlcN-GlcNAc but not GlcNAc-GlcNAc [853]. A member of the glycoside hydrolase family 2 (GH-2) [496].

**References:** [2055, 2104, 853, 496, 1228]

[EC 3.2.1.165 created 2008]

EC 3.2.1.166

**Accepted name:** heparanase

**Reaction:** endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-glycosidic bonds of heparan sulfate chains in heparan sulfate proteo-

glycan

Other name(s): Hpa1 heparanase; Hpa1; heparanase 1; heparanase-1; C1A heparanase; HPSE

**Systematic name:** heparan sulfate *N*-sulfo-D-glucosamine endoglucanase

**Comments:** Heparanase cleaves the linkage between a glucuronic acid unit and an N-sulfo glucosamine unit car-

rying either a 3-*O*-sulfo or a 6-*O*-sulfo group [2265]. Heparanase-1 cuts macromolecular heparin into fragments of 5000–20000 Da [3080]. The enzyme cleaves the heparan sulfate glycosamino-glycans from proteoglycan core proteins and degrades them to small oligosaccharides. Inside cells, the enzyme is important for the normal catabolism of heparan sulfate proteoglycans, generating glycosaminoglycan fragments that are then transported to lysosomes and completely degraded. When secreted, heparanase degrades basement membrane heparan sulfate glycosaminoglycans at sites of injury or inflammation, allowing extravasion of immune cells into nonvascular spaces and releasing

factors that regulate cell proliferation and angiogenesis [133].

**References:** [133, 2265, 2282, 2183, 3080, 955, 2956, 1900, 1038]

[EC 3.2.1.166 created 2010]

EC 3.2.1.167

**Accepted name:** baicalin-β-D-glucuronidase

**Reaction:** baicalin +  $H_2O$  = baicalein + D-glucuronate

Other name(s): baicalinase

**Systematic name:** 5,6,7-trihydroxyflavone-7-*O*-β-D-glucupyranosiduronate glucuronosylhydrolase

**Comments:** The enzyme also hydrolyses wogonin 7-*O*-β-D-glucuronide and oroxylin 7-*O*-β-D-glucuronide with

lower efficiency [1976]. Neglegible activity with *p*-nitrophenyl-β-D-glucuronide [3324].

**References:** [1229, 3324, 2534, 1976]

# [EC 3.2.1.167 created 2011]

EC 3.2.1.168

**Accepted name:** hesperidin 6-O- $\alpha$ -L-rhamnosyl- $\beta$ -D-glucosidase

**Reaction:** hesperidin +  $H_2O$  = hesperetin + rutinose

Systematic name: hesperetin 7-(6-O- $\alpha$ -L-rhamnopyranosyl- $\beta$ -D-glucopyranoside) 6-O- $\alpha$ -rhamnopyranosyl- $\beta$ -

glucohydrolase

**Comments:** The enzyme exhibits high specificity towards 7-O-linked flavonoid  $\beta$ -rutinosides.

**References:** [1859, 1860]

[EC 3.2.1.168 created 2011]

EC 3.2.1.169

**Accepted name:** protein *O*-GlcNAcase

**Reaction:** (1) [protein]-3-O-(N-acetyl- $\beta$ -D-glucosaminyl)-L-serine + H<sub>2</sub>O = [protein]-L-serine + N-acetyl-D-

glucosamine

(2) [protein]-3-O-(N-acetyl- $\beta$ -D-glucosaminyl)-L-theronine +  $H_2O$  = [protein]-L-threonine + N-acetyl-

D-glucosamine

**Other name(s):** OGA; glycoside hydrolase *O*-GlcNAcase; *O*-GlcNAcase; BtGH84; *O*-GlcNAc hydrolase

**Systematic name:** [protein]-3-O-(N-acetyl- $\beta$ -D-glucosaminyl)-L-serine/threonine N-acetylglucosaminyl hydrolase **Comments:** Within higher eukaryotes post-translational modification of protein serines/threonines with N-

acetylglucosamine (*O*-GlcNAc) is dynamic, inducible and abundant, regulating many cellular processes by interfering with protein phosphorylation. EC 2.4.1.255 (protein *O*-GlcNAc transferase)

transfers GlcNAc onto substrate proteins and EC 3.2.1.169 (protein *O*-GlcNAcase) cleaves GlcNAc

from the modified proteins.

**References:** [873, 3160, 387, 584, 1447, 633]

[EC 3.2.1.169 created 2011]

EC 3.2.1.170

**Accepted name:** mannosylglycerate hydrolase

**Reaction:** 2-O-( $\alpha$ -D-mannopyranosyl)-D-glycerate + H<sub>2</sub>O = D-mannopyranose + D-glycerate

Other name(s): MgH

**Systematic name:** 2-O-( $\alpha$ -D-mannopyranosyl)-D-glycerate D-mannohydrolase

Comments: The enzyme occurs in thermophilic bacteria and has been characterized in *Thermus thermophilus* and

Rubrobacter radiotolerans. It also has been identified in the moss Selaginella moellendorffii.

**References:** [28, 2102]

[EC 3.2.1.170 created 2011, modified 2018]

EC 3.2.1.171

Accepted name: rhamnogalacturonan hydrolase

**Reaction:** Endohydrolysis of  $\alpha$ -D-GalA- $(1\rightarrow 2)$ - $\alpha$ -L-Rha glycosidic bond in the rhamnogalacturonan I backbone

with initial inversion of anomeric configuration releasing oligosaccharides with β-D-GalA at the re-

ducing end.

**Other name(s):** rhamnogalacturonase A; RGase A; RG-hydrolase

**Systematic name:** rhamnogalacturonan  $\alpha$ -D-GalA- $(1\rightarrow 2)$ - $\alpha$ -L-Rha hydrolase

**Comments:** The enzyme is part of the degradation system for rhamnogalacturonan I in *Aspergillus aculeatus*.

**References:** [2264, 1494, 107, 2263, 2289]

[EC 3.2.1.171 created 2011]

Accepted name: unsaturated rhamnogalacturonyl hydrolase

**Reaction:** 2-O-(4-deoxy- $\beta$ -L-threo-hex-4-enopyranuronosyl)- $\alpha$ -L-rhamnopyranose + H<sub>2</sub>O = 5-dehydro-4-

deoxy-D-glucuronate + L-rhamnopyranose

Other name(s): YteR; YesR

Systematic name: 2-O-(4-deoxy-β-L-threo-hex-4-enopyranuronosyl)-α-L-rhamnopyranose hydrolase

**Comments:** The enzyme is part of the degradation system for rhamnogalacturonan I in *Bacillus subtilis* strain 168.

**References:** [1274, 3330, 1275]

[EC 3.2.1.172 created 2011, modified 2012]

EC 3.2.1.173

Accepted name: rhamnogalacturonan galacturonohydrolase

**Reaction:** Exohydrolysis of the  $\alpha$ -D-GalA- $(1\rightarrow 2)$ - $\alpha$ -L-Rha bond in rhamnogalacturonan oligosaccharides with

initial inversion of configuration releasing D-galacturonic acid from the non-reducing end of rhamno-

galacturonan oligosaccharides.

**Other name(s):** RG-galacturonohydrolase

Systematic name: rhamnogalacturonan oligosaccharide  $\alpha$ -D-GalA- $(1\rightarrow 2)$ - $\alpha$ -L-Rha galacturonohydrolase

**Comments:** The enzyme is part of the degradation system for rhamnogalacturonan I in *Aspergillus aculeatus*.

**References:** [2012]

[EC 3.2.1.173 created 2011]

EC 3.2.1.174

**Accepted name:** rhamnogalacturonan rhamnohydrolase

**Reaction:** Exohydrolysis of the  $\alpha$ -L-Rha- $(1\rightarrow 4)$ - $\alpha$ -D-GalA bond in rhamnogalacturonan oligosaccharides with

initial inversion of configuration releasing β-L-rhamnose from the non-reducing end of rhamnogalac-

turonan oligosaccharides.

**Other name(s):** RG-rhamnohydrolase; RG α-L-rhamnopyranohydrolase

**Systematic name:** rhamnogalacturonan oligosaccharide  $\alpha$ -L-Rha- $(1\rightarrow 4)$ - $\alpha$ -D-GalA rhamnohydrolase

**Comments:** The enzyme is part of the degradation system for rhamnogalacturonan I in *Aspergillus aculeatus*.

**References:** [2289, 2013]

[EC 3.2.1.174 created 2011]

EC 3.2.1.175

**Accepted name:** β-D-glucopyranosyl abscisate β-glucosidase

**Reaction:** D-glucopyranosyl abscisate  $+ H_2O = D$ -glucose + abscisate

Other name(s): AtBG1; ABA-β-D-glucosidase; ABA-specific β-glucosidase; ABA-GE hydrolase; β-D-

glucopyranosyl abscisate hydrolase

**Systematic name:** β-D-glucopyranosyl abscisate glucohydrolase

 $\textbf{Comments:} \quad \text{The enzyme hydrolzes the biologically inactive } \beta\text{-D-glucopyranosyl ester of abscisic acid to produce}$ 

active abscisate. Abscisate is a phytohormone critical for plant growth, development and adaption to

various stress conditions. The enzyme does not hydrolyse  $\beta$ -D-glucopyranosyl zeatin [1624].

**References:** [1624, 1403, 610]

[EC 3.2.1.175 created 2011]

EC 3.2.1.176

**Accepted name:** cellulose 1,4-β-cellobiosidase (reducing end)

**Reaction:** Hydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-glucosidic linkages in cellulose and similar substrates, releasing cellobiose

from the reducing ends of the chains.

Other name(s): CelS; CelSS; endoglucanase SS; cellulase SS; cellobiohydrolase CelS; Cel48A

**Systematic name:** 4-β-D-glucan cellobiohydrolase (reducing end)

**Comments:** Some exocellulases, most of which belong to the glycoside hydrolase family 48 (GH48, formerly

known as cellulase family L), act at the reducing ends of cellulose and similar substrates. The CelS enzyme from *Clostridium thermocellum* is the most abundant subunit of the cellulosome formed by the organism. It liberates cellobiose units from the reducing end by hydrolysis of the glycosidic bond, employing an inverting reaction mechanism [2493]. Different from EC 3.2.1.91, which attacks cellu-

lose from the non-reducing end.

**References:** [150, 2493]

[EC 3.2.1.176 created 2011]

EC 3.2.1.177

**Accepted name:** α-D-xyloside xylohydrolase

**Reaction:** Hydrolysis of terminal, non-reducing  $\alpha$ -D-xylose residues with release of  $\alpha$ -D-xylose.

Other name(s):  $\alpha$ -xylosidase

**Systematic name:**  $\alpha$ -D-xyloside xylohydrolase

**Comments:** The enzyme catalyses hydrolysis of a terminal, unsubstituted xyloside at the extreme reducing end of

a xylogluco-oligosaccharide. Representative  $\alpha$ -xylosidases from glycoside hydrolase family 31 utilize a two-step (double-displacement) mechanism involving a covalent glycosyl-enzyme intermediate, and

retain the anomeric configuration of the product.

**References:** [1961, 2514, 505, 1741, 1222, 2190, 1604]

[EC 3.2.1.177 created 2011]

EC 3.2.1.178

**Accepted name:** β-porphyranase

**Reaction:** Hydrolysis of  $\beta$ -D-galactopyranose- $(1\rightarrow 4)$ - $\alpha$ -L-galactopyranose-6-sulfate linkages in porphyran

**Other name(s):** porphyranase; PorA; PorB; endo-β-porphyranase

**Systematic name:** porphyran  $\beta$ -D-galactopyranose- $(1\rightarrow 4)$ - $\alpha$ -L-galactopyranose-6-sulfate 4-glycanohydrolase

Comments: The backbone of porphyran consists largely (70%) of (1 $\rightarrow$ 3)-linked  $\beta$ -D-galactopyranose fol-

lowed by  $(1\rightarrow 4)$ -linked  $\alpha$ -L-galactopyranose-6-sulfate [the other 30% are mostly agarobiose repeating units of  $(1\rightarrow 3)$ -linked  $\beta$ -D-galactopyranose followed by  $(1\rightarrow 4)$ -linked 3,6-anhydro- $\alpha$ -L-galactopyranose] [489]. This enzyme cleaves the  $(1\rightarrow 4)$  linkages between  $\beta$ -D-galactopyranose and  $\alpha$ -L-galactopyranose-6-sulfate, forming mostly the disaccharide  $\alpha$ -L-galactopyranose-6-sulfate- $(1\rightarrow 3)$ - $\beta$ -D-galactose, although some longer oligosaccharides of even number of residues are also observed. Since the enzyme is inactive on the non-sulfated agarose portion of the porphyran backbone, some agarose fragments are also included in the products [1098]. Methylation of the D-galactose prevents its binding at position -1 [489].

**References:** [1098, 489]

[EC 3.2.1.178 created 2011]

EC 3.2.1.179

**Accepted name:** gellan tetrasaccharide unsaturated glucuronosyl hydrolase

**Reaction:**  $\beta$ -D-4-deoxy- $\Delta^4$ -GlcAp-(1 $\rightarrow$ 4)- $\beta$ -D-Glcp-(1 $\rightarrow$ 4)- $\alpha$ -L-Rhap-(1 $\rightarrow$ 3)-D-Glcp + H<sub>2</sub>O = 5-dehydro-4-

deoxy-D-glucuronate +  $\beta$ -D-Glcp-(1 $\rightarrow$ 4)- $\alpha$ -L-Rhap-(1 $\rightarrow$ 3)-D-Glcp

Other name(s): UGL (ambiguous); unsaturated glucuronyl hydrolase (ambiguous); gellan tetrasaccharide unsaturated

glucuronyl hydrolase

Systematic name:  $\beta$ -D-4-deoxy- $\Delta^4$ -GlcAp-(1 $\rightarrow$ 4)- $\beta$ -D-Glcp-(1 $\rightarrow$ 4)- $\alpha$ -L-Rhap-(1 $\rightarrow$ 3)-D-Glcp-D-4-deoxy- $\Delta^4$ -GlcAp-GlcAp-

hydrolase

**Comments:** The enzyme releases 4-deoxy-4(5)-unsaturated D-glucuronic acid from oligosaccharides produced by

polysaccharide lyases, e.g. the tetrasaccharide β-D-4-deoxy- $\Delta^4$ -GlcAp-(1 $\rightarrow$ 4)-β-D-Glcp-(1 $\rightarrow$ 4)-α-L-Rhap-(1 $\rightarrow$ 3)-D-Glcp produced by EC 4.2.2.25, gellan lyase. The enzyme can also hydrolyse unsaturated chondroitin and hyaluronate disaccharides (β-D-4-deoxy- $\Delta^4$ -GlcAp-(1 $\rightarrow$ 3)-D-GalNAc, β-D-4-deoxy- $\Delta^4$ -GlcAp-(1 $\rightarrow$ 3)-D-GalNAc, β-D-4-deoxy- $\Delta^4$ -GlcAp-(1 $\rightarrow$ 3)-D-GlcNAc), preferring the unsulfated disaccharides to the sulfated disaccharides.

**References:** [1272, 1070, 1273]

[EC 3.2.1.179 created 2011, modified 2016]

EC 3.2.1.180

Accepted name: unsaturated chondroitin disaccharide hydrolase

**Reaction:**  $\beta$ -D-4-deoxy- $\Delta$ <sup>4</sup>-GlcAp-(1 $\rightarrow$ 3)- $\beta$ -D-GalNAc6S + H<sub>2</sub>O = 5-dehydro-4-deoxy-D-glucuronate + N-

acetyl- $\beta$ -D-galactosamine-6-O-sulfate

**Other name(s):** UGL (ambiguous); unsaturated glucuronyl hydrolase (ambiguous)

**Systematic name:**  $\beta$ -D-4-deoxy- $\Delta^4$ -GlcAp-(1 $\rightarrow$ 3)- $\beta$ -D-GalNAc6S hydrolase

**Comments:** The enzyme releases 4-deoxy-4,5-didehydro D-glucuronic acid or 4-deoxy-4,5-didehydro L-iduronic

acid from chondroitin disaccharides, hyaluronan disaccharides and heparin disaccharides and cleaves both glycosidic  $(1\rightarrow 3)$  and  $(1\rightarrow 4)$  bonds. It prefers the sulfated disaccharides to the unsulfated disaccharides

charides.

**References:** [1828, 2044]

[EC 3.2.1.180 created 2011]

EC 3.2.1.181

**Accepted name:** galactan endo-β-1,3-galactanase

**Reaction:** The enzyme specifically hydrolyses  $\beta$ -1,3-galactan and  $\beta$ -1,3-galactooligosaccharides

**Other name**(s): endo- $\beta$ -1,3-galactanase

**Systematic name:** arabinogalactan 3-β-D-galactanohydrolase

**Comments:** The enzyme from the fungus *Flammulina velutipes* (winter mushroom) hydrolyses the  $\beta(1\rightarrow 3)$  bonds

found in type II plant arabinogalactans, which occur in cell walls of dicots and cereals. The enzyme is an endohydrolase, and requires at least 3 contiguous  $\beta$ -1,3-residues. *cf.* EC 3.2.1.89, arabinogalactan

endo- $\beta$ -1,4-galactanase and EC 3.2.1.145, galactan 1,3- $\beta$ -galactosidase.

**References:** [1518]

[EC 3.2.1.181 created 2012]

EC 3.2.1.182

**Accepted name:** 4-hydroxy-7-methoxy-3-oxo-3,4-dihydro-2*H*-1,4-benzoxazin-2-yl glucoside β-D-glucosidase

**Reaction:** (1) (2*R*)-4-hydroxy-7-methoxy-3-oxo-3,4-dihydro-2*H*-1,4-benzoxazin-2-yl  $\beta$ -D-glucopyranoside +

 $H_2O = 2,4$ -dihydroxy-7-methoxy-2*H*-1,4-benzoxazin-3(4*H*)-one + D-glucose

(2) (2R)-4-hydroxy-3-oxo-3,4-dihydro-2H-1,4-benzoxazin-2-yl  $\beta$ -D-glucopyranoside +  $H_2O$  = 2,4-

dihydroxy-2*H*-1,4-benzoxazin-3(4*H*)-one + D-glucose

**Other name(s):** DIMBOAGlc hydrolase; DIMBOA glucosidase

**Systematic name:** (2R)-4-hydroxy-7-methoxy-3-oxo-3,4-dihydro-2H-1,4-benzoxazin-2-yl  $\beta$ -D-glucopyranoside  $\beta$ -D-

glucosidase

Comments: The enzyme from Triticum aestivum (wheat) has a higher affinity for DIMBOA glucoside than DI-

BOA glucoside. With Secale cereale (rye) the preference is reversed.

**References:** [2788, 2787, 525, 2087, 2790, 2789]

[EC 3.2.1.182 created 2012]

EC 3.2.1.183

**Accepted name:** UDP-*N*-acetylglucosamine 2-epimerase (hydrolysing)

**Reaction:** UDP-N-acetyl- $\alpha$ -D-glucosamine + H<sub>2</sub>O = N-acetyl-D-mannosamine + UDP

Other name(s): UDP-N-acetylglucosamine 2-epimerase (ambiguous); GNE (gene name); siaA (gene name); neuC

(gene name)

**Systematic name:** UDP-*N*-acetyl- $\alpha$ -D-glucosamine hydrolase (2-epimerising)

**Comments:** The enzyme is found in mammalian liver, as well as in some pathogenic bacteria including *Neisseria* 

meningitidis and Staphylococcus aureus. It catalyses the first step of sialic acid (N-acetylneuraminic acid) biosynthesis. The initial product formed is the  $\alpha$  anomer, which rapidly mutarotates to a mixture of anomers [440]. The mammalian enzyme is bifunctional and also catalyses EC 2.7.1.60, N-acetylmannosamine kinase. cf. EC 5.1.3.14, UDP-N-acetylglucosamine 2-epimerase (non-

hydrolysing).

**References:** [2756, 440, 247, 2007]

[EC 3.2.1.183 created 2012]

EC 3.2.1.184

**Accepted name:** UDP-*N*,*N'*-diacetylbacillosamine 2-epimerase (hydrolysing)

**Reaction:** UDP-N,N'-diacetylbacillosamine + H<sub>2</sub>O = UDP + 2,4-diacetamido-2,4,6-trideoxy-D-mannopyranose

**Other name(s):** UDP-Bac2Ac<sub>4</sub>Ac 2-epimerase; NeuC

**Systematic name:** UDP-*N*,*N*′-diacetylbacillosamine hydrolase (2-epimerising)

Comments: Requires  $Mg^{2+}$ . Involved in biosynthesis of legionaminic acid, a nonulosonate derivative that is in-

corporated by some bacteria into assorted virulence-associated cell surface glycoconjugates. The initial product formed by the enzyme from  $Legionella\ pneumophila$ , which incorporates legionaminic acid into the O-antigen moiety of its lipopolysaccharide, is 2,4-diacetamido-2,4,6-trideoxy- $\alpha$ -D-mannopyranose, which rapidly mutarotates to a mixture of anomers [928]. The enzyme from  $Campylobacter\ jejuni$ , which incorporates legionaminic acid into flagellin, prefers GDP-N,N'-

diacetylbacillosamine [2582].

**References:** [928, 2582]

[EC 3.2.1.184 created 2012]

EC 3.2.1.185

**Accepted name:** non-reducing end β-L-arabinofuranosidase

**Reaction:**  $\beta$ -L-arabinofuranosyl- $(1\rightarrow 2)$ - $\beta$ -L-arabinofuranose +  $H_2O = 2$   $\beta$ -L-arabinofuranose

**Other name(s):** HypBA1

Systematic name:  $\beta$ -L-arabinofuranoside non-reducing end  $\beta$ -L-arabinofuranosidase

**Comments:** The enzyme, which was identified in the bacterium *Bifidobacterium longum* JCM1217, removes

the  $\beta$ -L-arabinofuranose residue from the non-reducing end of multiple substrates, including  $\beta$ -L-arabinofuranosyl-hydroxyproline (Ara-Hyp), Ara<sub>2</sub>-Hyp, Ara<sub>3</sub>-Hyp, and  $\beta$ -L-arabinofuranosyl-(1 $\rightarrow$ 2)-1-O-methyl- $\beta$ -L-arabinofuranose. In the presence of 1-alkanols, the enzyme demonstrates transglycosylation activity, retaining the anomeric configuration of the arabinofuranose residue. *cf.* EC 3.2.1.55,

non-reducing end α-L-arabinofuranosidase

**References:** [852]

[EC 3.2.1.185 created 2013]

EC 3.2.1.186

**Accepted name:** protodioscin 26-*O*-β-D-glucosidase

**Reaction:** protodioscin +  $H_2O = 26$ -deglucoprotodioscin + D-glucose

Other name(s): F26G; torvosidase; CSF26G1; furostanol glycoside 26-O-β-D-glucosidase; furostanol 26-O-β-D-

glucoside glucohydrolase

**Systematic name:** protodioscin glucohydrolase

**Comments:** The enzyme has been characterized from the plants *Cheilocostus speciosus* and *Solanum torvum*. It

also hydrolyses the 26- $\beta$ -D-glucose group from related steroid glucosides such as protogracillin, tor-

voside A and torvoside H.

**References:** [1254, 77]

[EC 3.2.1.186 created 2013]

EC 3.2.1.187

**Accepted name:** (Ara-f)<sub>3</sub>-Hyp  $\beta$ -L-arabinobiosidase

**Reaction:** 4-O-( $\beta$ -L-arabinofuranosyl-( $1\rightarrow 2$ )- $\beta$ -L-arabinofuranosyl-( $1\rightarrow 2$ )- $\beta$ -L-arabinofuranosyl)-(2S,4S)-

4-hydroxyproline +  $H_2O = 4$ -O-( $\beta$ -L-arabinofuranosyl)-(2S,4S)-4-hydroxyproline +  $\beta$ -L-

arabinofuranosyl- $(1\rightarrow 2)$ - $\beta$ -L-arabinofuranose

Other name(s): hypBA2 (gene name);  $\beta$ -L-arabinobiosidase

**Systematic name:** 4-O-( $\beta$ -L-arabinofuranosyl-( $1 \rightarrow 2$ )- $\beta$ -L-arabinofuranosyl-( $1 \rightarrow 2$ )- $\beta$ -L-arabinofuranosyl)-(2S,4S)-4-

hydroxyproline  $\beta$ -L-arabinofuranosyl- $(1\rightarrow 2)$ - $\beta$ -L-arabinofuranose hydrolase

Comments: The enzyme, which was identified in the bacterium Bifidobacterium longum JCM1217, is specific for

(Ara-f)<sub>3</sub>-Hyp, a sugar chain found in hydroxyproline-rich glyoproteins such as extensin and lectin. The enzyme was not able to accept (Ara-f)<sub>2</sub>-Hyp or (Ara-f)<sub>4</sub>-Hyp as substrates. In the presence of 1-alkanols, the enzyme demonstrates transglycosylation activity, retaining the anomeric configuration of

the arabinofuranose residue.

**References:** [851]

[EC 3.2.1.187 created 2013]

EC 3.2.1.188

Accepted name: avenacosidase

**Reaction:** avenacoside B +  $H_2O$  = 26-desgluco-avenacoside B + D-glucose

Other name(s): As-P60

**Systematic name:** avenacoside B 26-β-D-glucohydrolase

Comments: Isolated from oat (Avena sativa) seedlings. The product acts as a defense system against fungal infec-

tion. Also acts on avenacoside A.

**References:** [1006, 1005]

[EC 3.2.1.188 created 2013]

EC 3.2.1.189

**Accepted name:** dioscin glycosidase (diosgenin-forming)

**Reaction:**  $3-O-[\alpha-L-Rha-(1\rightarrow 4)-[\alpha-L-Rha-(1\rightarrow 2)]-\beta-D-Glc]$ diosgenin + 3 H<sub>2</sub>O = D-glucose + 2 L-rhamnose +

diosgenin

Other name(s): dioscin glycosidase (aglycone-forming)

Systematic name:  $3-O-[\alpha-L-Rha-(1\rightarrow 4)-[\alpha-L-Rha-(1\rightarrow 2)]-\beta-D-Glc]$ diosgenin hydrolase (diosgenin-forming)

**Comments:** The enzyme is involved in degradation of the steroid saponin dioscin by some fungi of the *Absidia* 

genus. The enzyme can also hydrolyse 3-O-[ $\alpha$ -L-Ara-( $1 \rightarrow 4$ )-[ $\alpha$ -L-Rha-( $1 \rightarrow 2$ )]- $\beta$ -D-Glc]diosgenin into diosgenin and free sugars as the final products. cf. EC 3.2.1.190, dioscin glycosidase (3-O- $\beta$ -D-

Glc-diosgenin-forming).

**References:** [829]

[EC 3.2.1.189 created 2013]

EC 3.2.1.190

**Accepted name:** dioscin glycosidase (3-*O*-β-D-Glc-diosgenin-forming)

**Reaction:**  $3-O-[\alpha-L-Rha-(1\rightarrow 4)-[\alpha-L-Rha-(1\rightarrow 2)]-\beta-D-Glc]$ diosgenin + **2** H<sub>2</sub>O = **2** L-rhamnopyranose + dios-

genin 3-*O*-β-D-glucopyranoside

Other name(s): dioscin-α-L-rhamnosidase

**Systematic name:** 3-O-[ $\alpha$ -L-Rha-(1 $\rightarrow$ 4)-[ $\alpha$ -L-Rha-(1 $\rightarrow$ 2)]- $\beta$ -D-Glc]diosgenin (3-O- $\beta$ -D-Glc-diosgenin-forming)

**Comments:** The enzyme is involved in the hydrolysis of the steroid saponin dioscin by the digestive system of *Sus* 

scrofa (pig). cf. EC 3.2.1.189, dioscin glycosidase (diosgenin-forming).

**References:** [2336]

[EC 3.2.1.190 created 2013]

EC 3.2.1.191

Accepted name: ginsenosidase type III

**Reaction:** a protopanaxadiol-type ginsenoside with two glucosyl residues at position 3 + 2 H<sub>2</sub>O = a

protopanaxadiol-type ginsenoside with no glycosidic modification at position 3 + 2 D-glucopyranose

(overall reaction)

(1a) a protopanaxadiol-type ginsenoside with two glucosyl residues at position  $3 + H_2O$  a

protopanaxadiol-type ginsenoside with one glucosyl residue at position 3 + D-glucopyranose

(1b) a protopanaxadiol-type ginsenoside with one glucosyl residue at position  $3 + H_2O = a$  protopanaxadiol-type ginsenoside with no glycosidic modification at position 3 + D-glucopyranose

**Systematic name:** protopanaxadiol-type ginsenoside 3-β-D-hydrolase

**Comments:** Ginsenosidase type III catalyses the sequential hydrolysis of the 3-O- $\beta$ -D- $(1\rightarrow 2)$ -glucopyranosyl

bond followed by hydrolysis of the 3-O- $\beta$ -D-glucopyranosyl bond of protopanaxadiol ginsenosides. When acting for example on ginsenoside Rb1 the enzyme first generates ginsenoside XVII, and sub-

sequently ginsenoside LXXV.

**References:** [1323, 48, 1167]

[EC 3.2.1.191 created 2014]

EC 3.2.1.192

**Accepted name:** ginsenoside Rb1 β-glucosidase

**Reaction:** ginsenoside Rb1 + 2 H<sub>2</sub>O = ginsenoside Rg3 + 2 D-glucopyranose (overall reaction)

(1a) ginsenoside Rb1 +  $H_2O$  = ginsenoside Rd + D-glucopyranose (1b) ginsenoside Rd +  $H_2O$  = ginsenoside Rg3 + D-glucopyranose

**Systematic name:** ginsenoside Rb1 glucohydrolase

**Comments:** Ginsenosidases catalyse the hydrolysis of glycosyl moieties attached to the C-3, C-6 or C-20 position

of ginsenosides. They are specific with respect to the nature of the glycosidic linkage, the position and the order in which the linkages are cleaved. Ginsenoside Rb1  $\beta$ -glucosidase specifically and sequentially hydrolyses the 20-[ $\beta$ -D-glucopyranosyl-( $1\rightarrow 6$ )- $\beta$ -D glucopyranosyloxy] residues attached to position 20 by first hydrolysing the ( $1\rightarrow 6$ )-glucosidic bond to generate ginsenoside Rd as an interme-

diate, followed by hydrolysis of the remaining 20-O-β-D-glucosidic bond.

**References:** [3252]

[EC 3.2.1.192 created 2014]

EC 3.2.1.193

**Accepted name:** ginsenosidase type I

**Reaction:** (1) a protopanaxadiol-type ginsenoside with two glucosyl residues at position  $3 + H_2O = a$ 

protopanaxadiol-type ginsenoside with one glucosyl residue at position 3 + D-glucopyranose

(2) a protopanaxadiol-type ginsenoside with one glucosyl residue at position  $3 + H_2O = a$  protopanaxadiol-type ginsenoside with no glycosidic modifications at position 3 + D-glucopyranose (3) a protopanaxadiol-type ginsenoside with two glycosyl residues at position  $20 + H_2O = a$  protopanaxadiol-type ginsenoside with a single glucosyl residue at position 20 + a monosaccharide

**Systematic name:** ginsenoside glucohydrolase

**Comments:** Ginsenosidase type I is slightly activated by  $Mg^{2+}$  or  $Ca^{2+}$  [3309]. The enzyme hydrolyses the 3-O-

β-D-(1 $\rightarrow$ 2)-glucosidic bond, the 3-O-β-D-glucopyranosyl bond and the 20-O-β-D-(1 $\rightarrow$ 6)-glycosidic bond of protopanaxadiol-type ginsenosides. It usually leaves a single glucosyl residue attached at position 20 and one or no glucosyl residues at position 3. Starting with a ginsenoside that is glycosylated at both positions (e.g. ginsenoside Rb1, Rb2, Rb3, Rc or Rd), the most common products are ginseno-

side F2 and ginsenoside C-K, with low amounts of ginsenoside Rh2.

**References:** [3309]

[EC 3.2.1.193 created 2014]

EC 3.2.1.194

Accepted name: ginsenosidase type IV

**Reaction:** a protopanaxatriol-type ginsenoside with two glycosyl residues at position  $6 + 2 H_2O = a$ 

protopanaxatriol-type ginsenoside with no glycosidic modification at position 6 + D-glucopyranose

+ a monosaccharide (overall reaction)

(1a) a protopanaxatriol-type ginsenoside with two glycosyl residues at position  $6 + H_2O = a$ 

protopanaxatriol-type ginsenoside with a single glucosyl at position 6 + a monosaccharide

(1b) a protopanaxatriol-type ginsenoside with a single glucosyl at position  $6 + H_2O = a$  protopanaxatriol-type ginsenoside with no glycosidic modification at position 6 + D-glucopyranose

**Systematic name:** protopanaxatriol-type ginsenoside 6-β-D-glucohydrolase

**Comments:** Ginsenosidase type IV catalyses the sequential hydrolysis of the 6-O- $\beta$ -D- $(1\rightarrow 2)$ -glycosidic bond

or the 6-O- $\alpha$ -D-(1 $\rightarrow$ 2)-glycosidic bond in protopanaxatriol-type ginsenosides with a disacchride attached to the C<sub>6</sub> position, followed by the hydrolysis of the remaining 6-O- $\beta$ -D-glycosidic bond (e.g.

ginsenoside Re  $\rightarrow$  ginsenoside Rg1  $\rightarrow$  ginsenoside F1).

**References:** [3107, 3106]

[EC 3.2.1.194 created 2014]

EC 3.2.1.195

**Accepted name:** 20-*O*-multi-glycoside ginsenosidase

**Reaction:** a protopanaxadiol-type ginsenoside with two glycosyl residues at position  $20 + H_2O = a$ 

protopanaxadiol-type ginsenoside with a single glucosyl residue at position 20 + a monosaccharide

**Other name(s):** ginsenosidase type II (erroneous)

**Systematic name:** protopanaxadiol-type ginsenoside 20-β-D-glucohydrolase

Comments: The 20-O-multi-glycoside ginsenosidase catalyses the hydrolysis of the 20-O- $\alpha$ -(1 $\rightarrow$ 6)-glycosidic

bond and the 20-O- $\beta$ - $(1\rightarrow 6)$ -glycosidic bond of protopanaxadiol-type ginsenosides. The enzyme usually leaves a single glucosyl residue attached at position 20, although it can cleave the remaining glucosyl residue with a lower efficiency. Starting with a ginsenoside that is glycosylated at positions 3 and 20, such as ginsenosides Rb1, Rb2, Rb3 and Rc, the most common product is ginsenoside Rd,

with a low amount of ginsenoside Rg3 also formed.

**References:** [3308]

[EC 3.2.1.195 created 2014]

EC 3.2.1.196

**Accepted name:** limit dextrin  $\alpha$ -1,6-maltotetraose-hydrolase

**Reaction:** Hydrolysis of  $(1\rightarrow 6)$ - $\alpha$ -D-glucosidic linkages to branches with degrees of polymerization of three or

four glucose residues in limit dextrin.

Other name(s): glgX (gene name); glycogen debranching enzyme (ambiguous) Systematic name: glycogen phosphorylase-limit dextrin maltotetraose-hydrolase Comments: This bacterial enzyme catalyses a reaction similar to EC 3.2.1.33, amylo- $\alpha$ -1,6-glucosidase (one

of the activities of the eukaryotic glycogen debranching enzyme). However, while EC 3.2.1.33 removes single glucose residues linked by 1,6- $\alpha$ -linkage, and thus requires the additional activity of 4- $\alpha$ -glucanotransferase (EC 2.4.1.25) to act on limit dextrins formed by glycogen phosphorylase (EC 2.4.1.1), this enzyme removes maltotriose and maltotetraose chains that are attached by 1,6- $\alpha$ -linkage to the limit dextrin main chain, generating a debranched limit dextrin without a need for another en-

zyme.

**References:** [1313, 541, 2716]

[EC 3.2.1.196 created 2016]

EC 3.2.1.197

Accepted name:  $\beta$ -1,2-mannosidase

**Reaction:**  $\beta$ -D-mannopyranosyl- $(1\rightarrow 2)$ - $\beta$ -D-mannopyranosyl- $(1\rightarrow 2)$ -D-mannopyranose +  $H_2O = \beta$ -D-

mannopyranosyl- $(1\rightarrow 2)$ -D-mannopyranose +  $\alpha$ -D-mannopyranose

**Systematic name:**  $\beta$ -1,2-D-mannoside mannohydrolase

**Comments:** The enzyme, characterized from multiple bacterial species, catalyses the hydrolysis of terminal,

non-reducing D-mannose residues from  $\beta$ -1,2-mannotriose and  $\beta$ -1,2-mannobiose. The mechanism involves anomeric inversion, resulting in the release of  $\alpha$ -D-mannopyranose. Activity with  $\beta$ -1,2-

mannotriose or higher oligosaccharides is higher than that with  $\beta$ -1,2-mannobiose.

**References:** [523, 2083]

[EC 3.2.1.197 created 2016]

EC 3.2.1.198

**Accepted name:**  $\alpha$ -mannan endo-1,2- $\alpha$ -mannanase

**Reaction:** Hydrolysis of the terminal  $\alpha$ -D-mannosyl- $(1\rightarrow 3)$ - $\alpha$ -D-mannose disaccharide from  $\alpha$ -D-mannosyl-

 $(1\rightarrow 3)-\alpha$ -D-mannosyl- $(1\rightarrow 2)-\alpha$ -D-mannosyl- $(1\rightarrow 2)-\alpha$ -D-mannosyl side chains in fungal cell wall

α-mannans.

**Systematic name:**  $\alpha$ -mannan 1,2-[ $\alpha$ -D-mannosyl-( $1\rightarrow 3$ )- $\alpha$ -D-mannose] hydrolase

**Comments:** The enzyme, characterized from the gut bacteria *Bacteroides thetaiotaomicron* and *Bacteroides xy*-

lanisolvens, can also catalyse the reaction of EC 3.2.1.130, glycoprotein endo- $\alpha$ -1,2-mannosidase.

**References:** [1023, 524]

[EC 3.2.1.198 created 2016]

EC 3.2.1.199

Accepted name: sulfoquinovosidase

**Reaction:** an 6-sulfo- $\alpha$ -D-quinovosyl diacylglycerol + H<sub>2</sub>O = 6-sulfo-D-quinovose + a 1,2-diacylglycerol

Other name(s): yihQ (gene name)

**Systematic name:** 6-sulfo-α-D-quinovosyl diacylglycerol 6-sulfo-D-quinovohydrolase

**Comments:** The enzyme, characterized from the bacteria *Escherichia coli* and *Pseudomonas putida*, hydrolyses

terminal non-reducing  $\alpha$ -sulfoquinovoside residues in  $\alpha$ -sulfoquinovosyl diacylglycerides and  $\alpha$ -

sulfoquinovosyl glycerol.

**References:** [2629, 2738]

[EC 3.2.1.199 created 2016]

EC 3.2.1.200

**Accepted name:** exo-chitinase (non-reducing end)

**Reaction:** Hydrolysis of N,N'-diacetylchitobiose from the non-reducing end of chitin and chitodextrins.

Other name(s): chiB (gene name)

Systematic name:  $(1\rightarrow 4)$ -2-acetamido-2-deoxy- $\beta$ -D-glucan diacetylchitobiohydrolase (non-reducing end)

**Comments:** The enzyme hydrolyses the second glycosidic  $(1\rightarrow 4)$  linkage from non-reducing ends of chitin and

chitodextrin molecules, liberating N,N'-diacetylchitobiose disaccharides. cf. EC 3.2.1.201, exo-

chitinase (reducing end).

**References:** [2873, 1207, 2166, 1007]

[EC 3.2.1.200 created 2017]

EC 3.2.1.201

**Accepted name:** exo-chitinase (reducing end)

**Reaction:** Hydrolysis of N,N'-diacetylchitobiose from the reducing end of chitin and chitodextrins.

**Other name(s):** *chiA* (gene name)

Systematic name:  $(1\rightarrow 4)$ -2-acetamido-2-deoxy- $\beta$ -D-glucan diacetylchitobiohydrolase (reducing end)

**Comments:** The enzyme hydrolyses the second glycosidic  $(1\rightarrow 4)$  linkage from reducing ends of chitin and chi-

todextrin molecules, liberating N,N'-diacetylchitobiose disaccharides. cf. EC 3.2.1.200, exo-chitinase

(non-reducing end).

**References:** [1207, 2036, 1007, 321]

[EC 3.2.1.201 created 2017]

EC 3.2.1.202

**Accepted name:** endo-chitodextinase

**Reaction:** Hydrolysis of chitodextrins, releasing N,N'-diacetylchitobiose and small amounts of N,N',N''-

triacetylchitotriose.

Other name(s): endo I (gene name); chitodextrinase (ambiguous); endolytic chitodextrinase; periplasmic chitodextri

nase

Systematic name:  $(1\rightarrow 4)$ -2-acetamido-2-deoxy- $\beta$ -D-glucan diacetylchitobiohydrolase (endo-cleaving)

**Comments:** The enzyme, characterized from the bacterium *Vibrio furnissii*, is an endo-cleaving chitodextrinase

that participates in the the chitin catabolic pathway found in members of the *Vibrionaceae*. Unlike EC 3.2.1.14, chitinase, it has no activity on chitin. The smallest substrate is a tetrasaccharide, and the final products are N,N'-diacetylchitobiose and small amounts of N,N',N''-triacetylchitotriose. cf. EC

3.2.1.200, exo-chitinase (non-reducing end), and EC 3.2.1.201, exo-chitinase (reducing end).

**References:** [170, 1432]

[EC 3.2.1.202 created 2017]

EC 3.2.1.203

Accepted name: carboxymethylcellulase

**Reaction:** Endohydrolysis of  $(1\rightarrow 4)$ - $\beta$ -D-glucosidic linkages in (carboxymethyl)cellulose.

Other name(s): CMCase

Systematic name: 4-β-D-(carboxymethyl)glucan 4-(carboxymethyl)glucanohydrolase

**Comments:** The enzyme from the acidophilic bacterium *Alicyclobacillus acidocaldarius* is an endo-cleaving hy-

drolase that cleaves  $\beta(1\rightarrow 4)$ -linked residues. However, it is specific for (carboxymethyl)cellulose and

does not act on cellulosic substrates such as avicel.

References: [1962]

[EC 3.2.1.203 created 2017]

EC 3.2.1.204

**Accepted name:** 1,3-α-isomaltosidase

**Reaction:** cyclobis- $(1\rightarrow 6)$ - $\alpha$ -nigerosyl + 2 H<sub>2</sub>O = 2 isomaltose (overall reaction)

(1a) cyclobis- $(1\rightarrow 6)$ - $\alpha$ -nigerosyl +  $H_2O = \alpha$ -isomaltosyl- $(1\rightarrow 3)$ -isomaltose

(1b)  $\alpha$ -isomaltosyl-(1 $\rightarrow$ 3)-isomaltose + H<sub>2</sub>O = **2** isomaltose

**Systematic name:** 1,3- $\alpha$ -isomaltohydrolase (configuration-retaining)

**Comments:** The enzyme, characterized from the bacteria Bacillus sp. NRRL B-21195 and Kribbella flavida,

participates in the degradation of starch. The cyclic tetrasaccharide cyclobis- $(1\rightarrow 6)$ - $\alpha$ -nigerosyl is

formed from starch extracellularly and imported into the cell, where it is degraded to glucose.

**References:** [1457, 2828]

[EC 3.2.1.204 created 2017]

EC 3.2.1.205

Accepted name: isomaltose glucohydrolase

> **Reaction:** isomaltose +  $H_2O = \beta$ -D-glucose + D-glucose

**Systematic name:** isomaltose 6-α-glucohydrolase (configuration-inverting)

**Comments:** The enzyme catalyses the hydrolysis of  $\alpha$ -1,6-glucosidic linkages from the non-reducing end of its

substrate. Unlike EC 3.2.1.10, oligo-1,6-glucosidase, the enzyme inverts the anomeric configuration

of the released residue. The enzyme can also act on panose and maltotriose at a lower rate.

**References:** [2828]

[EC 3.2.1.205 created 2017]

EC 3.2.1.206

oleuropein β-glucosidase Accepted name:

> **Reaction:** oleuropein + H<sub>2</sub>O = oleuropein aglycone + D-glucopyranose

OeGLU (gene name) Other name(s):

**Systematic name:** oleuropein 2-β-D-glucohydrolase

> **Comments:** Oleuropein is a glycosylated secoiridoid exclusively biosynthesized by members of the Oleaceae plant

> > family where it is part of a defence system againt herbivores. The enzyme also hydrolyses ligstroside

and demethyloleuropein.

[451, 2455, 1008, 1520, 1521] References:

[EC 3.2.1.206 created 2017]

EC 3.2.1.207

Accepted name: mannosyl-oligosaccharide α-1,3-glucosidase

> (1)  $Glc_2Man_9GlcNAc_2$ -[protein] +  $H_2O$  =  $GlcMan_9GlcNAc_2$ -[protein] +  $\beta$ -D-glucopyranose **Reaction:**

> > (2) GlcMan<sub>9</sub>GlcNAc<sub>2</sub>-[protein] + H<sub>2</sub>O = Man<sub>9</sub>GlcNAc<sub>2</sub>-[protein] + β-D-glucopyranose

Other name(s): ER glucosidase II; α-glucosidase II; trimming glucosidase II; ROT2 (gene name); GTB1 (gene name);

GANAB (gene name); PRKCSH (gene name)

Glc<sub>2</sub>Man<sub>9</sub>GlcNAc<sub>2</sub>-[protein] 3-α-glucohydrolase (configuration-inverting) **Systematic name:** 

**Comments:** This eukaryotic enzyme cleaves off sequentially the two  $\alpha$ -1,3-linked glucose residues from the

Glc<sub>2</sub>Man<sub>9</sub>GlcNAc<sub>2</sub> oligosaccharide precursor of immature *N*-glycosylated proteins.

**References:** [2965, 3341, 3190, 1960]

[EC 3.2.1.207 created 2018]

EC 3.2.1.208

Accepted name: glucosylglycerate hydrolase

> $2-O-(\alpha-D-glucopyranosyl)-D-glycerate + H<sub>2</sub>O = D-glucopyranose + D-glycerate$ **Reaction:**

Other name(s): GG hydrolase; GgH

**Systematic name:** 2-O-(α-D-glucopyranosyl)-D-glycerate D-glucohydrolase

**Comments:** The enzyme has been isolated from nontuberculous mycobacteria (e.g. Mycobacterium hassiacum),

which accumulate 2-*O*-(α-D-glucopyranosyl)-D-glycerate during growth under nitrogen deprivation.

References: [27, 386]

[EC 3.2.1.208 created 2018]

# EC 3.2.2 Hydrolysing N-glycosyl compounds

EC 3.2.2.1

Accepted name: purine nucleosidase

**Reaction:** a purine nucleoside  $+ H_2O = D$ -ribose + a purine base

Other name(s): nucleosidase (misleading); purine  $\beta$ -ribosidase; purine nucleoside hydrolase; purine ribonucleosidase;

ribonucleoside hydrolase (misleading); nucleoside hydrolase (misleading); *N*-ribosyl purine ribohydrolase; nucleosidase g; *N*-D-ribosylpurine ribohydrolase; inosine-adenosine-guanosine preferring nucleoside hydrolase; purine-specific nucleoside *N*-ribohydrolase; IAG-nucleoside hydrolase; IAG-

NH

**Systematic name:** purine-nucleoside ribohydrolase

**Comments:** The enzyme from the bacterium *Ochrobactrum anthropi* specifically catalyses the irreversible *N*-

riboside hydrolysis of purine nucleosides. Pyrimidine nucleosides, purine and pyrimidine nucleotides,

NAD<sup>+</sup>, NADP<sup>+</sup> and nicotinaminde mononucleotide are not substrates [2148].

**References:** [1122, 1362, 2834, 2890, 2237, 2148, 3063, 1858]

[EC 3.2.2.1 created 1961, modified 2006, modified 2011]

EC 3.2.2.2

Accepted name: inosine nucleosidase

**Reaction:** inosine +  $H_2O$  = D-ribose + hypoxanthine **Other name(s):** inosinase; inosine-guanosine nucleosidase

**Systematic name:** inosine ribohydrolase

**References:** [1492, 2890]

[EC 3.2.2.2 created 1961]

EC 3.2.2.3

Accepted name: uridine nucleosidase

**Reaction:** uridine +  $H_2O$  = D-ribose + uracil

Other name(s): uridine hydrolase
Systematic name: uridine ribohydrolase

**References:** [377]

[EC 3.2.2.3 created 1961]

EC 3.2.2.4

Accepted name: AMP nucleosidase

**Reaction:** AMP +  $H_2O$  = D-ribose 5-phosphate + adenine

**Other name(s):** adenylate nucleosidase; adenosine monophosphate nucleosidase

**Systematic name:** AMP phosphoribohydrolase

**References:** [1211]

[EC 3.2.2.4 created 1961]

EC 3.2.2.5

**Accepted name:** NAD<sup>+</sup> glycohydrolase

**Reaction:**  $NAD^+ + H_2O = ADP$ -D-ribose + nicotinamide

**Other name(s):** NAD glycohydrolase; nicotinamide adenine dinucleotide glycohydrolase; β-NAD<sup>+</sup> glycohydrolase;

DPNase (ambiguous); NAD hydrolase (ambiguous); diphosphopyridine nucleosidase (ambiguous); nicotinamide adenine dinucleotide nucleosidase (ambiguous); NAD nucleosidase (ambiguous); DPN

hydrolase (ambiguous); NADase (ambiguous); nga (gene name); NAD<sup>+</sup> nucleosidase

**Systematic name:** NAD<sup>+</sup> glycohydrolase

**Comments:** This enzyme catalyses the hydrolysis of NAD<sup>+</sup>, without associated ADP-ribosyl cyclase activity (un-

like the metazoan enzyme EC 3.2.2.6, bifunctional ADP-ribosyl cyclase/cyclic ADP-ribose hydrolase). The enzyme from Group A streptococci has been implicated in the pathogenesis of diseases such as streptococcal toxic shock-like syndrome (STSS) and necrotizing fasciitis. The enzyme from

the venom of the snake Agkistrodon acutus also catalyses EC 3.6.1.5, apyrase [3329].

**References:** [750, 995, 3329, 905, 2696]

[EC 3.2.2.5 created 1961, modified 2013]

EC 3.2.2.6

**Accepted name:** ADP-ribosyl cyclase/cyclic ADP-ribose hydrolase

**Reaction:**  $NAD^+ + H_2O = ADP$ -D-ribose + nicotinamide (overall reaction)

(1a)  $NAD^+$  = cyclic ADP-ribose + nicotinamide (1b) cyclic ADP-ribose +  $H_2O$  = ADP-D-ribose

Other name(s): NAD<sup>+</sup> nucleosidase; NADase (ambiguous); DPNase (ambiguous); DPN hydrolase (ambiguous);

NAD hydrolase (ambiguous); nicotinamide adenine dinucleotide nucleosidase (ambiguous); NAD glycohydrolase (misleading); NAD nucleosidase (ambiguous); nicotinamide adenine dinucleotide gly-

cohydrolase (misleading); CD38 (gene name); BST1 (gene name)

**Systematic name:** NAD<sup>+</sup> glycohydrolase (cyclic ADP-ribose-forming)

**Comments:** This multiunctional enzyme acts on NAD<sup>+</sup>, catalysing both the synthesis and hydrolysis of cyclic

ADP-ribose, a calcium messenger that can mobilize intracellular Ca<sup>2+</sup> stores and activate Ca<sup>2+</sup> influx to regulate a wide range of physiological processes. In addition, the enzyme also catalyses EC 2.4.99.20, 2'-phospho-ADP-ribosyl cyclase/2'-phospho-cyclic-ADP-ribose transferase. It is also able

to act on β-nicotinamide D-ribonucleotide. *cf.* EC 3.2.2.5, NAD<sup>+</sup> glycohydrolase.

**References:** [1238, 1193, 2853, 2928, 827, 3250, 1724]

[EC 3.2.2.6 created 1961, modified 2004, modified 2014, modified 2018]

EC 3.2.2.7

Accepted name: adenosine nucleosidase

**Reaction:** adenosine +  $H_2O$  = D-ribose + adenine

**Other name(s):** adenosinase; *N*-ribosyladenine ribohydrolase; adenosine hydrolase; ANase

Systematic name: adenosine ribohydrolase

**Comments:** Also acts on adenosine *N*-oxide.

**References:** [1856]

[EC 3.2.2.7 created 1972]

EC 3.2.2.8

**Accepted name:** ribosylpyrimidine nucleosidase

**Reaction:** a pyrimidine nucleoside +  $H_2O$  = D-ribose + a pyrimidine base

**Other name(s):** *N*-ribosylpyrimidine nucleosidase; pyrimidine nucleosidase; *N*-ribosylpyrimidine ribohydrolase;

pyrimidine nucleoside hydrolase; RihB; YeiK; nucleoside ribohydrolase

**Systematic name:** pyrimidine-nucleoside ribohydrolase

**Comments:** Also hydrolyses purine D-ribonucleosides, but much more slowly. 2'-, 3'- and 5'-deoxynucleosides are

not substrates [910].

**References:** [2900, 2260, 910, 911]

[EC 3.2.2.8 created 1972]

EC 3.2.2.9

**Accepted name:** adenosylhomocysteine nucleosidase

**Reaction:** S-adenosyl-L-homocysteine +  $H_2O = S$ -(5-deoxy-D-ribos-5-yl)-L-homocysteine + adenine **Other name(s):** S-adenosylhomocysteine hydrolase (ambiguous); S-adenosylhomocysteine nucleosidase; 5'-

methyladenosine nucleosidase; S-adenosylhomocysteine/5'-methylthioadenosine nucleosidase; Ado-

Hcy/MTA nucleosidase

**Systematic name:** S-adenosyl-L-homocysteine homocysteinylribohydrolase

**Comments:** Also acts on S-methyl-5'-thioadenosine to give adenine and S-methyl-5-thioribose (cf. EC 3.2.2.16,

methylthioadenosine nucleosidase).

**References:** [662, 761]

[EC 3.2.2.9 created 1972, modified 2004]

EC 3.2.2.10

**Accepted name:** pyrimidine-5'-nucleotide nucleosidase

**Reaction:** a pyrimidine 5'-nucleotide +  $H_2O$  = D-ribose 5-phosphate + a pyrimidine base

**Other name(s):** pyrimidine nucleotide *N*-ribosidase; Pyr5N

**Systematic name:** pyrimidine-5'-nucleotide phosphoribo(deoxyribo)hydrolase

Comments: Also acts on dUMP, dTMP and dCMP.

**References:** [1233, 1234]

[EC 3.2.2.10 created 1972]

EC 3.2.2.11

**Accepted name:** β-aspartyl-*N*-acetylglucosaminidase

**Reaction:**  $1-\beta$ -aspartyl-*N*-acetyl-D-glucosaminylamine + H<sub>2</sub>O = L-asparagine + *N*-acetyl-D-glucosamine

**Other name(s):** β-aspartylacetylglucosaminidase

**Systematic name:** 1-β-aspartyl-*N*-acetyl-D-glucosaminylamine L-asparaginohydrolase

**References:** [728]

[EC 3.2.2.11 created 1972]

EC 3.2.2.12

Accepted name: inosinate nucleosidase

**Reaction:** IMP +  $H_2O$  = D-ribose 5-phosphate + hypoxanthine

Other name(s): 5'-inosinate phosphoribohydrolase

Systematic name: IMP phosphoribohydrolase

stematic name: Invir phosphorioonytholas

**References:** [1554]

[EC 3.2.2.12 created 1972]

EC 3.2.2.13

Accepted name: 1-methyladenosine nucleosidase

**Reaction:** 1-methyladenosine +  $H_2O = 1$ -methyladenine + D-ribose

Other name(s): 1-methyladenosine hydrolase
Systematic name: 1-methyladenosine ribohydrolase

**References:** [2891]

[EC 3.2.2.13 created 1976]

EC 3.2.2.14

**Accepted name:** NMN nucleosidase

**Reaction:**  $\beta$ -nicotinamide D-ribonucleotide +  $H_2O$  = D-ribose 5-phosphate + nicotinamide

Other name(s): NMNase; nicotinamide mononucleotide nucleosidase; nicotinamide mononucleotidase; NMN glyco-

hydrolase; NMNGhase

Systematic name: nicotinamide-nucleotide phosphoribohydrolase

**Comments:** The enzyme is thought to participate in an NAD<sup>+</sup>-salvage pathway. In eukaryotic organisms this ac-

tivity has been attributed to EC 3.2.2.6, ADP-ribosyl cyclase/cyclic ADP-ribose hydrolase.

**References:** [56, 1236, 1237]

[EC 3.2.2.14 created 1976, modified 2018]

EC 3.2.2.15

Accepted name: DNA-deoxyinosine glycosylase

**Reaction:** Hydrolyses DNA and polynucleotides, releasing free hypoxanthine

Other name(s): DNA(hypoxanthine) glycohydrolase; deoxyribonucleic acid glycosylase; hypoxanthine-DNA glyco-

sylase

Systematic name: DNA-deoxyinosine deoxyribohydrolase

**References:** [1391]

[EC 3.2.2.15 created 1980, modified 1982, modified 2000]

EC 3.2.2.16

Accepted name: methylthioadenosine nucleosidase

**Reaction:** S-methyl-5'-thioadenosine +  $H_2O = S$ -methyl-5-thio-D-ribose + adenine

Other name(s): 5'-methylthioadenosine nucleosidase; MTA nucleosidase; MeSAdo nucleosidase; methylthioadeno-

sine methylthioribohydrolase

**Systematic name:** S-methyl-5'-thioadenosine adeninehyrolase

**Comments:** Does not act on S-adenosylhomocysteine. cf. EC 3.2.2.9 adenosylhomocysteine nucleosidase.

References: [1003]

[EC 3.2.2.16 created 1983, modified 2004]

EC 3.2.2.17

Accepted name: deoxyribodipyrimidine endonucleosidase

**Reaction:** Cleaves the N-glycosidic bond between the 5'-pyrimidine residue in cyclobutadipyrimidine (in DNA)

and the corresponding deoxy-D-ribose residue

Other name(s): pyrimidine dimer DNA-glycosylase; endonuclease V; deoxyribonucleate pyrimidine dimer glycosi-

dase; pyrimidine dimer DNA glycosylase; T4-induced UV endonuclease; PD-DNA glycosylase

Systematic name: deoxy-D-ribocyclobutadipyrimidine polynucleotidodeoxyribohydrolase

**References:** [1069]

[EC 3.2.2.17 created 1983]

[3.2.2.18 Deleted entry. glycopeptide N-glycosidase. Now included with EC 3.5.1.52, peptide- $N^4$ -(N-acetyl- $\beta$ -glucosaminyl)asparagine amidase]

[EC 3.2.2.18 created 1984, deleted 1989]

EC 3.2.2.19

Accepted name: [protein ADP-ribosylarginine] hydrolase

**Reaction:** (1) protein- $N^{\omega}$ -(ADP-D-ribosyl)-L-arginine + H<sub>2</sub>O = ADP-D-ribose + protein-L-arginine

(2)  $N^{\omega}$ -(ADP-D-ribosyl)-L-arginine + H<sub>2</sub>O = ADP-D-ribose + L-arginine

Other name(s): ADP-ribose-L-arginine cleavage enzyme; ADP-ribosylarginine hydrolase;  $N^{\omega}$ -(ADP-D-ribosyl)-L-

arginine ADP-ribosylhydrolase; protein-ω-*N*-(ADP-D-ribosyl)-L-arginine ADP-ribosylhydrolase

**Systematic name:** protein- $N^{\omega}$ -(ADP-D-ribosyl)-L-arginine ADP-ribosylhydrolase

**Comments:** The enzyme will remove ADP-D-ribose from arginine residues in ADP-ribosylated proteins.

**References:** [1985, 1986, 1507, 2832, 2164]

# [EC 3.2.2.19 created 1989, modified 2004]

EC 3.2.2.20

**Accepted name:** DNA-3-methyladenine glycosylase I

**Reaction:** Hydrolysis of alkylated DNA, releasing 3-methyladenine

Other name(s): deoxyribonucleate 3-methyladenine glycosidase I; 3-methyladenine DNA glycosylase I; DNA-3-

methyladenine glycosidase I

Systematic name: alkylated-DNA glycohydrolase (releasing methyladenine and methylguanine)

Comments: Involved in the removal of alkylated bases from DNA in Escherichia coli (cf. EC 2.1.1.63 methylated-

DNA—[protein]-cysteine S-methyltransferase).

**References:** [725, 1390, 2914]

[EC 3.2.2.20 created 1990, modified 2000]

EC 3.2.2.21

Accepted name: DNA-3-methyladenine glycosylase II

Reaction: Hydrolysis of alkylated DNA, releasing 3-methyladenine, 3-methylguanine, 7-methylguanine and 7-

methyladenine

Other name(s): deoxyribonucleate 3-methyladenine glycosidase II; 3-methyladenine DNA glycosylase II; DNA-3-

methyladenine glycosidase II; AlkA

**Systematic name:** alkylated-DNA glycohydrolase (releasing methyladenine and methylguanine)

Comments: Involved in the removal of alkylated bases from DNA in Escherichia coli (cf. EC 2.1.1.63 methylated-

DNA—[protein]-cysteine S-methyltransferase).

**References:** [725, 1390, 2417, 2914]

[EC 3.2.2.21 created 1990, modified 2000]

EC 3.2.2.22

Accepted name: rRNA N-glycosylase

**Reaction:** Hydrolysis of the *N*-glycosylic bond at A-4324 in 28S rRNA from rat ribosomes

**Other name(s):** ribosomal ribonucleate *N*-glycosidase; nigrin b; RNA *N*-glycosidase; rRNA *N*-glycosidase; ricin;

momorcochin-S; Mirabilis antiviral protein; momorcochin-S; gelonin; saporins

**Systematic name:** rRNA *N*-glycohydrolase

Comments: Ricin A-chain and related toxins show this activity. Naked rRNA is attacked more slowly than rRNA

in intact ribosomes. Naked rRNA from  $\it Escherichia\ coli$  is cleaved at a corresponding position.

**References:** [704]

[EC 3.2.2.22 created 1990, modified 2000]

EC 3.2.2.23

**Accepted name:** DNA-formamidopyrimidine glycosylase

Reaction: Hydrolysis of DNA containing ring-opened 7-methylguanine residues, releasing 2,6-diamino-4-

hydroxy-5-(N-methyl)formamidopyrimidine

Other name(s): Fapy-DNA glycosylase; deoxyribonucleate glycosidase; 2,6-diamino-4-hydroxy-5*N*-

 $formamidopyrimidine-DNA\ glycosylase;\ 2,6-diamino-4-hydroxy-5(N-methyl) formamidopyrimidine-DNA\ glycosylase;\ DNA-formamidopyrimidine\ glycosidase;$ 

Fpg protein

**Systematic name:** DNA glycohydrolase [2,6-diamino-4-hydroxy-5-(*N*-methyl)formamidopyrimide releasing]

Comments: May play a significant role in processes leading to recovery from mutagenesis and/or cell death by

alkylating agents. Also involved in the GO system responsible for removing an oxidatively damaged

form of guanine (7,8-dihydro-8-oxoguanine) from DNA.

**References:** [255]

# [EC 3.2.2.23 created 1990, modified 2000]

EC 3.2.2.24

**Accepted name:** ADP-ribosyl-[dinitrogen reductase] hydrolase

**Reaction:** [dinitrogen reductase]- $N^{\omega}$ - $\alpha$ -(ADP-D-ribosyl)-L-arginine = ADP-D-ribose + [dinitrogen reductase]-

L-arginine

Other name(s): azoferredoxin glycosidase; azoferredoxin-activating enzymes; dinitrogenase reductase-activating gly-

cohydrolase; ADP-ribosyl glycohydrolase;  $draG\ ({\rm gene\ name})$ 

Systematic name: ADP-D-ribosyl-[dinitrogen reductase] ADP-ribosylhydrolase

**Comments:** The enzyme restores the activity of EC 1.18.6.1, nitrogenase, by catalysing the removal of ADP-

ribose from an arginine residue of the dinitrogenase reductase component of nitrogenase. This activity occurs only when the nitrogenase product, ammonium, is not available. The combined activity of this enzyme and EC 2.4.2.37, NAD<sup>+</sup>-dinitrogen-reductase ADP-D-ribosyltransferase, controls the level of

activity of nitrogenase.

**References:** [779, 1674, 206]

[EC 3.2.2.24 created 1992]

EC 3.2.2.25

**Accepted name:** *N*-methyl nucleosidase

**Reaction:** 7-methylxanthosine +  $H_2O = 7$ -methylxanthine + D-ribose

Other name(s): 7-methylxanthosine nucleosidase; N-MeNase; N-methyl nucleoside hydrolase; methylpurine nucle-

osidase

**Systematic name:** 7-methylxanthosine ribohydrolase

**Comments:** The enzyme preferentially hydrolyses 3- and 7-methylpurine nucleosides, such as 3-

methylxanthosine, 3-methyladenosine and 7-methylguanosine. Hydrolysis of 7-methylxanthosine

to form 7-methylxanthine is the second step in the caffeine-biosynthesis pathway.

**References:** [2066]

[EC 3.2.2.25 created 2007]

EC 3.2.2.26

**Accepted name:** futalosine hydrolase

**Reaction:** futalosine +  $H_2O$  = dehypoxanthine futalosine + hypoxanthine

Other name(s): futalosine nucleosidase; MqnB Systematic name: futalosine ribohydrolase

**Comments:** This enzyme, which is specific for futalosine, catalyses the second step of a novel menaquinone

biosynthetic pathway that is found in some prokaryotes.

**References:** [1149]

[EC 3.2.2.26 created 2008]

EC 3.2.2.27

Accepted name: uracil-DNA glycosylase

Reaction: Hydrolyses single-stranded DNA or mismatched double-stranded DNA and polynucleotides, releasing

free uracil

Other name(s): UdgB (ambiguous); uracil-DNA N-glycosylase; UDG (ambiguous); uracil DNA glycohydrolase

**Systematic name:** uracil-DNA deoxyribohydrolase (uracil-releasing)

**Comments:** Uracil-DNA glycosylases are widespread enzymes that are found in all living organisms. EC 3.2.2.27

and double-stranded uracil-DNA glycosylase (EC 3.2.2.28) form a central part of the DNA-repair machinery since they initiate the DNA base-excision repair pathway by hydrolysing the *N*-glycosidic bond between uracil and the deoxyribose sugar thereby catalysing the removal of mis-incorporated

uracil from DNA.

**References:** [1626, 1448, 2234, 2773]

# [EC 3.2.2.27 created 2009]

EC 3.2.2.28

Accepted name: double-stranded uracil-DNA glycosylase

**Reaction:** Specifically hydrolyses mismatched double-stranded DNA and polynucleotides, releasing free uracil **Other name(s):** Mug; double-strand uracil-DNA glycosylase; Dug; dsUDG; double-stranded DNA specific UDG;

dsDNA specific UDG; UdgB (ambiguous); G:T/U mismatch-specific DNA glycosylase; UDG (am-

biguous)

**Systematic name:** uracil-double-stranded DNA deoxyribohydrolase (uracil-releasing)

**Comments:** No activity on DNA containing a T/G mispair or single-stranded DNA containing either a site-

specific uracil or 3,*N*<sup>4</sup>-ethenocytosine residue [2804], significant role for double-stranded uracil-DNA glycosylase in mutation avoidance in non-dividing *E. coli* [1948]. Uracil-DNA glycosylases are widespread enzymes that are found in all living organisms. Uracil-DNA glycosylase (EC 3.2.2.27) and EC 3.2.2.28 form a central part of the DNA-repair machinery since they initiate the DNA base-excision repair pathway by hydrolysing the *N*-glycosidic bond between uracil and the deoxyribose

sugar thereby catalysing the removal of mis-incorporated uracil from DNA.

**References:** [161, 2804, 1948]

[EC 3.2.2.28 created 2009]

EC 3.2.2.29

**Accepted name:** thymine-DNA glycosylase

**Reaction:** Hydrolyses mismatched double-stranded DNA and polynucleotides, releasing free thymine. **Other name(s):** mismatch-specific thymine-DNA glycosylase; mismatch-specific thymine-DNA *N*-glycosylase;

hTDG; hsTDG; TDG; thymine DNA glycosylase; G/T glycosylase; uracil/thymine DNA glycosylase; T:G mismatch-specific thymidine-DNA glycosylase; G:T mismatch-specific thymine DNA-

glycosylase

**Systematic name:** thymine-DNA deoxyribohydrolase (thymine-releasing)

**Comments:** Thymine-DNA glycosylase is part of the DNA-repair machinery. Thymine removal is fastest when it

is from a G/T mismatch with a 5'-flanking C/G pair. The glycosylase removes uracil from G/U, C/U,

and T/U base pairs faster than it removes thymine from G/T [3134].

**References:** [3135, 2062, 3134]

[EC 3.2.2.29 created 2009]

EC 3.2.2.30

Accepted name: aminodeoxyfutalosine nucleosidase

**Reaction:** 6-amino-6-deoxyfutalosine +  $H_2O$  = dehypoxanthine futalosine + adenine

Other name(s): AFL nucleosidase; aminofutalosine nucleosidase; methylthioadenosine nucleosidase; MqnB

**Systematic name:** 6-amino-6-deoxyfutalosine ribohydrolase

**Comments:** The enzyme, found in several bacterial species, catalyses a step in a modified futalosine pathway

for menaquinone biosynthesis. While the enzyme from some organisms also has the activity of EC 3.2.2.9, adenosylhomocysteine nucleosidase, the enzyme from *Chlamydia trachomatis* is specific for

6-amino-6-deoxyfutalosine [164].

**References:** [1149, 1672, 65, 3117, 1927, 1454, 164]

[EC 3.2.2.30 created 2014]

EC 3.2.2.31

Accepted name: adenine glycosylase

**Reaction:** Hydrolyses free adenine bases from 7,8-dihydro-8-oxoguanine:adenine mismatched double-stranded

DNA, leaving an apurinic site.

**Other name(s):** mutY (gene name); A/G-specific adenine glycosylase

**Systematic name:** adenine-DNA deoxyribohydrolase (adenine-releasing)

**Comments:** The enzyme serves as a mismatch repair enzyme that works to correct 7,8-dihydro-8-

oxoguanine:adenine mispairs that arise in DNA when error-prone synthesis occurs past 7,8-dihydro-8-oxoguanine (GO) lesions in DNA. The enzyme excises the adenine of the mispair, producing an apurinic site sensitive to AP endonuclease activity. After removing the undamaged adenine the enzyme remains bound to the site to prevent EC 3.2.2.23 (MutM) from removing the GO lesion, which could lead to a double strand break. *In vitro* the enzyme is also active with adenine:guanine, adenine:cytosine, and adenine:7,8-dihydro-8-oxoadenine (AO) mispairs, removing the adenine in all

cases.

**References:** [96, 1901]

[EC 3.2.2.31 created 2018]

# EC 3.2.3 Hydrolysing S-glycosyl compounds (deleted sub-subclass)

[3.2.3.1 Transferred entry. thioglucosidase. Now EC 3.2.1.147, thioglucosidase]

[EC 3.2.3.1 created 1972, deleted 2001]

# EC 3.3 Acting on ether bonds

This subclass contains enzymes that act on ether bonds. It is subdivided into those hydrolysing thioether and trialkylsulfonium compounds (EC 3.3.1) and those acting on ethers (EC 3.3.2).

# EC 3.3.1 Thioether and trialkylsulfonium hydrolases

EC 3.3.1.1

Accepted name: adenosylhomocysteinase

**Reaction:** S-adenosyl-L-homocysteine +  $H_2O = L$ -homocysteine + adenosine

Other name(s): S-adenosylhomocysteine synthase; S-adenosylhomocysteine hydrolase (ambiguous); adenosylhomo-

cysteine hydrolase; S-adenosylhomocysteinase; SAHase; AdoHcyase

**Systematic name:** S-adenosyl-L-homocysteine hydrolase

**Comments:** The enzyme contains one tightly bound NAD<sup>+</sup> per subunit. This appears to bring about a transient

oxidation at C-3' of the 5'-deoxyadenosine residue, thus labilizing the thioether bond [2229] (for

mechanism, click here), cf. EC 5.5.1.4, inositol-3-phosphate synthase.

**References:** [559, 2229]

[EC 3.3.1.1 created 1961, modified 2004]

EC 3.3.1.2

**Accepted name:** S-adenosyl-L-methionine hydrolase (L-homoserine-forming)

**Reaction:** S-adenosyl-L-methionine +  $H_2O = L$ -homoserine + S-methyl-5'-thioadenosine

**Other name(s):** S-adenosylmethionine cleaving enzyme; methylmethionine-sulfonium-salt hydrolase; adenosyl-

methionine lyase; adenosylmethionine hydrolase; S-adenosylmethionine hydrolase; S-adenosyl-L-

methionine hydrolase

**Systematic name:** S-adenosyl-L-methionine hydrolase (L-homoserine-forming)

Comments: Also hydrolyses S-methyl-L-methionine to dimethyl sulfide and homoserine. cf. EC 3.13.1.8, S-

adenosyl-L-methionine hydrolase (adenosine-forming).

**References:** [1857]

[EC 3.3.1.2 created 1972, modified 1976, modified 2018]

[3.3.1.3 Deleted entry. ribosylhomocysteinase. This enzyme was transferred to EC 3.2.1.148, ribosylhomocysteinase, which has since been deleted. The activity is most probably attributable to EC 4.4.1.21, S-ribosylhomocysteine lyase]

[EC 3.3.1.3 created 1972, deleted 2001]

# EC 3.3.2 Ether hydrolases

#### EC 3.3.2.1

Accepted name: isochorismatase

**Reaction:** isochorismate  $+ H_2O = (2S,3S)-2,3$ -dihydroxy-2,3-dihydrobenzoate + pyruvate

Other name(s): 2,3-dihydro-2,3-dihydroxybenzoate synthase; 2,3-dihydroxy-2,3-dihydrobenzoate synthase; 2,3-

dihydroxy-2,3-dihydrobenzoic synthase

Systematic name: isochorismate pyruvate-hydrolase

**Comments:** The enzyme is involved in the biosynthesis of several siderophores, such as 2,3-

dihydroxybenzoylglycine, enterobactin, bacillibactin, and vibriobactin.

**References:** [3306]

[EC 3.3.2.1 created 1972]

## EC 3.3.2.2

Accepted name: lysoplasmalogenase

**Reaction:** (1) 1-(1-alkenyl)-sn-glycero-3-phosphocholine + H<sub>2</sub>O = an aldehyde + sn-glycero-3-phosphocholine

(2) 1-(1-alkenyl)-sn-glycero-3-phosphoethanolamine +  $H_2O$  = an aldehyde + sn-glycero-3-

phosphoethanolamine

Other name(s): alkenylglycerophosphocholine hydrolase; alkenylglycerophosphoethanolamine hydrolase; 1-(1-

alkenyl)-sn-glycero-3-phosphocholine aldehydohydrolase

Systematic name: lysoplasmalogen aldehydohydrolase

Comments: Lysoplasmalogenase is specific for the sn-2-deacylated (lyso) form of plasmalogen and catalyses hy-

drolytic cleavage of the vinyl ether bond, releasing a fatty aldehyde and sn-glycero-3-phosphocholine

or sn-glycero-3-phosphoethanolamine.

**References:** [3129, 691, 1000, 78, 3219]

[EC 3.3.2.2 created 1972, modified 1976, (EC 3.3.2.5 created 1984, incorporated 2016), modified 2016]

[3.3.2.3 Transferred entry. epoxide hydrolase. Now known to comprise two enzymes, microsomal epoxide hydrolase (EC 3.3.2.9) and soluble epoxide hydrolase (EC 3.3.2.10)]

[EC 3.3.2.3 created 1978, modified 1999, deleted 2006]

# EC 3.3.2.4

Accepted name: trans-epoxysuccinate hydrolase

**Reaction:** trans-2,3-epoxysuccinate + H<sub>2</sub>O = meso-tartrate **Other name(s):** trans-epoxysuccinate hydratase; tartrate epoxydase

**Systematic name:** *trans*-2,3-epoxysuccinate hydrolase

**Comments:** Acts on both optical isomers of the substrate.

**References:** [36]

[EC 3.3.2.4 created 1984]

[3.3.2.5 Transferred entry. alkenylglycerophosphoethanolamine hydrolase. Now included in EC 3.3.2.2, lysoplasmalogenase.]

[EC 3.3.2.5 created 1984, deleted 2016]

# EC 3.3.2.6

**Accepted name:** leukotriene-A<sub>4</sub> hydrolase

**Reaction:** leukotriene  $A_4 + H_2O = leukotriene B_4$ 

**Other name(s):** LTA<sub>4</sub> hydrolase; LTA<sub>4</sub>H; leukotriene A<sub>4</sub> hydrolase

**Systematic name:** (7E,9E,11Z,14Z)-(5S,6S)-5,6-epoxyicosa-7,9,11,14-tetraenoate hydrolase

Comments: This is a bifunctional zinc metalloprotease that displays both epoxide hydrolase and aminopeptidase

activities [2073, 2201]. It preferentially cleaves tripeptides at an arginyl bond, with dipeptides and tetrapeptides being poorer substrates [2201] (see EC 3.4.11.6, aminopeptidase B). It also converts leukotriene A<sub>4</sub> into leukotriene B<sub>4</sub>, unlike EC 3.3.2.10, soluble epoxide hydrolase, which converts leukotriene A<sub>4</sub> into 5,6-dihydroxy-7,9,11,14-icosatetraenoic acid [1017, 2073]. In vertebrates, five epoxide-hydrolase enzymes have been identified to date: EC 3.3.2.6 (leukotriene A<sub>4</sub> hydrolase), EC 3.3.2.7 (hepoxilin-epoxide hydrolase), EC 3.3.2.9 (microsomal epoxide hydrolase), EC 3.3.2.10 (solu-

ble epoxide hydrolase) and EC 3.3.2.11 (cholesterol-5,6-oxide hydrolase) [812].

**References:** [724, 1922, 1017, 2073, 812, 2201, 2161]

[EC 3.3.2.6 created 1989, modified 2006]

#### EC 3.3.2.7

**Accepted name:** hepoxilin-epoxide hydrolase **Reaction:** hepoxilin  $A_3 + H_2O = \text{trioxilin } A_3$ 

Other name(s): hepoxilin epoxide hydrolase; hepoxylin hydrolase; hepoxilin  $A_3$  hydrolase Systematic name:  $(5Z,9E,14Z)-(8\xi,11R,12S)-11,12$ -epoxy-8-hydroxyicosa-5,9,14-trienoate hydrolase

Comments: Converts hepoxilin  $A_3$  into trioxilin  $A_3$ . Highly specific for the substrate, having only slight activity

with other epoxides such as leukotriene A<sub>4</sub> and styrene oxide [2222]. Hepoxilin A<sub>3</sub> is an hydroxy-epoxide derivative of arachidonic acid that is formed via the 12-lipoxygenase pathway [2222]. It is probable that this enzyme plays a modulatory role in inflammation, vascular physiology, systemic glucose metabolism and neurological function [2073]. In vertebrates, five epoxide-hydrolase enzymes have been identified to date: EC 3.3.2.6 (leukotriene-A<sub>4</sub> hydrolase), EC 3.3.2.7 (hepoxilin-epoxide hydrolase), EC 3.3.2.9 (microsomal epoxide hydrolase), EC 3.3.2.10 (soluble epoxide hydrolase) and

EC 3.3.2.11 (cholesterol 5,6-oxide hydrolase) [812].

**References:** [2221, 2222, 812, 2073]

[EC 3.3.2.7 created 1992, modified 2006]

# EC 3.3.2.8

**Accepted name:** limonene-1,2-epoxide hydrolase

**Reaction:** 1,2-epoxymenth-8-ene +  $H_2O$  = menth-8-ene-1,2-diol

Other name(s): limonene oxide hydrolase
Systematic name: 1,2-epoxymenth-8-ene hydrolase

Comments: Involved in the monoterpene degradation pathway of the actinomycete *Rhodococcus erythropolis*. The

enzyme hydrolyses several alicyclic and 1-methyl-substituted epoxides, such as 1-methylcyclohexene oxide, indene oxide and cyclohexene oxide. It differs from the previously described epoxide hydrolases [EC 3.3.2.4 (*trans*-epoxysuccinate hydrolase), EC 3.3.2.6 (leukotriene-A<sub>4</sub> hydrolase), EC 3.3.2.7 (hepoxilin-epoxide hydrolase), EC 3.3.2.9 (microsomal epoxide hydrolase) and EC 3.3.2.10 (soluble epoxide hydrolase)] as it is not inhibited by 2-bromo-4'-nitroacetophenone, diethyl dicarbonate, 4-fluorochalcone oxide or 1,10-phenanthroline. Both enantiomers of menth-8-ene-1,2-diol [i.e.

(1R,2R,4S)-menth-8-ene-1,2-diol and (1S,2S,4R)-menth-8-ene-1,2-diol] are metabolized.

**References:** [3041, 138, 3042]

[EC 3.3.2.8 created 2001]

EC 3.3.2.9

**Accepted name:** microsomal epoxide hydrolase

**Reaction:** (1) *cis*-stilbene oxide +  $H_2O = (1R,2R)$ -1,2-diphenylethane-1,2-diol

(2) 1-(4-methoxyphenyl)-N-methyl-N-[(3-methyloxetan-3-yl)methyl]methanamine +  $H_2O = 2$ -([(4-methoxyphenyl)-N-methyl-N-[(3-methyloxetan-3-yl)methyl]methanamine

methoxyphenyl)methyl](methyl)aminomethyl)-2-methylpropane-1,3-diol

Other name(s): microsomal oxirane/oxetane hydrolase; epoxide hydratase (ambiguous); microsomal epoxide hy-

dratase (ambiguous); epoxide hydrase; microsomal epoxide hydrase; arene-oxide hydratase (ambiguous); benzo[a]pyrene-4,5-oxide hydratase; benzo(a)pyrene-4,5-epoxide hydratase; aryl epoxide

hydrase (ambiguous); cis-epoxide hydrolase; mEH; EPHX1 (gene name)

**Systematic name:** *cis*-stilbene-oxide hydrolase

**Comments:** This is a key hepatic enzyme that catalyses the hydrolytic ring opening of oxiranes (epoxides) and

oxetanes to give the corresponding diols. The enzyme is involved in the metabolism of numerous substrates including the stereoselective hydrolytic ring opening of 7-oxabicyclo[4.1.0]hepta-2,4-dienes (arene oxides) to the corresponding *trans*-dihydrodiols. The reaction proceeds via a triad mechanism and involves the formation of an hydroxyalkyl-enzyme intermediate. Five epoxide-hydrolase enzymes have been identified in vertebrates to date: EC 3.3.2.6 (leukotriene-A<sub>4</sub> hydrolase), EC 3.3.2.7 (hepoxilin-epoxide hydrolase), EC 3.3.2.9 (microsomal epoxide hydrolase), EC 3.3.2.10 (soluble

epoxide hydrolase) and EC 3.3.2.11 (cholesterol-5,6-oxide hydrolase).

**References:** [2142, 1301, 2140, 2141, 1743, 188, 812, 1977, 2073, 2950]

[EC 3.3.2.9 created 2006 (EC 3.3.2.3 created 1978, modified 1999, part incorporated 2006), modified 2017]

#### EC 3.3.2.10

**Accepted name:** soluble epoxide hydrolase **Reaction:** an epoxide  $+ H_2O = a$  glycol

**Other name(s):** epoxide hydrase (ambiguous); epoxide hydratase (ambiguous); arene-oxide hydratase (ambiguous);

aryl epoxide hydrase (ambiguous); trans-stilbene oxide hydrolase; sEH; cytosolic epoxide hydrolase

**Systematic name:** epoxide hydrolase

Comments: Catalyses the hydrolysis of trans-substituted epoxides, such as trans-stilbene oxide, as well as var-

ious aliphatic epoxides derived from fatty-acid metabolism [812]. It is involved in the metabolism of arachidonic epoxides (epoxyicosatrienoic acids; EETs) and linoleic acid epoxides. The EETs, which are endogenous chemical mediators, act at the vascular, renal and cardiac levels to regulate blood pressure [1977, 3312]. The enzyme from mammals is a bifunctional enzyme: the C-terminal domain exhibits epoxide-hydrolase activity and the N-terminal domain has the activity of EC 3.1.3.76, lipid-phosphate phosphatase [2074, 507]. Like EC 3.3.2.9, microsomal epoxide hydrolase, it is probable that the reaction involves the formation of an hydroxyalkyl—enzyme intermediate [1977, 1587]. The enzyme can also use leukotriene A4, the substrate of EC 3.3.2.6, leukotriene-A4 hydrolase, but it forms 5,6-dihydroxy-7,9,11,14-icosatetraenoic acid rather than leukotriene B4 as the product [1017, 2073]. In vertebrates, five epoxide-hydrolase enzymes have been identified to date: EC 3.3.2.6 (leukotriene-A4 hydrolase), EC 3.3.2.7 (hepoxilin-epoxide hydrolase), EC 3.3.2.9 (microsomal epoxide hydrolase), EC 3.3.2.10 (soluble epoxide hydrolase) and EC 3.3.2.11 (cholesterol 5,6-oxide hydrolase)

drolase) [812].

**References:** [2074, 507, 2140, 1977, 3312, 1587, 812, 3320, 1017, 2073]

[EC 3.3.2.10 created 2006 (EC 3.3.2.3 created 1978, part incorporated 2006)]

#### EC 3.3.2.11

**Accepted name:** cholesterol-5,6-oxide hydrolase

**Reaction:** (1)  $5.6\alpha$ -epoxy- $5\alpha$ -cholestan- $3\beta$ -ol +  $H_2O = 5\alpha$ -cholestane- $3\beta$ , $5\alpha$ , $6\beta$ -triol

(2) 5,6 $\beta$ -epoxy-5 $\beta$ -cholestan-3 $\beta$ -ol + H<sub>2</sub>O = 5 $\alpha$ -cholestane-3 $\beta$ ,5 $\alpha$ ,6 $\beta$ -triol

Other name(s): cholesterol-epoxide hydrolase; ChEH Systematic name: 5,6α-epoxy-5α-cholestan-3β-ol hydrolase

**Comments:** The enzyme appears to work equally well with either epoxide as substrate [2614]. The product is a

competitive inhibitor of the reaction. In vertebrates, five epoxide-hydrolase enzymes have been identified to date: EC 3.3.2.6 (leukotriene-A<sub>4</sub> hydrolase), EC 3.3.2.7 (hepoxilin-epoxide hydrolase), EC 3.3.2.9 (microsomal epoxide hydrolase), EC 3.3.2.10 (soluble epoxide hydrolase) and EC 3.3.2.11

(cholesterol 5,6-oxide hydrolase) [2614].

**References:** [1655, 2143, 2614, 812, 2073]

#### [EC 3.3.2.11 created 2006]

EC 3.3.2.12

**Accepted name:** oxepin-CoA hydrolase

**Reaction:** 2-oxepin-2(3*H*)-ylideneacetyl-CoA + H<sub>2</sub>O = 3-oxo-5,6-dehydrosuberyl-CoA semialdehyde

**Other name(s):** paaZ (gene name)

**Systematic name:** 2-oxepin-2(3*H*)-ylideneacetyl-CoA hydrolase

**Comments:** The enzyme from *Escherichia coli* is a bifunctional fusion protein that also catalyses EC 1.17.1.7, 3-

oxo-5,6-dehydrosuberyl-CoA semialdehyde dehydrogenase.Combined the two activities result in a two-step conversion of oxepin-CoA to 3-oxo-5,6-dehydrosuberyl-CoA, part of an aerobic phenylacetate degradation pathway [1,3,4]. The enzyme from *Escherichia coli* also exhibits enoyl-CoA hy-

dratase activity utilizing crotonyl-CoA as a substrate [2235].

**References:** [758, 2235, 1265, 2905]

[EC 3.3.2.12 created 2011 as EC 3.7.1.16, transferred 2013 to EC 3.3.2.12]

EC 3.3.2.13

Accepted name: chorismatase

**Reaction:** chorismate +  $H_2O = (4R,5R)$ -4,5-dihydroxycyclohexa-1(6),2-diene-1-carboxylate + pyruvate **Other name(s):** chorismate/3,4-dihydroxycyclohexa-1,5-dienoate synthase; *fkbO* (gene name); *rapK* (gene name)

**Systematic name:** chorismate pyruvate-hydrolase

Comments: The enzyme found in several bacterial species is involved in the biosynthesis of macrocyclic polyke-

tides.

**References:** [54, 1348]

[EC 3.3.2.13 created 2013]

EC 3.3.2.14

**Accepted name:** 2,4-dinitroanisole *O*-demethylase

**Reaction:** 2,4-dinitroanisole +  $H_2O$  = methanol + 2,4-dinitrophenol

**Other name(s):** 2,4-dinitroanisole ether hydrolase; *dnhA* (gene name); *dnhB* (gene name); DNAN demethylase

**Systematic name:** 2,4-dinitroanisole methanol hydrolase

Comments: The enzyme, characterized from the bacterium *Nocardioides* sp. JS1661, is involved in the degra-

dation of 2,4-dinitroanisole. Unlike other known *O*-demethylases, such as EC 1.14.99.15, 4-methoxybenzoate monooxygenase (*O*-demethylating), or EC 1.14.11.32, codeine 3-*O*-demethylase, it does not require oxygen or electron donors, and produces methanol rather than formaldehyde.

**References:** [762]

[EC 3.3.2.14 created 2015]

EC 3.3.2.15

**Accepted name:** *trans*-2,3-dihydro-3-hydroxyanthranilic acid synthase

**Reaction:** (2S)-2-amino-4-deoxychorismate +  $H_2O = (5S,6S)$ -6-amino-5-hydroxycyclohexa-1,3-diene-1-

carboxylate + pyruvate

Other name(s): isochorismatase (ambiguous); *phzD* (gene name)

Systematic name: (2S)-2-amino-4-deoxychorismate pyruvate-hydrolase

**Comments:** Isolated from the bacterium *Pseudomonas aeruginosa*. Involved in phenazine biosynthesis.

**References:** [1852, 2240]

[EC 3.3.2.15 created 2016]

# EC 3.4 Acting on peptide bonds (peptidases)

It is recommended that the term "peptidase" be used as being synonymous with "peptide hydrolase" for any enzyme that hydrolyses peptide bonds. Peptidases are recommended to be further divided into "exopeptidases" that act only near a terminus of a polypeptide chain and "endopeptidases" that act internally in polypeptide chains. The types of exopeptidases and endopeptidases are described more fully below. The usage of "peptidase", which is now recommended, is synonymous with "protease" as it was originally used [1] as a general term for both exopeptidases and endopeptidases, but it should be noted that previously, in Enzyme Nomenclature (1984), "peptidase" was restricted to the enzymes included in sub-subclasses EC 3.4.11 and EC 3.4.13-19, the exopeptidases. Also, the term "proteinase" used previously for the enzymes included in sub-subclasses EC 3.4.21-25 carried the same meaning as "endopeptidase", and has been replaced by "endopeptidase", for consistency.¡p;

The nomenclature of the peptidases is troublesome. Their specificity is commonly difficult to define, depending upon the nature of several amino-acid residues around the peptide bond to be hydrolysed and also on the conformation of the substrate's polypeptide chain. A classification involving the additional criterion of catalytic mechanism is therefore used.;p;

Two sets of sub-subclasses of peptidases are recognized, those of the exopeptidases (EC 3.4.11 and EC 3.4.13-19) and those of the endopeptidases (EC 3.4.21-25). The exopeptidases act only near the ends of polypeptide chains, and those acting at a free N-terminus liberate a single amino-acid residue (aminopeptidases; EC 3.4.11), or a dipeptide or a tripeptide (dipeptidyl-peptidases and tripeptidyl-peptidases; EC 3.4.14). The exopeptidases that act at a free C-terminus liberate a single residue (carboxypeptidases, EC 3.4.16-18), or a dipeptide (peptidyl-dipeptidases; EC 3.4.15). The carboxypeptidases are allocated to three groups on the basis of catalytic mechanism: the serine-type carboxypeptidases (EC 3.4.16), the metallocarboxypeptidases (EC 3.4.17) and the cysteine-type carboxypeptidases (EC 3.4.18). Other exopeptidases are specific for dipeptides (dipeptidases, EC 3.4.13), or for removal of terminal residues that are substituted, cyclized or linked by isopeptide bonds (peptide linkages other than those of alpha-carboxyl to alpha-amino groups) (omega peptidases; EC 3.4.19).;p;

The endopeptidases are divided into sub-subclasses on the basis of catalytic mechanism, and specificity is used only to identify individual enzymes within the groups. The sub-subclasses are: serine endopeptidases (EC 3.4.21), cysteine endopeptidases (EC 3.4.22), aspartic endopeptidases (EC 3.4.23), metalloendopeptidases (EC 3.4.24) and threonine endopeptidases (EC 3.4.25). ¡p¿

There are characteristic inhibitors of the members of each catalytic type of endopeptidase; to save space, these have not been listed separately for each individual enzyme but are reviewed in [2] and [3]. A general source of information on peptidases that similarly has not been cited for each individual enzyme is reference [4].;p;

In describing the specificity of peptidases, use is made of a model in which the catalytic site is considered to be flanked on one or both sides by specificity subsites, each able to accommodate the sidechain of a single amino-acid residue (based on [5]). These sites are numbered from the catalytic site, S1...Sn towards the N-terminus of the substrate, and S1'...Sn' towards the C-terminus. The residues they accommodate are numbered P1...Pn, and P1'...Pn', respectively, as follows: $p_{ij}$ 

```
Substrate: - P3 - P2 - P1 + P1'- P2'- P3'-jp;
Enzyme: - S3 - S2 - S1 * S1'- S2'- S3'-jp;
```

In this representation, the catalytic site of the enzyme is marked by an asterisk (\*). The peptide bond cleaved (the scissile bond) is indicated by the symbol '+' or a hyphen in the structural formula of the substrate, or a hyphen in the name of the enzyme.;p $\xi$ 

Finally, in describing the specificity of endopeptidases, the term oligopeptidase' is used to refer to those that act optimally on substrates smaller than proteins.;  $p_{ij}$ 

Families of peptidases are referred to by use of the numbering system of Rawlings & Barrett [6,7].;p;

# References;p;

- 1. Grassmann, W. & Dyckerhoff, H. Über die Proteinase und die Polypeptidase der Hefe. 13. Abhandlung über Pflanzenproteasen in der von R. Willstätter und Mitarbeitern begonnenen Untersuchungsreihe. Hoppe-Seyler's Z. Physiol. Chem. 179 (1928) 41-78.
  - 2. Barrett, A. J. & Salvesen, G. S. (eds) Proteinase Inhibitors, Elsevier Science Publishers, Amsterdam (1986).
  - 3. Beynon, R. J. & Bond, J. S. Proteolytic Enzymes. A Practical Approach. IRL Press, Oxford (1989).
  - 4. Barrett, A. J., Rawlings, N. D. & Woessner, J. F. (eds) Handbook of Proteolytic Enzymes, Academic Press, London (1998).
- 5. Berger, A. and Schechter, I. Mapping the active site of papain with the aid of peptide substrates and inhibitors. Philos. Trans. R. Soc. London, Ser. B Biol. Sci. 257 (1970) 249-264.
- 6. Rawlings, N.D. and Barrett, A.J. In: Methods Enzymol. 244 (1994) 19-61 and 461-486; Methods Enzymol. 248 (1995) 105-120 and 183-228.

7. Rawlings, N. D. and Barrett, A. J. MEROPS: the peptidase database. Nucleic Acids Res. 27 (1999) 325-331.

[EC 3.4.1.4 created 1965, deleted 1972]

# EC 3.4.1 $\alpha$ -Amino-acyl-peptide hydrolases (deleted sub-subclass)

[3.4.1.1	Transferred entry. leucyl aminopeptidase. Now EC 3.4.11.1, leucyl aminopeptidase]
	[EC 3.4.1.1 created 1961, deleted 1972]
[3.4.1.2	Transferred entry. aminopeptidase. Now EC 3.4.11.2, membrane alanyl aminopeptidase]
	[EC 3.4.1.2 created 1961, deleted 1972]
[3.4.1.3	Transferred entry. aminotripeptidase. Now EC 3.4.11.4, tripeptide aminopeptidase]
	[EC 3.4.1.3 created 1961, deleted 1972]
[3.4.1.4	Transferred entry. proline iminopeptidase. Now EC 3.4.11.5, prolyl aminopeptidase]

# EC 3.4.2 Peptidyl-amino-acid hydrolases (deleted sub-subclass)

[3.4.2.1	Transferred entry. carboxypeptidase A. Now EC 3.4.17.1, carboxypeptidase A]
	[EC 3.4.2.1 created 1961, deleted 1972]
[3.4.2.2	Transferred entry. carboxypeptidase B. Now EC 3.4.17.2, carboxypeptidase B]
	[EC 3.4.2.2 created 1961, deleted 1972]
[3.4.2.3	Transferred entry. yeast carboxypeptidase. Now EC 3.4.17.4, Gly-Xaa carboxypeptidase]
	[EC 3.4.2.3 created 1961, deleted 1972]

# EC 3.4.3 Dipeptide hydrolases (deleted sub-subclass)

[3.4.3.1	Transferred entry. glycyl-glycine dipeptidase. Now EC 3.4.13.18, cytosol nonspecific dipeptidase]
	[EC 3.4.3.1 created 1961, deleted 1972]
[3.4.3.2	Transferred entry. glycyl-leucine dipeptidase. Now EC 3.4.13.18, cytosol nonspecific dipeptidase]
	[EC 3.4.3.2 created 1961, deleted 1972]
[3.4.3.3	Transferred entry. aminoacyl-histidine dipeptidase. Now EC 3.4.13.3, Xaa-His dipeptidase]
	[EC 3.4.3.3 created 1961, deleted 1972]
[3.4.3.4	Transferred entry. aminoacyl-methylhistidine dipeptidase. Now EC 3.4.13.5, Xaa-methyl-His dipeptidase]
	[EC 3.4.3.4 created 1961, deleted 1972]
[3.4.3.5	Transferred entry. cysteinylglycine dipeptidase. Now EC 3.4.11.2, membrane alanyl aminopeptidase]
	[EC 3.4.3.5 created 1961, deleted 1972]
[3.4.3.6	Transferred entry. iminodipeptidase. Now EC 3.4.13.18, cytosol nonspecific dipeptidase]
	[EC 3.4.3.6 created 1961, deleted 1972]
[3.4.3.7	Transferred entry. iminodipeptidase. Now EC 3.4.13.9, Xaa-Pro dipeptidase]

[EC 3.4.3.7 created 1961, deleted 1972]

# EC 3.4.4 Peptidyl peptide hydrolases (deleted sub-subclass)

[3.4.4.1	Transferred entry. pepsin. Now EC 3.4.23.1, pepsin A]
	[EC 3.4.4.1 created 1961, deleted 1972]
[3.4.4.2	Transferred entry. pepsin B. Now EC 3.4.23.2, pepsin B]
	[EC 3.4.4.2 created 1961, deleted 1972]
[3.4.4.3	Transferred entry. rennin. Now EC 3.4.23.4, chymosin]
	[EC 3.4.4.3 created 1961, deleted 1972]
[3.4.4.4	Transferred entry. trypsin. Now EC 3.4.21.4, trypsin]
	[EC 3.4.4.4 created 1961, deleted 1972]
[3.4.4.5	Transferred entry. chymotrypsin. Now EC 3.4.21.1, chymotrypsin]
	[EC 3.4.4.5 created 1961, deleted 1972]
[3.4.4.6	Transferred entry. chymotrypsin B. Now EC 3.4.21.1, chymotrypsin]
	[EC 3.4.4.6 created 1961, deleted 1972]
[3.4.4.7	Transferred entry. elastase. Now covered by EC 3.4.21.36, pancreatic elastase and EC 3.4.21.37, leukocyte elastase]
	[EC 3.4.4.7 created 1961, deleted 1972]
[3.4.4.8	Transferred entry. enteropeptidase. Now EC 3.4.21.9, enteropeptidase]
	[EC 3.4.4.8 created 1961, deleted 1972]
[3.4.4.9	Transferred entry. cathepsin C. Now EC 3.4.14.1, dipeptidyl-peptidase I]
	[EC 3.4.4.9 created 1961, deleted 1972]
[3.4.4.10	Transferred entry. papain. Now EC 3.4.22.2, papain]
	[EC 3.4.4.10 created 1961, deleted 1972]
[3.4.4.11	Transferred entry. chymopapain. Now EC 3.4.22.6, chymopapain]
	[EC 3.4.4.11 created 1961, deleted 1972]
[3.4.4.12	Transferred entry. ficin. Now EC 3.4.22.3, ficain]
	[EC 3.4.4.12 created 1961, deleted 1972]
[3.4.4.13	Transferred entry. thrombin. Now EC 3.4.21.5, thrombin]
	[EC 3.4.4.13 created 1961, deleted 1972]
[3.4.4.14	Transferred entry. plasmin. Now EC 3.4.21.7, plasmin]
	[EC 3.4.4.14 created 1961, deleted 1972]
[3.4.4.15	Transferred entry. renin. Now EC 3.4.23.15, renin]
	[EC 3.4.4.15 created 1961, deleted 1972]
[3.4.4.16	Transferred entry. subtilopeptidase A. Now covered by the microbial serine proteinases EC 3.4.21.62 (subtil-

[3.4.4.16 Transferred entry. subtilopeptidase A. Now covered by the microbial serine proteinases EC 3.4.21.62 (subtilisin), EC 3.4.21.63 (oryzin), EC 3.4.21.64 (endopeptidase K), EC 3.4.21.65 (thermomycolin), EC 3.4.21.66 (thermitase) and EC 3.4.21.67 (ndopeptidase So)]

[EC 3.4.4.16 created 1961, deleted 1972]

[3.4.4.17 Transferred entry. aspergillopeptidase A. Now covered by the microbial aspartic proteinases EC 3.4.23.20 (penicillopepsin), EC 3.4.23.21 (rhizopuspepsin), EC 3.4.23.22 (endothiapepsin), EC 3.4.23.23 (mucorpepsin), EC 3.4.23.24 (candidapepsin), EC 3.4.23.25 (saccharopepsin), EC 3.4.23.26 (rhodotorulapepsin), EC 3.4.21.103 (physarolisin), EC 3.4.23.28 (acrocylindropepsin), EC 3.4.23.29 (polyporopepsin) and EC 3.4.23.30 (pycnoporopepsin)]

[EC 3.4.4.17 created 1961, deleted 1972]

[3.4.4.18 Transferred entry. streptococcus peptidase A. Now EC 3.4.22.10, streptopain]

[EC 3.4.4.18 created 1961, deleted 1972]

[3.4.4.19 Transferred entry, clostridiopeptidase A. Now EC 3.4.24.3, microbial collagenase]

[EC 3.4.4.19 created 1961, deleted 1972]

[3.4.4.20 Transferred entry. clostridiopeptidase B. Now EC 3.4.22.8, clostripain]

[EC 3.4.4.20 created 1961, deleted 1972]

[3.4.4.21 Transferred entry. kallikrein. Now EC 3.4.21.34 (plasma kallikrein) and EC 3.4.21.35 (tissue kallikrein)]

[EC 3.4.4.21 created 1965, deleted 1972]

[3.4.4.22 Transferred entry. now EC 3.4.23.3, gastricsin]

[EC 3.4.4.22 created 1965, deleted 1972]

[3.4.4.23 Transferred entry. now EC 3.4.23.5, cathepsin D]

[EC 3.4.4.23 created 1965, deleted 1972]

[3.4.4.24 Transferred entry, now covered by EC 3.4.22.32 (stem bromelain) and EC 3.4.22.33 (fruit bromelain)]

[EC 3.4.4.24 created 1965, deleted 1972]

[3.4.4.25 Deleted entry. Streptomyces alkalophilic keratinase]

[EC 3.4.4.25 created 1965, deleted 1972]

# EC 3.4.11 Aminopeptidases

EC 3.4.11.1

**Accepted name:** leucyl aminopeptidase

**Reaction:** Release of an N-terminal amino acid, Xaa—Yaa-, in which Xaa is preferably Leu, but may be other

amino acids including Pro although not Arg or Lys, and Yaa may be Pro. Amino acid amides and

methyl esters are also readily hydrolysed, but rates on arylamides are exceedingly low

Other name(s): leucine aminopeptidase; leucyl peptidase; peptidase S; cytosol aminopeptidase; cathepsin III; L-

leucine aminopeptidase; leucinaminopeptidase; leucinamide aminopeptidase; FTBL proteins; pro-

teinates FTBL; aminopeptidase II; aminopeptidase III; aminopeptidase I

**Comments:** A zinc enzyme isolated from pig kidney and cattle lens; activated by heavy metal ions. Type example

of peptidase family M17.

**References:** [1142, 574, 3047]

[EC 3.4.11.1 created 1961 as EC 3.4.1.1, transferred 1972 to EC 3.4.11.1]

EC 3.4.11.2

Accepted name: membrane alanyl aminopeptidase

**Reaction:** Release of an N-terminal amino acid, Xaa—Yaa- from a peptide, amide or arylamide. Xaa is prefer-

ably Ala, but may be most amino acids including Pro (slow action). When a terminal hydrophobic residue is followed by a prolyl residue, the two may be released as an intact Xaa-Pro dipeptide

Other name(s): microsomal aminopeptidase; aminopeptidase M; aminopeptidase N; particle-bound aminopepti-

dase; amino-oligopeptidase; alanine aminopeptidase; membrane aminopeptidase I; pseudo leucine aminopeptidase; alanyl aminopeptidase; alanine-specific aminopeptidase; cysteinylglycine dipepti-

dase; cysteinylglycinase; L-alanine aminopeptidase; CD13

**Comments:** A zinc enzyme, not activated by heavy metal ions. Type example of peptidase family M1.

**References:** [3081, 1458, 968, 2679, 756]

[EC 3.4.11.2 created 1961 as EC 3.4.1.2, transferred 1972 to EC 3.4.11.2 (EC 3.4.13.6 created 1961 as EC 3.4.3.5, transferred 1972 to EC 3.4.13.6, incorporated 1997)]

#### EC 3.4.11.3

Accepted name: cystinyl aminopeptidase

Reaction: Release of an N-terminal amino acid, Cys—Xaa-, in which the half-cystine residue is involved in

a disulfide loop, notably in oxytocin or vasopressin. Hydrolysis rates on a range of aminoacyl ary-

lamides exceed that for the cystinyl derivative, however [4]

Other name(s): cystyl-aminopeptidase; oxytocinase; cystine aminopeptidase; L-cystine aminopeptidase; oxytocin pep-

tidase; vasopresssinase

Comments: A zinc-containing sialoglycoprotein in peptidase family M1 (membrane alanyl aminopeptidase fam-

ily)

**References:** [2676, 2677, 3285, 2507]

[EC 3.4.11.3 created 1972]

#### EC 3.4.11.4

**Accepted name:** tripeptide aminopeptidase

**Reaction:** Release of the N-terminal residue from a tripeptide

Other name(s): tripeptidase; aminotripeptidase; aminoexotripeptidase; lymphopeptidase; imidoendopeptidase; pepti-

dase B; alanine-phenylalanine-proline arylamidase; peptidase T

**Comments:** A zinc enzyme, widely distributed in mammalian tissues. Formerly EC 3.4.1.3

**References:** [641, 2486]

[EC 3.4.11.4 created 1961 as EC 3.4.1.3, transferred 1972 to EC 3.4.11.4]

#### EC 3.4.11.5

Accepted name: prolyl aminopeptidase

**Reaction:** Release of N-terminal proline from a peptide

Other name(s): proline aminopeptidase; Pro-X aminopeptidase; cytosol aminopeptidase V; proline iminopeptidase

Comments: A Mn<sup>2+</sup>-requiring enzyme present in the cytosol of mammalian and microbial cells. In contrast to the

mammalian form, the bacterial form of the enzyme (type example of peptidase family S33) hydrolyses both polyproline and prolyl-2-naphthylamide. The mammalian enzyme, which is not specific for

prolyl bonds, is possibly identical with EC 3.4.11.1, leucyl aminopeptidase.

**References:** [2530, 2112, 3004]

[EC 3.4.11.5 created 1965 as EC 3.4.1.4, transferred 1972 to EC 3.4.11.5]

## EC 3.4.11.6

**Accepted name:** aminopeptidase B

**Reaction:** Release of N-terminal Arg and Lys from oligopeptides when P1' is not Pro. Also acts on arylamides

of Arg and Lys

Other name(s): arylamidase II; arginine aminopeptidase; arginyl aminopeptidase; Cl<sup>-</sup>-activated arginine aminopepti-

dase; cytosol aminopeptidase IV; L-arginine aminopeptidase

**Comments:** Cytosolic or membrane-associated enzyme from mammalian tissues, activated by chloride ions and

low concentrations of thiol compounds. This is one of the activities of the bifunctional enzyme EC

3.3.2.6 (membrane alanyl aminopeptidase family) [855, 349].

**References:** [869, 187, 350, 855, 349, 2201]

[EC 3.4.11.6 created 1972, modified 1997]

EC 3.4.11.7

Accepted name: glutamyl aminopeptidase

Reaction: Release of N-terminal glutamate (and to a lesser extent aspartate) from a peptide

Other name(s): aminopeptidase A; aspartate aminopeptidase; angiotensinase A; glutamyl peptidase; Ca<sup>2+</sup>-activated

glutamate aminopeptidase; membrane aminopeptidase II; antigen BP-1/6C3 of mouse B lymphocytes;

L-aspartate aminopeptidase; angiotensinase A2

**Comments:** Ca<sup>2+</sup>-activated and generally membrane-bound. A zinc-metallopeptidase in family M1 (membrane

alanyl aminopeptidase family)

**References:** [929, 446, 536, 2926, 3221]

[EC 3.4.11.7 created 1972]

[3.4.11.8 Transferred entry. pyroglutamyl aminopeptidase. Now EC 3.4.19.3, pyroglutamyl-peptidase I]

[EC 3.4.11.8 created 1972, deleted 1981]

EC 3.4.11.9

Accepted name: Xaa-Pro aminopeptidase

Reaction: Release of any N-terminal amino acid, including proline, that is linked to proline, even from a dipep-

tide or tripeptide

**Other name(s):** proline aminopeptidase; aminopeptidase P; aminoacylproline aminopeptidase; X-Pro aminopeptidase

**Comments:** A Mn<sup>2+</sup>-dependent, generally membrane-bound enzyme present in both mammalian and bacterial

cells. In peptidase family M24 (methionyl aminopeptidase family)

**References:** [3269, 3268, 782, 2199, 1172]

[EC 3.4.11.9 created 1972]

EC 3.4.11.10

**Accepted name:** bacterial leucyl aminopeptidase

Reaction: Release of an N-terminal amino acid, preferentially leucine, but not glutamic or aspartic acids

**Other name(s):** Aeromonas proteolytica aminopeptidase

**Comments:** A zinc enzyme. Forms of the enzyme have been isolated from *Aeromonas proteolytica*, *Escherichia* 

coli and Streptococcus thermophilus. Examples are known from peptidase families M17 and M28 (of

leucyl aminopeptidase and aminopeptidase Y, respectively)

**References:** [2323, 604, 2347]

[EC 3.4.11.10 created 1972]

[3.4.11.11 Deleted entry. aminopeptidase]

[EC 3.4.11.11 created 1978, deleted 1992]

[3.4.11.12 Deleted entry. thermophilic aminopeptidase]

[EC 3.4.11.12 created 1978, deleted 1997]

EC 3.4.11.13

Accepted name: clostridial aminopeptidase

**Reaction:** Release of any N-terminal amino acid, including proline and hydroxyproline, but no cleavage of Xaa-

Pro-

**Other name(s):** Clostridium histolyticum aminopeptidase

**Comments:** A secreted enzyme from *Clostridium histolyticum*, requiring Mn<sup>2+</sup> or Co<sup>2+</sup>

**References:** [1427, 1428, 1429]

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## [EC 3.4.11.13 created 1978]

EC 3.4.11.14

Accepted name: cytosol alanyl aminopeptidase

Reaction: Release of an N-terminal amino acid, preferentially alanine, from a wide range of peptides, amides

and arylamides

Other name(s): arylamidase; aminopolypeptidase; thiol-activated aminopeptidase; human liver aminopeptidase;

puromycin-sensitive aminopeptidase; soluble alanyl aminopeptidase; cytosol aminopeptidase III; ala-

nine aminopeptidase

**Comments:** A puromycin-sensitive, Co<sup>2+</sup>-activated zinc-sialoglycoprotein that is generally cytosolic. Multiple

forms are widely distributed in mammalian tissues and body fluids. In peptidase family M1 (mem-

brane alanyl aminopeptidase family)

**References:** [2755, 1387, 2654]

[EC 3.4.11.14 created 1978]

EC 3.4.11.15

Accepted name: aminopeptidase Y

Reaction: Preferentially, release of N-terminal lysine

Other name(s): aminopeptidase Co; aminopeptidase (cobalt-activated); lysyl aminopeptidase

**Comments:** Requires  $Co^{2+}$ ; inhibited by  $Zn^{2+}$  and  $Mn^{2+}$ . An enzyme best known from *Saccharomyces cerevisiae* 

that hydrolyses Lys-NHPhNO2 and, more slowly, Arg-NHPhNO2. Type example of peptidase family

M28

**References:** [8, 3270, 2097]

[EC 3.4.11.15 created 1989, modified 1997]

EC 3.4.11.16

**Accepted name:** Xaa-Trp aminopeptidase

Reaction: Release of a variety of N-terminal residues (especially glutamate and leucine) from peptides, provided

tryptophan (or at least phenylalanine or tyrosine) is the penultimate residue. Also acts on Glu-Trp,

Leu—Trp and a number of other dipeptides

**Other name(s):** aminopeptidase W; aminopeptidase X-Trp; X-Trp aminopeptidase

**Comments:** A glycoprotein containing Zn<sup>2+</sup>, from renal and intestinal brush border membranes

**References:** [885, 886]

[EC 3.4.11.16 created 1989]

EC 3.4.11.17

Accepted name: tryptophanyl aminopeptidase

**Reaction:** Preferential release of N-terminal tryptophan

**Other name(s):** tryptophan aminopeptidase; L-tryptophan aminopeptidase

**Comments:** From *Trichosporon cutaneum*. Also acts on L-tryptophanamide. Requires Mn<sup>2+</sup>

**References:** [1283]

[EC 3.4.11.17 created 1989]

EC 3.4.11.18

**Accepted name:** methionyl aminopeptidase

**Reaction:** Release of N-terminal amino acids, preferentially methionine, from peptides and arylamides

Other name(s): methionine aminopeptidase; peptidase M; L-methionine aminopeptidase; MAP

Comments: This membrane-bound enzyme, which is present in both prokaryotes and eukaryotes, releases the ini-

tiator methionine from nascent peptides. The activity is dependent on the identity of the second, third and fourth amino acid residues of the target protein, but in general the enzyme acts only when the

penultimate residue is small and uncharged (e.g. Gly, Ala, Cys, Ser, Thr, and Val).

**References:** [3298, 2985, 806, 189, 2446]

[EC 3.4.11.18 created 1990]

EC 3.4.11.19

**Accepted name:** D-stereospecific aminopeptidase

**Reaction:** Release of an N-terminal D-amino acid from a peptide, Xaa—Yaa-, in which Xaa is preferably D-Ala,

D-Ser or D-Thr. D-Amino acid amides and methyl esters also are hydrolysed, as is glycine amide

Other name(s): D-aminopeptidase

Comments: Known from the bacterium Ochrobactrum anthropi. In peptidase family S12 (D-Ala-D-Ala car-

boxypeptidase family) [82]

**References:** [83, 82]

[EC 3.4.11.19 created 1993]

EC 3.4.11.20

**Accepted name:** aminopeptidase Ey

Reaction: Differs from other aminopeptidases in broad specificity for amino acids in the P1 position and the

ability to hydrolyse peptides of four or five residues that contain Pro in the P1' position

Comments: A zinc glycoprotein in peptidase family M1 (membrane alanyl aminopeptidase family), composed of

two 150 kDa subunits. From the plasma fraction of hen egg yolk

**References:** [1220, 2875, 2874]

[EC 3.4.11.20 created 1995]

EC 3.4.11.21

**Accepted name:** aspartyl aminopeptidase

**Reaction:** Release of an N-terminal aspartate or glutamate from a peptide, with a preference for aspartate **Comments:** Aminoacyl-arylamides are poor substrates. This is an abundant cytosolic enzyme in mammalian cells,

in peptidase family M18 of aminopeptidase I

**References:** [1416, 3186]

[EC 3.4.11.21 created 2000]

EC 3.4.11.22

Accepted name: aminopeptidase I

**Reaction:** Release of an N-terminal amino acid, preferably a neutral or hydrophobic one, from a polypeptide.

Aminoacyl-arylamides are poor substrates

Other name(s): aminopeptidase III; aminopeptidase yscI; leucine aminopeptidase IV; yeast aminopeptidase I

**Comments:** A 640-kDa, dodecameric enzyme best known as the major vacuolar aminopeptidase of yeast, *Sac*-

*charomyces cervisiae*, in which species it was first given the name aminopeptidase I (one), amongst others. Activity is stimulated by both Zn<sup>2+</sup> and Cl<sup>-</sup> ions. Type example of peptidase family M18

**References:** [1330, 1892, 398, 2137]

[EC 3.4.11.22 created 1997]

EC 3.4.11.23

Accepted name: PepB aminopeptidase

Reaction: Release of an N-terminal amino acid, Xaa, from a peptide or arylamide. Xaa is preferably Glu or Asp

but may be other amino acids, including Leu, Met, His, Cys and Gln

**Other name(s):** Salmonella enterica serovar Typhimurium peptidase B

**Comments:** A 270-kDa protein composed of six 46.3-kDa subunits. The pH optimum is in the alkaline range and

activity is stimulated by KCl. In peptidase family M17.

**References:** [1835]

[EC 3.4.11.23 created 2003]

EC 3.4.11.24

Accepted name: aminopeptidase S

**Reaction:** Release of an N-terminal amino acid with a preference for large hydrophobic amino-terminus residues **Other name(s):** Mername-AA022 peptidase; SGAP; aminopeptidase (*Streptomyces griseus*); *Streptomyces griseus* 

aminopeptidase; S. griseus AP; double-zinc aminopeptidase

**Comments:** Aminopeptidases are associated with many biological functions, including protein maturation, pro-

tein degradation, cell-cycle control and hormone-level regulation [72, 863]. This enzyme contains two zinc molecules in its active site and is activated by Ca<sup>2+</sup> [863]. In the presence of Ca<sup>2+</sup>, the best substrates are Leu-Phe, Leu-Ser, Leu-pNA (aminoacyl-*p*-nitroanilide), Phe-Phe-Phe and Phe-Phe [72]. Peptides with proline in the P1' position are not substrates [72]. Belongs in peptidase family M28.

**References:** [2744, 190, 72, 863, 921]

[EC 3.4.11.24 created 2008]

EC 3.4.11.25

**Accepted name:** β-peptidyl aminopeptidase

**Reaction:** Cleaves N-terminal β-homoamino acids from peptides composed of 2 to 6 amino acids

**Other name(s):** BapA (ambiguous)

**Comments:** Sphingosinicella xenopeptidilytica strain 3-2W4 is able to utilize the  $\beta$ -peptides  $\beta$ -homoVal- $\beta$ -

homoAla-β-homoLeu and β-homoAla-β-homoLeu as sole carbon and energy sources [898].

**References:** [1095, 898, 897, 1094]

[EC 3.4.11.25 created 2011]

EC 3.4.11.26

**Accepted name:** intermediate cleaving peptidase 55

**Reaction:** The enzyme cleaves the Pro<sup>36</sup>-Pro<sup>37</sup> bond of cysteine desulfurase (EC 2.8.1.7) removing three amino

acid residues (Tyr-Ser-Pro) from the N-terminus after cleavage by mitochondrial processing pepti-

dase.

Other name(s): Icp55; mitochondrial intermediate cleaving peptidase 55 kDa

Comments: Icp55 removes the destabilizing N-terminal amino acid residues that are left after cleavage by the mi-

tochondrial processing peptidase, leading to the stabilisation of the substrate. The enzyme can remove single amino acids or a short peptide, as in the case of cysteine desulfurase (EC 2.8.1.7), where three

amino acids are removed.

**References:** [2020, 3077]

[EC 3.4.11.26 created 2011]

# EC 3.4.12 Peptidylamino-acid hydrolases or acylamino-acid hydrolases (deleted sub-subclass)

[3.4.12.1 Transferred entry, now EC 3.4.16.5 (carboxypeptidase C) and EC 3.4.16.6 (carboxypeptidase D)]

[EC 3.4.12.1 created 1972, deleted 1978]

[3.4.12.2 Transferred entry. now EC 3.4.17.1, carboxypeptidase A]

[EC 3.4.12.2 created 1972, deleted 1978] [3.4.12.3 *Transferred entry. now EC 3.4.17.2, carboxypeptidase B*] [EC 3.4.12.3 created 1972, deleted 1978] [3.4.12.4 Transferred entry. now EC 3.4.16.2, lysosomal Pro-Xaa carboxypeptidase] [EC 3.4.12.4 created 1972, modified 1976, deleted 1978] [3.4.12.5 Transferred entry. now EC 3.5.1.28, N-acetylmuramoyl-L-alanine amidase] [EC 3.4.12.5 created 1972, deleted 1978] [3.4.12.6] *Transferred entry. now EC 3.4.17.8, muramoyl-pentapeptidase carboxypeptidase]* [EC 3.4.12.6 created 1972, deleted 1978] [3.4.12.7 Transferred entry. now EC 3.4.17.3, lysine carboxypeptidase] [EC 3.4.12.7 created 1972, deleted 1978] [3.4.12.8] Transferred entry. now EC 3.4.17.4, Gly-Xaa carboxypeptidase] [EC 3.4.12.8 created 1972, deleted 1978] [3.4.12.9] Deleted entry. aspartate carboxypeptidase] [EC 3.4.12.9 created 1972, deleted 1978] [3.4.12.10 Transferred entry. now EC 3.4.19.9, γ-glutamyl hydrolase] [EC 3.4.12.10 created 1972, modified 1976, deleted 1978] [3.4.12.11 Transferred entry. now EC 3.4.17.6, alanine carboxypeptidase] [EC 3.4.12.11 created 1972, deleted 1978] Transferred entry. now EC 3.4.16.5 (carboxypeptidase C) and EC 3.4.16.6 (carboxypeptidase D)] [3.4.12.12 [EC 3.4.12.12 created 1972, deleted 1978] [3.4.12.13 Deleted entry. Y-glutamylglutamate carboxypeptidase] [EC 3.4.12.13 created 1975, modified 1976, deleted 1978]

# EC 3.4.13 Dipeptidases

[3.4.13.1 Transferred entry. glycyl-glycine dipeptidase. Now EC 3.4.13.18, cytosol nonspecific dipeptidase]

[EC 3.4.13.1 created 1972, deleted 1978 [transferred to EC 3.4.13.11, deleted 1992]]

[3.4.13.2 Transferred entry. glycyl-leucine dipeptidase. Now EC 3.4.13.18, cytosol nonspecific dipeptidase]

[EC 3.4.13.2 created 1972, deleted 1978 [transferred to EC 3.4.13.11, deleted 1992]]

[3.4.13.3 Deleted entry. Xaa-His dipeptidase. The activity is covered by EC 3.4.13.18, cytosol nonspecific dipeptidase and EC 3.4.13.20,  $\beta$ -Ala-His dipeptidase.]

[EC 3.4.13.3 created 1961 as EC 3.4.3.3, transferred 1972 to EC 3.4.13.3, modified 1989 (EC 3.4.13.13 created 1981, incorporated 1992), deleted 2011]

#### EC 3.4.13.4

**Accepted name:** Xaa-Arg dipeptidase

**Reaction:** Preferential hydrolysis of Xaa—Arg, Xaa—Lys or Xaa—ornithine dipeptides

Other name(s): aminoacyl-lysine dipeptidase;  $N^2$ -(4-amino-butyryl)-L-lysine hydrolase; X-Arg dipeptidase

**Comments:** Widely distributed in mammals

**References:** [1552]

## [EC 3.4.13.4 created 1972]

EC 3.4.13.5

**Accepted name:** Xaa-methyl-His dipeptidase

**Reaction:** Hydrolysis of anserine ( $\beta$ -alanyl $+N^{\pi}$ -methyl-L-histidine), carnosine, homocarnosine,

glycyl—leucine and other dipeptides with broad specificity

Other name(s): anserinase; aminoacyl-methylhistidine dipeptidase; acetylhistidine deacetylase; N-acetylhistidine

deacetylase; α-N-acetyl-L-histidine aminohydrolase; X-methyl-His dipeptidase

**References:** [1338, 168, 1645]

[EC 3.4.13.5 created 1961 as EC 3.4.3.4, transferred 1972 to EC 3.4.13.5, modified 1981 (EC 3.5.1.34 created 1972, incorporated 1981)]

[3.4.13.6 Transferred entry. Cys-Gly dipeptidase. Now EC 3.4.11.2, membrane alanyl aminopeptidase]

[EC 3.4.13.6 created 1961 as EC 3.4.3.5, transferred 1972 to EC 3.4.13.6]

EC 3.4.13.7

Accepted name: Glu-Glu dipeptidase

**Reaction:** Hydrolysis of the Glu—Glu dipeptide

Other name(s):  $\alpha$ -glutamyl-glutamate dipeptidase; glutamylglutamic arylamidase

**Comments:** It is unclear whether the specificity of this enzyme extends to other  $\alpha$ -glutamyl dipeptides

**References:** [2322]

[EC 3.4.13.7 created 1972]

[3.4.13.8] Transferred entry. Pro-X dipeptidase. Now EC 3.4.13.18, cytosol nonspecific dipeptidase.]

[EC 3.4.13.8 created 1961 as EC 3.4.3.6, transferred 1972 to EC 3.4.13.8]

EC 3.4.13.9

Accepted name: Xaa-Pro dipeptidase

**Reaction:** Hydrolysis of Xaa—Pro dipeptides; also acts on aminoacyl-hydroxyproline analogs. No action on

Pro-Pro

Other name(s): prolidase; imidodipeptidase; proline dipeptidase; peptidase D; γ-peptidase; X-Pro dipeptidase

Comments: A Mn<sup>2+</sup>-activated enzyme, in peptidase family M24 (methionyl aminopeptidase family); cytosolic

from most animal tissues.

**References:** [549, 2680, 127, 311]

[EC 3.4.13.9 created 1961 as EC 3.4.3.7, transferred 1972 to EC 3.4.13.9]

[3.4.13.10 Transferred entry. \(\beta\)-aspartyldipeptidase. Now EC 3.4.19.5, \(\beta\)-aspartyl-peptidase]

[EC 3.4.13.10 created 1972, deleted 1992]

[3.4.13.11 Transferred entry. dipeptidase. Now EC 3.4.13.19, membrane dipeptidase]

[EC 3.4.13.11 created 1972, deleted 1992]

EC 3.4.13.12

Accepted name: Met-Xaa dipeptidase

**Reaction:** Hydrolysis of Met—Xaa dipeptides

Other name(s): methionyl dipeptidase; dipeptidase M; Met-X dipeptidase

**Comments:** A Mn<sup>2+</sup>-activated *Escherichia coli* enzyme with thiol dependence

**References:** [309]

## [EC 3.4.13.12 created 1976]

[3.4.13.13 Transferred entry. homocarnosinase. Now EC 3.4.13.3, X-His dipeptidase]

[EC 3.4.13.13 created 1981, deleted 1992]

[3.4.13.14 Deleted entry.  $\gamma$ -glutamyldipeptidase]

[EC 3.4.13.14 created 1989, deleted 1992]

[3.4.13.15] Transferred entry,  $N^2$ - $\beta$ -alanylarginine dipeptidase, Now EC 3.4.13.18, cytosol nonspecific dipeptidase]

[EC 3.4.13.15 created 1989, deleted 1992]

[3.4.13.16 Deleted entry. aspartylphenylalanine dipeptidase]

[EC 3.4.13.16 created 1989, deleted 1992]

#### EC 3.4.13.17

**Accepted name:** non-stereospecific dipeptidase

**Reaction:** Hydrolysis of dipeptides containing either D- or L-amino acids or both **Other name(s):** peptidyl-D-amino acid hydrolase; D-(or L-)aminoacyl-dipeptidase

**Comments:** A digestive enzyme of cephalopods

**References:** [535]

[EC 3.4.13.17 created 1990]

## EC 3.4.13.18

Accepted name: cytosol nonspecific dipeptidase

**Reaction:** Hydrolysis of dipeptides, preferentially hydrophobic dipeptides including prolyl amino acids

Other name(s):  $N^2$ - $\beta$ -alanylarginine dipeptidase; glycyl-glycine dipeptidase; glycyl-leucine dipeptidase; iminodipep-

tidase; peptidase A; Pro-X dipeptidase; prolinase; prolyl dipeptidase; prolylglycine dipeptidase; iminodipeptidase; prolinase; L-prolylglycine dipeptidase; prolylglycine dipeptidase; Gly-Leu hydrolase; glycyl-L-leucine dipeptidase; glycyl-L-leucine hydrolase; glycyl-L-leucine peptidase; L-amino-acyl-L-amino-acid hydrolase; glycylleucine peptidase; glycylleucine hydrolase; glycylleucine dipeptidase; non-specific dipeptidase; human cytosolic non-specific dipeptidase; glycyl-L-

leucine hydrolase; glycyl-glycine dipeptidase

Comments: A zinc enzyme with broad specificity, varying somewhat with source species. Activated and stabilized

by dithiothreitol and Mn<sup>2+</sup>. Inhibited by bestatin and leucine.

**References:** [173]

[EC 3.4.13.18 created 1961 as EC 3.4.3.1 and EC 3.4.3.2, transferred 1972 to EC 3.4.13.1 and EC 3.4.13.2, transferred 1978 to EC 3.4.13.11, part transferred 1992 to EC 3.4.13.18, modified 2000 (EC 3.4.13.15 created 1989, incorporated 1992)]

## EC 3.4.13.19

**Accepted name:** membrane dipeptidase **Reaction:** Hydrolysis of dipeptides

Other name(s): renal dipeptidase; dehydropeptidase I (DPH I); dipeptidase (ambiguous); aminodipeptidase; dipep-

tide hydrolase (ambiguous); dipeptidyl hydrolase (ambiguous); nonspecific dipeptidase; glycosyl-

phosphatidylinositol-anchored renal dipeptidase; MDP

Comments: A membrane-bound, zinc enzyme with broad specificity. Abundant in the kidney cortex. Inhibited by

bestatin and cilastatin. Type example of peptidase family M19.

**References:** [359, 360, 1539, 1173]

[EC 3.4.13.19 created 1961 as EC 3.4.3.1 and EC 3.4.3.2, transferred 1972 to EC 3.4.13.1 and EC 3.4.13.2, transferred 1978 to EC 3.4.13.11, part transferred 1992 to EC 3.4.13.19, modified 2011]

## EC 3.4.13.20

**Accepted name:** β-Ala-His dipeptidase

**Reaction:** Preferential hydrolysis of the  $\beta$ -Ala—His dipeptide (carnosine), and also anserine, Xaa—His dipep-

tides and other dipeptides including homocarnosine

Other name(s): serum carnosinase

**Comments:** Present in the serum of humans and higher primates, but not in the serum of other mammals. Acti-

vated by Cd<sup>2+</sup> and citrate. Belongs in peptidase family M20.

**References:** [1646, 1290]

[EC 3.4.13.20 created 1992]

#### EC 3.4.13.21

Accepted name: dipeptidase E

Reaction: Dipeptidase E catalyses the hydrolysis of dipeptides Asp—Xaa. It does not act on peptides with N-

terminal Glu, Asn or Gln, nor does it cleave isoaspartyl peptides

Other name(s): aspartyl dipeptidase; peptidase E; PepE gene product (Salmonella typhimurium)

Comments: A free carboxy group is not absolutely required in the substrate since Asp-Phe-NH<sub>2</sub> and Asp-Phe-

OMe are hydrolysed somewhat more slowly than dipeptides with free C-termini. No peptide larger than a C-blocked dipeptide is known to be a substrate. Asp-NH-Np is hydrolysed and is a convenient substrate for routine assay. The enzyme is most active near pH 7.0, and is not inhibited by disopropylfluorophosphate or phenylmethanesulfonyl fluoride. Belongs in peptidase family S51.

**References:** [1012, 1609]

[EC 3.4.13.21 created 2001]

# EC 3.4.13.22

**Accepted name:** D-Ala-D-Ala dipeptidase **Reaction:** D-Ala-D-Ala +  $H_2O = 2$  D-Ala

**Other name(s):** D-alanyl-D-alanine dipeptidase; *vanX* D-Ala-D-Ala dipeptidase; VanX

**Comments:** A  $Zn^{2+}$ -dependent enzyme [335]. The enzyme protects *Enterococcus faecium* from the antibiotic

vancomycin, which can bind to the -D-Ala-D-Ala sequence at the C-terminus of the peptidoglycan pentapeptide (see diagram). This enzyme reduces the availability of the free dipeptide D-Ala-D-Ala, which is the precursor for this pentapeptide sequence, allowing D-Ala-(*R*)-lactate (for which vancomycin has much less affinity) to be added to the cell wall instead [3224, 1861]. The enzyme is stereospecific, as L-Ala-L-Ala, D-Ala-L-Ala and L-Ala-D-Ala are not substrates [3224]. Belongs in

peptidase family M15.

**References:** [2413, 3224, 1861, 335, 2869, 1847]

[EC 3.4.13.22 created 2006]

# EC 3.4.14 Dipeptidyl-peptidases and tripeptidyl-peptidases

## EC 3.4.14.1

**Accepted name:** dipeptidyl-peptidase I

**Reaction:** Release of an N-terminal dipeptide, Xaa-Yaa—Zaa-, except when Xaa is Arg or Lys, or Yaa or Zaa is

Pro

Other name(s): cathepsin C; dipeptidyl aminopeptidase I; dipeptidyl transferase; cathepsin C; dipeptidyl transferase;

dipeptide arylamidase I; DAP I

**Comments:** A Cl<sup>-</sup>-dependent, lysosomal cysteine-type peptidase maximally active at acidic pH. Also polymerizes

dipeptide amides, arylamides and esters at neutral pH. In peptidase family C1 (papain family).

**References:** [2291, 1891, 1870, 1869]

[EC 3.4.14.1 created 1961 as EC 3.4.4.9, transferred 1972 to EC 3.4.14.1]

## EC 3.4.14.2

Accepted name: dipeptidyl-peptidase II

Reaction: Release of an N-terminal dipeptide, Xaa-Yaa—, preferentially when Yaa is Ala or Pro. Substrates are

oligopeptides, preferentially tripeptides

Other name(s): dipeptidyl aminopeptidase II; dipeptidyl arylamidase II; carboxytripeptidase; dipeptidyl peptidase II;

dipeptidyl arylamidase II; DAP II; dipeptidyl(amino)peptidase II; dipeptidylarylamidase

Comments: A lysosomal serine-type peptidase in family S28 (Pro-X carboxypeptidase family); maximally active

at acidic pH

**References:** [1868, 1869]

[EC 3.4.14.2 created 1978]

[3.4.14.3 Transferred entry. acylamino-acid-releasing enzyme. Now EC 3.4.19.1, acylaminoacyl-peptidase]

[EC 3.4.14.3 created 1978, deleted 1981]

## EC 3.4.14.4

Accepted name: dipeptidyl-peptidase III

**Reaction:** Release of an N-terminal dipeptide from a peptide comprising four or more residues, with broad

specificity. Also acts on dipeptidyl 2-naphthylamides.

**Other name(s):** dipeptidyl aminopeptidase III; dipeptidyl arylamidase III; enkephalinase B; red cell angiotensinase

**Comments:** A cytosolic peptidase that is active at neutral pH. It has broad activity on peptides, although it is

highly selective for Arg-Arg-2-naphthylamide, at pH 9.2. Active in the hydrolysis of enkephalins.

A metallopeptidase, the type example of peptidase family M49.

**References:** [1866, 854]

[EC 3.4.14.4 created 1981, modified 2001]

#### EC 3.4.14.5

**Accepted name:** dipeptidyl-peptidase IV

Reaction: Release of an N-terminal dipeptide, Xaa-Yaa—Zaa-, from a polypeptide, preferentially when Yaa is

Pro, provided Zaa is neither Pro nor hydroxyproline

Other name(s): dipeptidyl aminopeptidase IV; Xaa-Pro-dipeptidyl-aminopeptidase; Gly-Pro naphthylamidase; post-

proline dipeptidyl aminopeptidase IV; lymphocyte antigen CD26; glycoprotein GP110; dipeptidyl peptidase IV; glycylproline aminopeptidase; glycylproline aminopeptidase; X-prolyl dipeptidyl aminopeptidase; pep X; leukocyte antigen CD26; glycylprolyl dipeptidylaminopeptidase; dipeptidyl-peptide hydrolase; glycylprolyl aminopeptidase; dipeptidyl-aminopeptidase IV; DPP IV/CD26; amino

acyl-prolyl dipeptidyl aminopeptidase; T cell triggering molecule Tp103; X-PDAP

**Comments:** A homodimer. An integral protein of the plasma membrane of lymphocytes and other mammalian

cells, in peptidase family S9 (prolyl oligopeptidase family). The reaction is similar to that of the unre-

lated EC 3.4.14.11 Xaa-Pro dipeptidyl-peptidase of lactococci

**References:** [1930, 543, 1230]

[EC 3.4.14.5 created 1981, modified 1996]

## EC 3.4.14.6

Accepted name: dipeptidyl-dipeptidase

**Reaction:** Preferential release of dipeptides from a tetrapeptide, e.g. Ala-Gly—Ala-Gly. Acts more slowly on

Ala-Ala Ala-Ala and Gly-Gly—Gly-Gly

Other name(s): dipeptidyl tetrapeptide hydrolase; dipeptidyl ligase; tetrapeptide dipeptidase

Comments: A thiol-activated peptidase from cabbage (Brassica oleracea). Tetrapeptides are formed from Ala-

Ala, Gly-Gly, Ala-Gly and Gly-Ala

**References:** [705]

#### [EC 3.4.14.6 created 1989]

[3.4.14.7 Deleted entry. tetralysine endopeptidase]

[EC 3.4.14.7 created 1989, deleted 1992]

[3.4.14.8 Transferred entry. tripeptidyl peptidase. Now EC 3.4.14.10, tripeptidyl-peptidase II]

[EC 3.4.14.8 created 1989, deleted 1992]

EC 3.4.14.9

Accepted name: tripeptidyl-peptidase I

**Reaction:** Release of an N-terminal tripeptide from a polypeptide, but also has endopeptidase activity.

**Other name(s):** tripeptidyl aminopeptidase; tripeptidyl peptidase

Comments: A lysosomal enzyme that is active at acidic pH. Deficient in classical late-infantile neuronal ceroid

lipofuscinosis brain tissue. Belongs in peptidase family S53. Formerly included in EC 3.4.14.8.

**References:** [730, 2384, 729, 1347, 1692]

[EC 3.4.14.9 created 1992 (part of EC 3.4.14.8 created 1989, incorporated 1992), modified 2000, modified 2001, modified 2003]

## EC 3.4.14.10

Accepted name: tripeptidyl-peptidase II

**Reaction:** Release of an N-terminal tripeptide from a polypeptide

Other name(s): tripeptidyl aminopeptidase; tripeptidyl peptidase; tripeptidyl aminopeptidase II; tripeptidyl peptidase

II; TPP

Comments: A cytosolic enzyme in peptidase family S8 (subtilisin family). Active at neutral pH. Inhibited by di-

isopropyl fluorophosphate. Formerly included in EC 3.4.14.8

**References:** [131, 132, 2937]

[EC 3.4.14.10 created 1992 (part of EC 3.4.14.8 created 1989, incorporated 1992)]

#### EC 3.4.14.11

Accepted name: Xaa-Pro dipeptidyl-peptidase

Reaction: Hydrolyses Xaa-Pro bonds to release unblocked, N-terminal dipeptides from substrates including

Ala-Pro p-nitroanilide and (sequentially) Tyr-Pro Phe-Pro Gly-Pro Ile

**Other name(s):** X-prolyl dipeptidyl aminopeptidase; PepX; X-prolyl dipeptidyl peptidase; X-Pro dipeptidyl-peptidase

Comments: The intracellular enzyme from Lactococcus lactis (190-kDa) is the type example of peptidase family

S15. The reaction is similar to that catalysed by dipeptidyl-peptidase IV of animals

**References:** [3322, 1897, 1015, 430, 429]

[EC 3.4.14.11 created 1996]

#### EC 3.4.14.12

**Accepted name:** Xaa-Xaa-Pro tripeptidyl-peptidase

Reaction: Hydrolysis of Xaa-Xaa-Pro—Yaa- releasing the N-terminal tripeptide of a peptide with Pro as the

third residue (position P1) and where Yaa is not proline

Other name(s): prolyltripeptidyl amino peptidase; prolyl tripeptidyl peptidase; prolyltripeptidyl aminopeptidase; PTP-

A; TPP

Comments: This cell-surface-associated serine exopeptidase is found in the Gram-negative, anaerobic bacterium

*Porphyromonas gingivalis*, which has been implicated in adult periodontal disease [134]. The enzyme releases the N-terminal tripeptide of peptides, such as interleukin-6. It has an absolute requirement for a proline residue at the P1 position but is completely inactivated by a proline residue at the P1'

position [134]. The size of the peptide does not affect the rate of reaction [134].

**References:** [134, 843]

#### [EC 3.4.14.12 created 2006]

EC 3.4.14.13

**Accepted name:** γ-D-glutamyl-L-lysine dipeptidyl-peptidase

**Reaction:** The enzyme releases L-Ala-γ-D-Glu dipeptides from cell wall peptides via cleavage of an L-Ala-γ-D-

Glu—L-Lys bond.

Other name(s): YkfC

Comments: The enzyme, characterized from the bacterium *Bacillus subtilis*, is involved in the recycling of the

murein peptide.

**References:** [2571, 3234]

[EC 3.4.14.13 created 2015]

# EC 3.4.15 Peptidyl-dipeptidases

EC 3.4.15.1

**Accepted name:** peptidyl-dipeptidase A

**Reaction:** Release of a C-terminal dipeptide, oligopeptide—Xaa-Yaa, when Xaa is not Pro, and Yaa is neither

Asp nor Glu. Thus, conversion of angiotensin I to angiotensin II, with increase in vasoconstrictor ac-

tivity, but no action on angiotensin II

Other name(s): dipeptidyl carboxypeptidase I; peptidase P; dipeptide hydrolase (ambiguous); peptidyl dipeptidase;

angiotensin converting enzyme; kininase II; angiotensin I-converting enzyme; carboxycathepsin; dipeptidyl carboxypeptidase; peptidyl dipeptidase I; peptidyl-dipeptide hydrolase; peptidyl dipeptidase; ACE; peptidyl dipeptidase-4; PDH; peptidyl dipeptidyl dipeptidase-4; PDH; peptidyl dipeptidyl dip

tide hydrolase; DCP

**Comments:** A Cl<sup>-</sup>-dependent, zinc glycoprotein that is generally membrane-bound. A potent inhibitor is cap-

topril. Important in elevation of blood pressure, through formation of angiotensin II (vasoconstrictor) and destruction of bradykinin (vasodilator). Two molecular forms exist in mammalian tissues, a widely-distributed somatic form of 150- to 180-kDa that contains two non-identical catalytic sites, and a testicular form of 90- to 100-kDa that contains only a single catalytic site. Type example of pep-

tidase family M2

**References:** [2730, 682, 3147, 490]

[EC 3.4.15.1 created 1972, modified 1981, modified 1989, modified 1996, modified 2011]

[3.4.15.2 Transferred entry. pepdidyl carboxyamidase. Now EC 3.4.19.2, peptidyl-glycinamidase]

[EC 3.4.15.2 created 1978, deleted 1981]

[3.4.15.3] Transferred entry, dipeptidyl carboxypeptidase, Now EC 3.4.15.5, peptidyl-dipeptidase Dcp]

[EC 3.4.15.3 created 1981, modified 1989, deleted 1996]

EC 3.4.15.4

Accepted name: peptidyl-dipeptidase B

**Reaction:** Release of a C-terminal dipeptide or exceptionally a tripeptide

Other name(s): dipeptidyl carboxyhydrolase; atriopeptin convertase; atrial di-(tri)peptidyl carboxyhydrolase; pep-

tidyldipeptidase B; atrial dipeptidyl carboxyhydrolase; atrial peptide convertase

Comments: A membrane-bound, zinc metallopeptidase located in mammalian atrial, but not ventricular, my-

ocytes. Although it is capable of converting the 126-residue atriopeptin III directly to atriopeptin I by releasing a C-terminal tripeptide Phe-Arg-Tyr, it is generally restricted to the release of dipeptides. In contrast to peptidyl-dipeptidase A (EC 3.4.15.1) it displays no Cl<sup>-</sup> dependence and shows no action on angiotensin I. Conversely, peptidyl-dipeptidase A is unable to release Phe-Arg from the C-terminus

of atriopeptin II

**References:** [1061, 1062, 2711, 2712]

## [EC 3.4.15.4 created 1992]

EC 3.4.15.5

**Accepted name:** peptidyl-dipeptidase Dcp

Reaction: Hydrolysis of unblocked, C-terminal dipeptides from oligopeptides, with broad specificity. Does not

hydrolyse bonds in which P1' is Pro, or both P1 and P1' are Gly

**Other name(s):** dipeptidyl carboxypeptidase (Dcp); dipeptidyl carboxypeptidase

Comments: Known from Escherichia coli and Salmonella typhimurium. A zinc metallopeptidase in peptidase

family M3 (thimet oligopeptidase family). Ac-Ala—Ala is a good test substrate [474]. Inhibited by captopril, as is peptidyl-dipeptidase A. Formerly EC 3.4.15.3, and included in EC 3.4.15.1,

peptidyl-dipeptidase A.

**References:** [3267, 1114, 474]

[EC 3.4.15.5 created 1981 as EC 3.4.15.3, modified 1989, transferred 1996 to EC 3.4.15.5]

EC 3.4.15.6

Accepted name: cyanophycinase

**Reaction:**  $[L-Asp(4-L-Arg)]_n + H_2O = [L-Asp(4-L-Arg)]_{n-1} + L-Asp(4-L-Arg)$ 

Other name(s): cyanophycin degrading enzyme; β-Asp-Arg hydrolysing enzyme; CGPase; CphB; CphE;

cyanophycin granule polypeptidase; extracellular CGPase

**Comments:** The enzyme is highly specific for the branched polypeptide cyanophycin and does not hydrolyse poly-

L-aspartate or poly-L-arginine [2423]. A serine-type exopeptidase that belongs in peptidase family

S51.

**References:** [2125, 2126, 2423]

[EC 3.4.15.6 created 2007]

# EC 3.4.16 Serine-type carboxypeptidases

[3.4.16.1] Transferred entry, serine carboxypeptidase, Now EC 3.4.16.6, carboxypeptidase D]

[EC 3.4.16.1 created 1972 as EC 3.4.12.1 and EC 3.4.21.13, both transferred 1978 to EC 3.4.16.1, deleted 1993]

EC 3.4.16.2

Accepted name: lysosomal Pro-Xaa carboxypeptidase

**Reaction:** Cleavage of a -Pro—Xaa bond to release a C-terminal amino acid

**Other name(s):** angiotensinase C; lysosomal carboxypeptidase C; peptidylprolylamino acid carboxypeptidase;

aminoacylproline carboxypeptidase; prolyl carboxypeptidase; carboxypeptidase P; proline-specific

carboxypeptidase P; PCP; lysosomal Pro-Xaa carboxypeptidase

**Comments:** A lysosomal peptidase active at acidic pH that inactivates angiotensin II. Inhibited by diisopropyl flu-

orophosphate. In peptidase family S28 (Pro-X carboxypeptidase family).

**References:** [3102, 2139]

[EC 3.4.16.2 created 1972 as EC 3.4.12.4, transferred 1978 to EC 3.4.16.2]

[3.4.16.3 Transferred entry. tyrosine carboxypeptidase. Now included with EC 3.4.16.5, carboxypeptidase C]

[EC 3.4.16.3 created 1972 as EC 3.4.12.12, transferred 1978 to EC 3.4.16.3, deleted 1992]

EC 3.4.16.4

Accepted name: serine-type D-Ala-D-Ala carboxypeptidase

**Reaction:** Preferential cleavage: (Ac)<sub>2</sub>-L-Lys-D-Ala. Also transpeptidation of peptidyl-alanyl moieties

that are N-acyl substituents of D-alanine

Other name(s): DD-peptidase; D-alanyl-D-alanine-carboxypeptidase; D-alanyl-D-alanine-cleaving-peptidase;

D-alanyl-D-alanine-cleaving peptidase; DD-transpeptidase; D-alanine carboxypeptidase; DD-

carboxypeptidase; D-alanyl carboxypeptidase

**Comments:** A membrane-bound, bacterial enzyme inhibited by penicillin and other β-lactam antibiotics, which

acylate the active site serine. Examples are known from peptidase families S11, S12 and S13. Distinct

from EC 3.4.17.14, zinc D-Ala-D-Ala carboxypeptidase

**References:** [909, 810]

[EC 3.4.16.4 created 1989]

#### EC 3.4.16.5

Accepted name: carboxypeptidase C

**Reaction:** Release of a C-terminal amino acid with broad specificity

Other name(s): carboxypeptidase Y; serine carboxypeptidase I; cathepsin A; lysosomal protective protein; deamidase;

lysosomal carboxypeptidase A; phaseolin

**Comments:** A carboxypeptidase with optimum pH 4.5-6.0, inhibited by diisopropyl fluorophosphate, and sensitive

to thiol-blocking reagents (reviewed in [283]). Widely distributed in eukaryotes. Type example of

peptidase family S10.

**References:** [283, 3030, 1287, 1914]

[EC 3.4.16.5 created 1972 as EC 3.4.12.1, transferred 1978 to EC 3.4.16.1, part transferred 1993 to EC 3.4.16.5 (EC 3.4.16.3 created 1972 as EC 3.4.12.12, transferred 1978 to EC 3.4.16.3, transferred 1992 to EC 3.4.16.1), (EC 3.4.21.13 created 1972, transferred 1978 to EC 3.4.16.1)]

## EC 3.4.16.6

Accepted name: carboxypeptidase D

**Reaction:** Preferential release of a C-terminal arginine or lysine residue

Other name(s): cereal serine carboxypeptidase II; Saccharomyces cerevisiae KEX1 gene product; carboxypep-

tidase Kex1; gene KEX1 serine carboxypeptidase; KEX1 carboxypeptidase; KEX1 proteinase; KEX1DELTAp; CPDW-II; serine carboxypeptidase (misleading); *Phaseolus* proteinase

**Comments:** A carboxypeptidase with optimum pH 4.5-6.0, inhibited by diisopropyl fluorophosphate, and sensitive

to thiol-blocking reagents (reviewed in [283]). In peptidase family S10 (carboxypeptidase C family).

**References:** [283, 285, 621, 1678]

[EC 3.4.16.6 created 1972 as EC 3.4.12.1, transferred 1978 to EC 3.4.16.1, part transferred 1993 to EC 3.4.16.6 (EC 3.4.16.3 created 1972 as EC 3.4.12.12, transferred 1978 to EC 3.4.16.3, transferred 1992 to EC 3.4.16.1), (EC 3.4.21.13 created 1972, transferred 1978 to EC 3.4.16.1), modified 2011]

# EC 3.4.17 Metallocarboxypeptidases

#### EC 3.4.17.1

Accepted name: carboxypeptidase A

**Reaction:** Release of a C-terminal amino acid, but little or no action with -Asp, -Glu, -Arg, -Lys or -Pro

Other name(s): carboxypolypeptidase; pancreatic carboxypeptidase A; tissue carboxypeptidase A

**Comments:** A zinc enzyme formed from procarboxypeptidase A. Isolated from cattle, pig and dogfish pancreas,

and other sources including mast cells [726] and skeletal muscle [252]. Type example of peptidase

family M14.

**References:** [2267, 2394, 726, 252]

 $[EC\ 3.4.17.1\ created\ 1961\ as\ EC\ 3.4.2.1,\ transferred\ 1972\ to\ EC\ 3.4.12.2,\ transferred\ 1978\ to\ EC\ 3.4.17.1]$ 

#### EC 3.4.17.2

Accepted name: carboxypeptidase B

**Reaction:** Preferential release of a C-terminal lysine or arginine amino acid

Other name(s): protaminase; pancreatic carboxypeptidase B; tissue carboxypeptidase B; peptidyl-L-lysine [L-

arginine]hydrolase

**Comments:** A zinc enzyme formed from procarboxypeptidase B. Isolated from cattle, pig and dogfish pancreas

and other sources, including skin fibroblasts [338] and adrenal medulla [3098]. In peptidase family

M14 (carboxypeptidase A family).

**References:** [788, 299, 338, 3098]

[EC 3.4.17.2 created 1961 as EC 3.4.2.2, transferred 1972 to EC 3.4.12.3, transferred 1978 to EC 3.4.17.2]

#### EC 3.4.17.3

Accepted name: lysine carboxypeptidase

**Reaction:** Release of a C-terminal basic amino acid, preferentially lysine

Other name(s): carboxypeptidase N; arginine carboxypeptidase; kininase I; anaphylatoxin inactivator; plasma car-

boxypeptidase B; creatine kinase conversion factor; bradykinase; kininase Ia; hippuryllysine hydrolase; bradykinin-decomposing enzyme; protaminase; CPase N; creatinine kinase convertase; peptidyl-

L-lysine(-L-arginine) hydrolase; CPN

Comments: A zinc enzyme found in plasma. Inactivates bradykinin and anaphylatoxins in blood plasma. In pepti-

dase family M14 (carboxypeptidase A family).

**References:** [2294, 1656, 2681]

[EC 3.4.17.3 created 1972 as EC 3.4.12.7, transferred 1978 to EC 3.4.17.3, modified 1989]

#### EC 3.4.17.4

Accepted name: Gly-Xaa carboxypeptidase

**Reaction:** Release of a C-terminal amino acid from a peptide in which glycine is the penultimate amino acid,

e.g. Z-Gly—Leu

Other name(s): glycine carboxypeptidase; carboxypeptidase a; carboxypeptidase S; peptidase  $\alpha$ ; yeast carboxypepti-

dase; Gly-X carboxypeptidase

**Comments:** From yeast. In peptidase family M20 (glutamate carboxypeptidase family).

**References:** [751, 3206]

[EC 3.4.17.4 created 1961 as EC 3.4.2.3, transferred 1972 to EC 3.4.12.8, transferred 1978 to EC 3.4.17.4 (EC 3.4.17.9 created 1981, incorporated 1992)]

[3.4.17.5 Deleted entry. aspartate carboxypeptidase]

[EC 3.4.17.5 created 1972 as EC 3.4.12.9, transferred 1978 to EC 3.4.17.5, deleted 1992]

## EC 3.4.17.6

**Accepted name:** alanine carboxypeptidase

**Reaction:** Release of a C-terminal alanine from a peptide or a variety of pteroyl or acyl groups

**Other name(s):** *N*-benzoyl-L-alanine-amidohydrolase

**Comments:** From soil bacteria. The enzyme from *Corynebacterium* equi also hydrolyses *N*-benzoylglycine and

N-benzoyl-L-aminobutyric acid.

**References:** [1659, 1936]

[EC 3.4.17.6 created 1972 as EC 3.4.12.11, transferred 1978 to EC 3.4.17.6]

[3.4.17.7 Transferred entry. acylmuramoyl-alanine carboxypeptidase. Now EC 3.5.1.28, N-acetylmuramoyl-L-alanine amidase]

[EC 3.4.17.7 created 1978, deleted 1992]

**Accepted name:** muramoylpentapeptide carboxypeptidase

**Reaction:** Cleavage of the bond UDP-*N*-acetylmuramoyl-L-alanyl-γ-D-glutamyl-6-carboxy-L-lysyl-D-

alanyl-D-alanine

Other name(s): D-alanine carboxypeptidase I; DD-carboxypeptidase; D-alanine carboxypeptidase; D-alanine

carboxypeptidase; D-alanine-D-alanine-carboxypeptidase; carboxypeptidase D-alanyl-D-alanine; carboxypeptidase I; UDP-*N*-acetylmuramoyl-tetrapeptidyl-D-alanine alanine-hydrolase; D-alanyl-D-alanyl-D-alanine

alanine peptidase; DD-peptidase; penicillin binding protein 5; PBP5; PdcA; VanY

Comments: A bacterial enzyme that requires a divalent cation for activity. Does not cleave the C-terminal D-

alanine from the product of the above reaction, UDP-N-acetyl-muramoyl-L-alanyl-γ-D-glutamyl-6-

carboxy-L-lysyl-D-alanine. Competitively inhibited by penicillins and cephalosporins.

**References:** [1285]

[EC 3.4.17.8 created 1972 as EC 3.4.12.6, transferred 1978 to EC 3.4.17.8]

[3.4.17.9 Transferred entry, carboxypeptidase S. Now included with EC 3.4.17.4, Gly-Xaa carboxypeptidase]

[EC 3.4.17.9 created 1981, deleted 1992]

## EC 3.4.17.10

Accepted name: carboxypeptidase E

**Reaction:** Release of C-terminal arginine or lysine residues from polypeptides

Other name(s): carboxypeptidase H; enkephalin convertase; cobalt-stimulated chromaffin granule carboxypeptidase;

insulin granule-associated carboxypeptidase; enkephalin convertase; membrane-bound carboxypeptidase; carboxypeptidase E; enkephalin-precursor endopeptidase; enkephalin precursor carboxypepti-

dase; peptidyl-L-lysine(-L-arginine) hydrolase

**Comments:** A zinc enzyme, activated by Co<sup>2+</sup>. Inhibited by 1,10-phenanthroline and other chelating agents. pH

optimum 5.6. Located in storage granules of secretory cells, and active in processing of protein hor-

mones and bioactive peptides. In peptidase family M14 (carboxypeptidase A family).

**References:** [2337, 818, 817, 1803, 816]

[EC 3.4.17.10 created 1986, modified 2000]

# EC 3.4.17.11

**Accepted name:** glutamate carboxypeptidase

**Reaction:** Release of C-terminal glutamate residues from a wide range of N-acylating moieties, including pep-

tidyl, aminoacyl, benzoyl, benzyloxycarbonyl, folyl and pteroyl groups

**Other name(s):** carboxypeptidase  $G_1$ ; carboxypeptidase  $G_2$ ; glutamyl carboxypeptidase;  $N_2$ 

pteroyl-L-glutamate hydrolase

**Comments:** A zinc enzyme produced by pseudomonads, *Flavobacterium* sp. and *Acinetobacter* sp. Its ability to

hydrolyse pteroyl-L-glutamate (folic acid) has led to its use as a folate-depleting, antitumour agent.

Type example of peptidase family M20

**References:** [947, 1865, 29, 2627]

[EC 3.4.17.11 created 1992]

## EC 3.4.17.12

Accepted name: carboxypeptidase M

**Reaction:** Cleavage of C-terminal arginine or lysine residues from polypeptides

Other name(s): CPM

Comments: A membrane-bound enzyme optimally active at neutral pH. In peptidase family M14 (carboxypepti-

dase A family)

**References:** [2682, 566, 2683]

[EC 3.4.17.12 created 1992]

## EC 3.4.17.13

**Accepted name:** muramoyltetrapeptide carboxypeptidase

**Reaction:** Hydrolysis of the bond: *N*-acetyl-D-glucosaminyl-*N*-acetylmuramoyl-L-Ala-D-glutamyl-6-carboxy-L-

lysyl\\_D-alanine

Other name(s): carboxypeptidase IIW; carboxypeptidase II; lysyl-D-alanine carboxypeptidase; L-lysyl-D-alanine car-

boxypeptidase; LD-carboxypeptidase

**Comments:** Variants are known from various microorganisms. Involved in peptidoglycan synthesis, catalysing

both decarboxylation and transpeptidation. Stimulated by divalent cations such as Mg<sup>2+</sup> and Ca<sup>2+</sup>,

but not by Zn<sup>2+</sup>. Inhibited by thiol-blocking reagents, but unaffected by penicillin

**References:** [539, 2467, 1893]

[EC 3.4.17.13 created 1992]

# EC 3.4.17.14

**Accepted name:** zinc D-Ala-D-Ala carboxypeptidase

**Reaction:** Cleavage of the bond: (Ac)<sub>2</sub>-L-lysyl-D-alanyl—D-alanine

**Other name(s):** Zn<sup>2+</sup> G peptidase, D-alanyl-D-alanine hydrolase; D-alanyl-D-alanine-cleaving carboxypeptidase; DD-

carboxypeptidase; G enzyme; DD-carboxypeptidase-transpeptidase

**Comments:** A zinc enzyme. Catalyses carboxypeptidation but not transpeptidation reactions involved in bacterial

cell wall metabolism. Weakly inhibited by  $\beta$ -lactams. In peptidase family M15. Distinct from EC

3.4.16.4, serine-type D-Ala-D-Ala carboxypeptidase.

**References:** [606, 1340, 909]

[EC 3.4.17.14 created 1992]

## EC 3.4.17.15

**Accepted name:** carboxypeptidase A<sub>2</sub>.

**Reaction:** Similar to that of carboxypeptidase A (EC 3.4.17.1), but with a preference for bulkier C-terminal

residues

Other name(s): CPA2

**Comments:** Isolated from rat pancreas but not present in cattle pancreas. In peptidase family M14 (carboxypepti-

dase A family).

**References:** [878]

[EC 3.4.17.15 created 1992]

# EC 3.4.17.16

Accepted name: membrane Pro-Xaa carboxypeptidase

**Reaction:** Release of a C-terminal residue other than proline, by preferential cleavage of a prolyl bond **Other name(s):** carboxypeptidase P; microsomal carboxypeptidase; membrane Pro-X carboxypeptidase

**Comments:** One of the renal brush border exopeptidases

**References:** [569, 263, 1096]

[EC 3.4.17.16 created 1992]

# EC 3.4.17.17

**Accepted name:** tubulinyl-Tyr carboxypeptidase

**Reaction:** Cleavage of the -Glu—Tyr bond to release the C-terminal tyrosine residue from the native tyrosinated

tubulin. Inactive on Z-Glu-Tyr

Other name(s): carboxypeptidase-tubulin; soluble carboxypeptidase; tubulin-tyrosine carboxypeptidase; tubulin car-

boxypeptidase; tubulinyltyrosine carboxypeptidase; tyrosinotubulin carboxypeptidase; tyrosyltubulin

carboxypeptidase; TTCPase; brain I carboxypeptidase

**Comments:** Active at neutral pH, from brain

**References:** [2019, 1549, 70]

## [EC 3.4.17.17 created 1992]

EC 3.4.17.18

**Accepted name:** carboxypeptidase T

**Reaction:** Releases a C-terminal residue, which may be hydrophobic or positively charged

Other name(s): CPT

**Comments:** Known from *Thermoactinomyces vulgaris*. In peptidase family M14 (carboxypeptidase A family)

**References:** [2208, 2703, 2898]

[EC 3.4.17.18 created 1993]

EC 3.4.17.19

Accepted name: carboxypeptidase Taq

**Reaction:** Release of a C-terminal amino acid with broad specificity, except for -Pro

**Comments:** A 56-kDa enzyme from *Thermus aquaticus*. Most active at 80° C. Type example of peptidase family

M32

**References:** [1627, 1628]

[EC 3.4.17.19 created 1996]

EC 3.4.17.20

**Accepted name:** carboxypeptidase U

**Reaction:** Release of C-terminal Arg and Lys from a polypeptide

Other name(s): arginine carboxypeptidase; carboxypeptidase R; plasma carboxypeptidase B (misleading, since the

term carboxypeptidase B is used for other enzymes); thrombin-activatable fibrinolysis inhibitor

Comments: Pro-carboxypeptidase U in (human) plasma is activated by thrombin or plasmin during clotting to

form the unstable carboxypeptidase U, with activity similar to that of the more stable lysine carboxypeptidase, except that no preference is shown for Lys over Arg. A zinc enzyme, in peptidase

family M14 (carboxypeptidase A family)

**References:** [674, 2640, 3121, 2868, 312]

[EC 3.4.17.20 created 1997]

EC 3.4.17.21

Accepted name: glutamate carboxypeptidase II

**Reaction:** Release of an unsubstituted, C-terminal glutamyl residue, typically from Ac-Asp-Glu or folylpoly- $\gamma$ -

glutamates

Other name(s): N-acetylated-γ-linked-acidic dipeptidase (NAALADase); folate hydrolase; prostate-specific mem-

brane antigen; pteroylpoly- $\gamma$ -glutamate carboxypeptidase; microsomal  $\gamma$ -glutamyl carboxypeptidase; pteroylpolyglutamate hydrolase; pteroylpoly- $\gamma$ -glutamate hydrolase; pteroylpolyglutamate hydrolase; pteroylpolyglutamic acid hydrolase; pteroylpolyglutamate hydrolase; pteroylpolyglutamic acid hydrolase; PSM antigen; acetylaspartylglutamate dipeptidase; NAALA dipeptidase; rat NAAG peptidase; mGCP; membrane glutamate carboxypeptidase; N-acetylated- $\alpha$ -linked-amino dipeptidase; prostrate-

specific membrane antigen; N-Acetylated α-linked acidic dipeptidase; PSMA

Comments: A metallo-carboxypeptidase that is predominantly expressed as a membrane-bound enzyme of 94-

100 kDa , but also exists in a soluble form. Hydrolyses  $\alpha$ -peptide bonds in Ac-Asp-Glu, Asp-Glu, and Glu-Glu, but also  $\gamma$ -glutamyl bonds in  $\gamma$ -Glu-Glu, and folylpoly- $\gamma$ -glutamates. With folylpoly- $\gamma$ -glutamates, shows processive carboxypeptidase activity to produce pteroylmonoglutamate [1758]. Does not hydrolyse Ac- $\beta$ -Asp-Glu. Known inhibitors: quisqualic acid, Ac- $\beta$ -Asp-Glu, and 2-phosphonomethyl-pentanedioate. In peptidase family M28 of *Vibrio* leucyl aminopeptidase. The release of C-terminal glutamate from folylpoly- $\gamma$ -glutamates is also catalysed by EC 3.4.17.11 (gluta-

mate carboxypeptidase) and EC 3.4.19.9 (γ-Glu-X carboxypeptidase).

**References:** [1134, 2383, 1032, 1758]

[EC 3.4.17.21 created 1997, modified 2000 (EC 3.4.13.8 created 1972 and EC 3.4.19.8 created 1992, incorporated 2000)]

EC 3.4.17.22

Accepted name: metallocarboxypeptidase D

**Reaction:** Releases C-terminal Arg and Lys from polypeptides

Other name(s): carboxypeptidase D (cattle, human, mouse, rat); gp180 (duck)

**Comments:** Activated by Co<sup>2+</sup>; inhibited by guanidinoethylmercaptosuccinic acid. Large molecule (180 kDa) because of presence of three copies of metallopeptidase domain. The product of the silver gene

because of presence of three copies of metallopeptidase domain. The product of the silver gene (*Drosophila*) is similar. A zinc metallopeptidase in peptidase family M14 (carboxypeptidase A fam-

ily)

**References:** [1573, 2718, 2719]

[EC 3.4.17.22 created 1997]

EC 3.4.17.23

Accepted name: angiotensin-converting enzyme 2

**Reaction:** angiotensin II +  $H_2O$  = angiotensin-(1-7) + L-phenylalanine

Other name(s): ACE-2; ACE2; hACE2; angiotensin converting enzyme 2; angiotensin converting enzyme-2; Tmem27 Comments: A transmembrane glycoprotein with an extracellular catalytic domain. Angiotensin-converting en-

zyme 2 functions as a carboxypeptidase, cleaving a single C-terminal residue from a distinct range of substrates [1594]. Catalytic efficiency is 400-fold higher with angiotensin II (1–8) as a substrate than with angiotensin I (1–10). Angiotensin-converting enzyme 2 also efficiently hydrolyses des-Arg<sup>9</sup>-

bradykinin, but it does not hydrolyse bradykinin [3066]. In peptidase family M2.

**References:** [3066, 1594, 2954]

[EC 3.4.17.23 created 2009]

# EC 3.4.18 Cysteine-type carboxypeptidases

EC 3.4.18.1

**Accepted name:** cathepsin X

**Reaction:** Release of C-terminal amino acid residues with broad specificity, but lacks action on C-terminal pro-

line. Shows weak endopeptidase activity

Other name(s): cathepsin B2; cysteine-type carboxypeptidase; cathepsin IV; cathepsin Z; acid carboxypeptidase; lyso-

somal carboxypeptidase B

**Comments:** Cathepsin X is a lysosomal cysteine peptidase of family C1 (papain family). The pH optimum is de-

pendent on the substrate and is 5.0 for the carboxypeptidase activity. Unstable above pH 7.0. Compound E-64, leupeptin and antipain are inhibitors, but not cystatin C. Cathepsin X is ubiquitously distributed in mammalian tissues. The propeptide is extremely short (38 amino acid residues) and the

proenzyme is catalytically active. Human gene locus: 20q13.

**References:** [2027, 2026, 2526, 1867, 2216, 2091]

[EC 3.4.18.1 created 1981, modified 2000]

# EC 3.4.19 Omega peptidases

EC 3.4.19.1

Accepted name: acylaminoacyl-peptidase

**Reaction:** Cleavage of an N-acetyl or N-formyl amino acid from the N-terminus of a polypeptide

Other name(s): acylamino-acid-releasing enzyme; N-acylpeptide hydrolase; N-formylmethionine (fMet) aminopepti-

dase; α-N-acylpeptide hydrolase

Comments: Active at neutral pH. Several variants of this enzyme exist; the human erythrocyte enzyme is rela-

tively specific for removal of N-acetylalanine from peptides. Displays dipeptidyl-peptidase activity on glycyl-peptides, perhaps as a result of mis-recognition of the glycyl residue as an uncharged N-acyl group. Inhibited by diisopropyl fluorophosphate. In peptidase family S9 (prolyl oligopeptidase fam-

ily).

**References:** [2986, 3012, 1490]

[EC 3.4.19.1 created 1978 as EC 3.4.14.3, transferred 1981 to EC 3.4.19.1]

EC 3.4.19.2

**Accepted name:** peptidyl-glycinamidase

**Reaction:** Cleavage of C-terminal glycinamide from polypeptides

Other name(s): carboxyamidase; peptidyl carboxy-amidase; peptidyl-aminoacylamidase; carboxamidopeptidase; pep-

tidyl amino acid amide hydrolase

**Comments:** Inactivates vasopressin and oxytocin by splitting off glycinamide. Also cleaves ester substrates of

trypsin and chymotrypsin. Although glycinamide is by far the preferred leaving group, other aminoacylamides may also be released, e.g. phenylalaninamide. The toad skin enzyme is inhibited by diiso-

propyl fluorophosphate.

**References:** [825, 2057, 2666]

[EC 3.4.19.2 created 1978 as EC 3.4.15.2, transferred 1981 to EC 3.4.19.2]

EC 3.4.19.3

Accepted name: pyroglutamyl-peptidase I

**Reaction:** Release of an N-terminal pyroglutamyl group from a polypeptide, the second amino acid generally

not being Pro

Other name(s): 5-oxoprolyl-peptidase; pyroglutamate aminopeptidase; pyroglutamyl aminopeptidase; L-

pyroglutamyl peptide hydrolase; pyrrolidone-carboxyl peptidase; pyrrolidone-carboxylate peptidase; pyrrolidonecarboxylate peptidase; pyrrolidonecarboxylyl

peptidase

Comments: A cysteine peptidase, known from bacteria, plants and animals. The enzyme from bacterial sources is

used in protein sequencing, and is the type example of peptidase family C15.

**References:** [2988, 106, 2247, 2536]

[EC 3.4.19.3 created 1972 as EC 3.4.11.8, transferred 1981 to EC 3.4.19.3, modified 1997]

[3.4.19.4 Deleted entry. N-acetylmethionylpeptide peptidase]

[EC 3.4.19.4 created 1989, deleted 1992]

EC 3.4.19.5

**Accepted name:** β-aspartyl-peptidase

**Reaction:** Cleavage of a β-linked Asp residue from the N-terminus of a polypeptide **Other name(s):** β-aspartyl dipeptidase; β-aspartyl peptidase; β-aspartyldipeptidase

Comments: Other isopeptide bonds, e.g.  $\gamma$ -glutamyl and  $\beta$ -alanyl, are not hydrolysed. A mammalian, cytosolic

enzyme.

**References:** [1025]

[EC 3.4.19.5 created 1972 as EC 3.4.13.10, transferred 1992 to EC 3.4.19.5, modified 1997]

EC 3.4.19.6

Accepted name: pyroglutamyl-peptidase II

**Reaction:** Release of the N-terminal pyroglutamyl group from pGlu—His-Xaa tripeptides and pGlu—His-Xaa-

Gly tetrapeptides

Other name(s): thyroliberinase; pyroglutamyl aminopeptidase II; , thyrotropin-releasing factor pyroglutamate

aminopeptidase; pyroglutamate aminopeptidase II; pyroglutamyl peptidase II; thyroliberin-

hydrolyzing pyroglutamate aminopeptidase; thyrotropin-releasing hormone-degrading pyroglutamate

aminopeptidase; thyrotropin-releasing hormone-degrading peptidase; TRH aminopeptidase

Comments: Highly specific for thyrotropin releasing hormone (pyroglutamyl-histidyl-prolylamide). Will not

cleave the pyroglutamyl-histidyl bond of luteinizing hormone releasing hormone. Found in serum and brain. Inhibited by metal chelators. In peptidase family M1 (membrane alanyl aminopeptidase family)

**References:** [174, 2128, 3187]

[EC 3.4.19.6 created 1992]

## EC 3.4.19.7

**Accepted name:** *N*-formylmethionyl-peptidase

**Reaction:** Release of an N-terminal, formyl-methionyl residue from a polypeptide

Other name(s): (fMet)-releasing enzyme; formylmethionine aminopeptidase

**Comments:** Highly specific for N-formylmethionyl peptides. Will not cleave methionyl peptides or N-formyl

derivatives of amino acids other than methionine. Isolated from rat liver. Inhibited by heavy metals

and activated by Cl-

**References:** [2786]

[EC 3.4.19.7 created 1992]

[3.4.19.8 Transferred entry. now EC 3.4.17.21, glutamate carboxypeptidase II]

[EC 3.4.19.8 created 1992, deleted 2000]

#### EC 3.4.19.9

**Accepted name:** folate γ-glutamyl hydrolase

**Reaction:** tetrahydropteroyl-(γ-glutamyl)<sub>n</sub> + (n-1) H<sub>2</sub>O = 5,6,7,8-tetrahydrofolate + (n-1) L-glutamate **Other name(s):** GGH (gene name); conjugase; folate conjugase; lysosomal γ-glutamyl carboxypeptidase; γ-Glu-X

carboxypeptidase; pteroyl-poly- $\gamma$ -glutamate hydrolase; carboxypeptidase G; folic acid conjugase; poly( $\gamma$ -glutamic acid) endohydrolase; polyglutamate hydrolase; poly(glutamic acid) hydrolase II;

pteroylpoly-γ-glutamyl hydrolase; γ-glutamyl hydrolase

**Systematic name:** tetrahydropteroyl-poly- $\gamma$ -glutamyl  $\gamma$ -glutamyl hydrolase

**Comments:** The enzyme, which occurs only in animals and plants, can be either endo- and/or exopeptidase. It acts

on tetrahydropteroyl polyglutamates and their modified forms, as well as the polyglutamates of the folate breakdown product N-(4-aminobenzoyl)-L-glutamate (pABA-Glu). The initial cleavage may release either monoglutamate or poly- $\gamma$ -glutamate of two or more residues, depending on the specific enzyme. For example, GGH1 from the plant Arabidopsis thaliana cleaves pentaglutamates, mainly to di- and triglutamates, whereas GGH2 from the same organism yields mainly monoglutamates. The

enzyme is lysosomal (and secreted) in animals and vacuolar in plants.

**References:** [1873, 3124, 3264, 3265, 3263, 2204, 25]

[EC 3.4.19.9 created 1972 as EC 3.4.12.10, transferred 1978 to EC 3.4.22.12, transferred 1992 to EC 3.4.19.9, modified 1997, modified 2018]

[3.4.19.10 Transferred entry, acylmuramoyl-Ala peptidase, Now EC 3.5.1.28, N-acetylmuramoyl-L-alanine amidase]

[EC 3.4.19.10 created 1972 as EC 3.4.12.5, transferred 1978 to EC 3.4.17.7, transferred 1992 to EC 3.4.19.10, deleted 1997]

#### EC 3.4.19.11

**Accepted name:** γ-D-glutamyl-*meso*-diaminopimelate peptidase

**Reaction:** Hydrolysis of γ-D-glutamyl bonds to the L-terminus (position 7) of meso-diaminopimelic acid (meso-

A2pm) in 7-(L-Ala-\gamma-D-Glu)-meso-A2pm and 7-(L-Ala-\gamma-Glu)-7-(D-Ala)-meso-A2pm. It is re-

quired that the D-terminal amino and carboxy groups of meso-A2pm are unsubstituted

Other name(s): endopeptidase Ι; γ-D-glutamyldiaminopimelate endopeptidase; γ-D-glutamyl-L-meso-diaminopimelate

peptidoglycan hydrolase;  $\gamma$ -glutamyl-meso-diaminopimelyl endopeptidase;  $\gamma$ -D-glutamyl-meso-diaminopimelate endopeptidase;  $\gamma$ -D-glutamyl-meso-diaminopimelic peptidoglycan hydrolase;  $\gamma$ -D-glutamyl-meso-D-aminopimelic endopeptidase

**Comments:** A 45-kDa metallopeptidase from *Bacillus sphaericus*, the substrates being components of the bacte-

rial spore wall. A member of peptidase family M14 (carboxypeptidase A family). Endopeptidase II has similar activity, but differs in cellular location, molecular mass and catalytic mechanism [1188]

**References:** [74, 880, 1188]

[EC 3.4.19.11 created 1996]

EC 3.4.19.12

Accepted name: ubiquitinyl hydrolase 1

Reaction: Thiol-dependent hydrolysis of ester, thioester, amide, peptide and isopeptide bonds formed by the C-

terminal Gly of ubiquitin (a 76-residue protein attached to proteins as an intracellular targeting signal)

**Other name**(s): ubiquitin C-terminal hydrolase; yeast ubiquitin hydrolase

**Comments:** Links to polypeptides smaller than 60 residues are hydrolysed more readily than those to larger

polypeptides. Isoforms exist with quantitatively different specificities, amongst the best known being UCH-L1 and UCH-L3, which are major proteins of the brain of mammals [1333]. Inhibited by ubiquitin aldehyde (in which Gly<sup>76</sup> is replaced by aminoacetaldehyde). Ubiquitinyl hydrolase 1 is the type example of peptidase family C12, with a similar protein fold to papain and catalytic amino acids Cys, His and Asp. There is a separate family (C19) of enzymes that also hydrolyse ubiquitinyl bonds, and it is thought that all the ubiquitinyl hydrolases are also ubiquitin thiolesterases (EC 3.1.2.15)

**References:** [1333, 3191]

[EC 3.4.19.12 created 2000]

EC 3.4.19.13

Accepted name: glutathione hydrolase

**Reaction:** glutathione +  $H_2O = L$ -cysteinylglycine + L-glutamate

**Other name(s):** glutathionase; GGT (ambiguous); γ-glutamyltranspeptidase (ambiguous)

Comments: This protein also has EC 2.3.2.2 ( $\gamma$ -glutamyltransferase) activity. The enzyme consists of two chains

that are created by the proteolytic cleavage of a single precursor polypeptide. The N-terminal L-threonine of the C-terminal subunit functions as the active site for both the cleavage and the hydrolysis reactions [2810, 2179, 248, 2180]. The human enzyme also hydrolyses oxidized glutathione and

leukotriene C<sub>4</sub> with similar efficiency, while the mouse enzyme does not [3177, 376].

**References:** [1044, 2810, 2179, 248, 2180, 3177, 376]

[EC 3.4.19.13 created 2011]

EC 3.4.19.14

**Accepted name:** leukotriene-C<sub>4</sub> hydrolase

**Reaction:** leukotriene  $C_4 + H_2O$  = leukotriene  $D_4 + L$ -glutamate

**Other name**(s):  $\gamma$ -glutamyl leukotrienase; GGT5

Comments: The mouse enzyme is specific for leukotriene  $C_4$ , while the human enzyme also has considerable

activity towards glutathione and oxidized glutathione (cf. EC 3.4.19.13, glutathione hydrolase)

[1039, 3177].

**References:** [376, 2628, 1039, 3177]

[EC 3.4.19.14 created 2012]

EC 3.4.19.15

**Accepted name:** desampylase

**Reaction:** an  $N^6$ -[small archaeal modifier protein]-[protein]-L-lysine + H<sub>2</sub>O = a [protein]-L-lysine + a small ar-

chaeal modifier protein

Other name(s): SAMP-protein conjugate cleaving protease; HvJAMM1

**Systematic name:**  $N^6$ -[small archaeal modifier protein]-[protein]-L-lysine hydrolase

**Comments:** The enzyme, characterized from the archaeon *Haloferax volcanii*, specifically cleaves the ubiquitin-

like small modifier proteins SAMP1 and SAMP2 from protein conjugates, hydrolysing the isopeptide bond between a lysine residue of the target protein and the C-terminal glycine of the modifier protein. The enzyme contains  $Zn^{2+}$ . *cf.* EC 3.4.19.12, ubiquitinyl hydrolase 1. In peptidase family M67.

**References:** [1120]

[EC 3.4.19.15 created 2015 as EC 3.4.24.88, transferred 2016 to EC 3.4.19.15]

# EC 3.4.19.16

**Accepted name:** glucosinolate γ-glutamyl hydrolase

**Reaction:** (1) an (E)-1-(glutathion-S-vl)-N-hydroxy- $\omega$ -(methylsulfanyl)alkan-1-imine + H<sub>2</sub>O = an (E)-1-(L-

cysteinylglycin-S-yl)-N-hydroxy- $\omega$ -(methylsulfanyl)alkan-1-imine + L-glutamate

(2) (E)-1-(glutathion-S-yl)-N-hydroxy-2-(1H-indol-3-yl)ethan-1-imine +  $H_2O = (E)$ -1-(L-

cysteinylglycin-S-yl)-N-hydroxy-2-(1H-indol-3-yl)ethan-1-imine + L-glutamate

(3) (glutathion-S-yl)(1H-indol-3-yl)acetonitrile +  $H_2O$  = (L-cysteinylglycin-S-yl)(1H-indol-3-yl)

yl)acetonitrile + L-glutamate

(4) (Z)-1-(glutathion-S-yl)-N-hydroxy-2-phenylethan-1-imine + H<sub>2</sub>O = (Z)-1-(L-cysteinyglycin-S-yl)-

*N*-hydroxy-2-phenylethan-1-imine + L-glutamate

Other name(s): GGP1 (gene name); GGP3 (gene name)

**Comments:** This enzyme, characterized from the plant *Arabidopsis thaliana*, participates in the biosynthesis of the

plant defense compounds glucosinolates and camalexin. It is the only known plant enzyme capable of

hydrolysing the  $\gamma$ -glutamyl residue of glutathione in the cytosol.

**References:** [896]

[EC 3.4.19.16 created 2017]

# EC 3.4.21 Serine endopeptidases

## EC 3.4.21.1

Accepted name: chymotrypsin

**Reaction:** Preferential cleavage: Tyr—, Trp—, Phe—, Leu—

 $\textbf{Other name}(s) \textbf{:} \quad \text{chymotrypsins A and B; } \alpha \textbf{-chymar ophth; avazyme; chymar; chymotest; enzeon; quimar; quimotrase;}$ 

 $\alpha$ -chymar;  $\alpha$ -chymotrypsin A;  $\alpha$ -chymotrypsin

Comments: Chymotrypsin A is formed from cattle and pig chymotrypsinogen A, several iso-forms being pro-

duced according to the number of bonds hydrolysed in the precursor. Chymotrypsin B (formerly listed as EC 3.4.4.6), formed from chymotrypsinogen B, is homologous with chymotrypsin A. Enzymes with specificity similar to that of chymotrypsins A and B have been isolated from many species. In

peptidase family S1 (trypsin family)

**References:** [3184, 245, 172, 2303, 2935]

[EC 3.4.21.1 created 1961 as EC 3.4.4.5 and EC 3.4.4.6, transferred 1972 to EC 3.4.21.1]

# EC 3.4.21.2

**Accepted name:** chymotrypsin C

**Reaction:** Preferential cleavage: Leu+, Tyr+, Phe+, Met+, Trp+, Gln+, Asn+

**Comments:** Formed from pig chymotrypsinogen C, and from cattle subunit II of procarboxypeptidase A. Reacts

more readily with Tos-Leu-CH<sub>2</sub>Cl than Tos-Phe-CH<sub>2</sub>Cl in contrast to chymotrypsin. In peptidase

family S1 (trypsin family)

**References:** [2251, 789, 3184]

EC 3.4.21.3

Accepted name: metridin

**Reaction:** Preferential cleavage: Leu—, Tyr—, Phe—, Met—, Trp—, Gln—, Asn— **Other name(s):** Metridium proteinase A; sea anemone protease A; sea anemone proteinase A

**Comments:** Digestive enzyme from the sea anemone *Metridium senile*.

**References:** [912, 2769]

[EC 3.4.21.3 created 1972]

EC 3.4.21.4

Accepted name: trypsin

**Reaction:** Preferential cleavage: Arg—, Lys—

Other name(s):  $\alpha$ -trypsin;  $\beta$ -trypsin; cocoonase; parenzyme; parenzymol; tryptar; trypure; pseudotrypsin; tryptase;

tripcellim; sperm receptor hydrolase

Comments: The single polypeptide chain cattle  $\beta$ -trypsin is formed from trypsinogen by cleavage of one peptide

bond. Further peptide bond cleavages produce  $\alpha$  and other iso-forms. Isolated as multiple cationic and anionic trypsins [783] from the pancreas of many vertebrates and from lower species including crayfish, insects (cocoonase) and microorganisms (*Streptomyces griseus*) [2389]. Type example of

peptidase family S1.

**References:** [1205, 3101, 2389, 763, 783, 2303, 2883]

[EC 3.4.21.4 created 1961 as EC 3.4.4.4, transferred 1972 to EC 3.4.21.4]

EC 3.4.21.5

Accepted name: thrombin

**Reaction:** Selective cleavage of Arg—Gly bonds in fibrinogen to form fibrin and release fibrinopeptides A and

В

Other name(s): fibrinogenase; thrombose; thrombose; thrombose; thrombin-C; tropostasin; activated blood-coagulation

factor II; blood-coagulation factor IIa; factor IIa; E thrombin; β-thrombin; γ-thrombin

**Comments:** Formed from prothrombin. More selective than trypsin and plasmin. In peptidase family S1 (trypsin

family).

**References:** [175, 1772, 1915, 1753, 1799, 546, 437, 1764]

[EC 3.4.21.5 created 1961 as EC 3.4.4.13, transferred 1972 to EC 3.4.21.5]

EC 3.4.21.6

Accepted name: coagulation factor Xa

**Reaction:** Selective cleavage of Arg—Thr and then Arg—Ile bonds in prothrombin to form thrombin

Other name(s): thrombokinase; prothrombinase; activated blood-coagulation factor X; autoprothrom-

bin C; thromboplastin; plasma thromboplastin; factor Xa; activated Stuart-Prower factor; activated

factor X

**Comments:** A blood coagulation factor formed from the proenzyme factor X by limited proteolysis. Factor X is

a glycoprotein composed of a heavy chain and a light chain, which are generated from a precursor protein by the excision of the tripeptide RKR and held together by one or more disulfide bonds. The activated factor Xa converts prothrombin to thrombin in the presence of factor Va,  $Ca^{2+}$  and phospho-

lipids. Scutelarin (EC 3.4.21.60) has similar specificity, but does not require factor Va.

**References:** [836, 1320, 546, 1289, 1874, 437]

[EC 3.4.21.6 created 1972, modified 2011]

Accepted name: plasmin

**Reaction:** Selective cleavage of Arg—Thr and then Arg—Ile bonds in prothrombin to form thrombin

Other name(s): fibrinase; fibrinolysin; actase; serum tryptase; thrombolysin

Comments: Formed from plasminogen by proteolysis which results in multiple forms of the active plasmin. In

peptidase family S1 (trypsin family).

**References:** [381, 380, 2433]

[EC 3.4.21.7 created 1961 as EC 3.4.4.14, transferred 1972 to EC 3.4.21.7]

[3.4.21.8 Transferred entry, kallikrein, Now EC 3.4.21.34 (plasma kallikrein) and EC 3.4.21.35 (tissue kallikrein)]

[EC 3.4.21.8 created 1972, deleted 1981]

EC 3.4.21.9

Accepted name: enteropeptidase

**Reaction:** Activation of trypsinogen by selective cleavage of Lys<sup>6</sup>—Ile bond

Other name(s): enterokinase

**Comments:** Is not inhibited by protein inhibitors of trypsin. In peptidase family S1 (trypsin family).

**References:** [1685]

[EC 3.4.21.9 created 1961 as EC 3.4.4.8, transferred 1972 to EC 3.4.21.9]

EC 3.4.21.10

Accepted name: acrosin

**Reaction:** Preferential cleavage: Arg—, Lys—

Other name(s): acrosomal proteinase; α-acrosin; β-acrosin; upsilon-acrosin; acrosomal protease; acrosin

amidase

Comments: Occurs in spermatozoa; formed from proacrosin by limited proteolysis. Inhibited by naturally occur-

ring trypsin inhibitors. In peptidase family S1 (trypsin family)

**References:** [2001, 2684, 1413]

[EC 3.4.21.10 created 1972]

[3.4.21.11 Transferred entry. elastase. Now EC 3.4.21.37, leukocyte elastase]

[EC 3.4.21.11 created 1972, deleted 1981]

EC 3.4.21.12

**Accepted name:** α-lytic endopeptidase

**Reaction:** Preferential cleavage: Ala—, Val— in bacterial cell walls, elastin and other proteins

Other name(s): myxobacter  $\alpha$ -lytic proteinase;  $\alpha$ -lytic proteinase;  $\alpha$ -lytic protease; Mycobacterium sorangium  $\alpha$ -lytic

proteinase; Myxobacter 495 α-lytic proteinase; α-lytic proteinase; Myxobacter α-lytic proteinase;

Mycobacterium sorangium α-lytic proteinase

**Comments:** From the myxobacterium *Lysobacter enzymogenes*. In peptidase family S1 (trypsin family)

**References:** [2195, 2303, 708, 262]

[EC 3.4.21.12 created 1972]

[3.4.21.13 Transferred entry. Phaseolus proteinase. Now EC 3.4.16.6, carboxypeptidase D]

[EC 3.4.21.13 created 1972, deleted 1978]

[3.4.21.14 Transferred entry. now EC 3.4.21.67 endopeptidase So]

[EC 3.4.21.14 created 1961 as EC 3.4.4.16, transferred 1972 to EC 3.4.21.14, modified 1986, deleted 1992]

[3.4.21.15] Transferred entry. Aspergillus alkaline proteinase. Now EC 3.4.21.63, oryzin]

[EC 3.4.21.15 created 1972, deleted 1978 (transferred to EC 3.4.21.14, deleted 1992)]

[3.4.21.16 Deleted entry. Alternaria serine proteinase]

[EC 3.4.21.16 created 1972, deleted 1992]

[3.4.21.17 Deleted entry. Arthrobacter serine proteinase]

[EC 3.4.21.17 created 1972, deleted 1978 [transferred to EC 3.4.21.14, deleted 1992]]

[3.4.21.18 Deleted entry. Tenebrio \alpha-proteinase]

[EC 3.4.21.18 created 1972 [EC 3.4.99.24 created 1972, incorporated 1978], deleted 1992]

EC 3.4.21.19

**Accepted name:** glutamyl endopeptidase

**Reaction:** Preferential cleavage: Glu—, Asp—

Other name(s): V8 proteinase; endoproteinase Glu-C; staphylococcal serine proteinase

**Comments:** From *Staphylococcus aureus* strain V8. In appropriate buffer the specificity is restricted to Glu—. In

peptidase family S1 (trypsin family)

**References:** [646, 648, 374]

[EC 3.4.21.19 created 1978]

EC 3.4.21.20

**Accepted name:** cathepsin G

**Reaction:** Specificity similar to chymotrypsin C

**Other name(s):** chymotrypsin-like proteinase; neutral proteinase

**Comments:** From azurophil granules of polymorphonuclear leukocytes. In peptidase family S1 (trypsin family)

**References:** [155, 2879, 1159]

[EC 3.4.21.20 created 1978]

EC 3.4.21.21

Accepted name: coagulation factor VIIa

**Reaction:** Selective cleavage of Arg—Ile bond in factor X to form factor Xa **Other name(s):** blood-coagulation factor VIIa; activated blood coagulation factor VII

**Comments:** Formed from the precursor factor VII. The cattle enzyme is more readily inhibited by diisopropyl flu-

orophosphate than the human [2069]. In peptidase family S1 (trypsin family)

**References:** [2069, 546, 1289, 313]

[EC 3.4.21.21 created 1978]

EC 3.4.21.22

Accepted name: coagulation factor IXa

**Reaction:** Selective cleavage of Arg—Ile bond in factor X to form factor Xa

Other name(s): activated Christmas factor; blood-coagulation factor IXa; activated blood-coagulation factor IX; auto-

prothrombin II; blood platelet cofactor II; activated blood coagulation factor XI

Comments: A chymotrypsin homologue, and one of the  $\gamma$ -carboxyglutamic acid-containing blood coagulation

factors. The proenzyme factor IX is activated by factor XIa. In peptidase family S1 (trypsin family)

**References:** [835, 546, 1706, 437]

[EC 3.4.21.22 created 1978]

[3.4.21.23 Deleted entry. Vipera russelli proteinase]

#### [3.4.21.24 Deleted entry. red cell neutral endopeptidase]

[EC 3.4.21.24 created 1978, deleted 1992]

## EC 3.4.21.25

Accepted name: cucumisin

**Reaction:** Hydrolysis of proteins with broad specificity **Other name(s):** euphorbain; solanain; hurain; tabernamontanain

**Comments:** From the sarcocarp of the musk melon (*Cucumis melo*). In peptidase family S8 (subtilisin family).

Other endopeptidases from plants, which are less well characterized but presumably of serine-type, include euphorbain from *Euphorbia cerifera* [1759], solanain from horse-nettle *Solanum elaeagni-folium* [973], hurain from *Hura crepitans* [1296] and tabernamontanain from *Tabernamontana gran-*

diflora [1295].

**References:** [973, 1296, 1295, 1380, 1379, 1759, 1381]

[EC 3.4.21.25 created 1978 (EC 3.4.21.56 created 1972 as EC 3.4.99.7 transferred 1989 to EC 3.4.21.56, deleted 1992; EC 3.4.99.9 created 1972 deleted 1992; EC 3.4.99.21 created 1972 deleted 1992; EC 3.4.99.23 created 1972 deleted 1992; all covered by EC 3.4.21.25)]

#### EC 3.4.21.26

Accepted name: prolyl oligopeptidase

**Reaction:** Hydrolysis of —Pro— and to a lesser extent —Ala— in oligopeptides

Other name(s): post-proline cleaving enzyme; proline-specific endopeptidase; post-proline endopeptidase; proline

endopeptidase; endoprolylpeptidase; prolyl endopeptidase

Comments: Found in vertebrates, plants and Flavobacterium. Generally cytosolic, commonly activated by thiol

compounds. Type example of peptidase family S9.

**References:** [3103, 2107, 1978, 2409]

[EC 3.4.21.26 created 1978, modified 1981 (EC 3.4.22.18 created 1981, incorporated 1992)]

## EC 3.4.21.27

Accepted name: coagulation factor XIa

**Reaction:** Selective cleavage of Arg—Ala and Arg—Val bonds in factor IX to form factor IXa

Other name(s): blood-coagulation factor XIa; activated blood-coagulation factor XI; activated plasma thromboplastin

antecedent

**Comments:** In peptidase family S1 (trypsin family), and one of the γ-carboxyglutamic acid-containing blood coag-

ulation factors. The proenzyme factor XI is activated by factor XIIa

**References:** [1563, 437, 834]

[EC 3.4.21.27 created 1978]

[3.4.21.28 Transferred entry. Agkistrodon serine proteinase. Now EC 3.4.21.74, venombin A]

[EC 3.4.21.28 created 1978, deleted 1992]

[3.4.21.29 Transferred entry. Bothrops atrox serine proteinase. Now EC 3.4.21.74, venombin A]

[EC 3.4.21.29 created 1978, deleted 1992]

[3.4.21.30 Transferred entry. Crotalus adamanteus serine proteinase. Now EC 3.4.21.74, venombin A]

[EC 3.4.21.30 created 1978, deleted 1992]

[3.4.21.31 Transferred entry. urokinase. Now EC 3.4.21.73, u-plasminogen activator]

[EC 3.4.21.31 created 1972 as EC 3.4.99.26, transferred 1978 to EC 3.4.21.31, deleted 1992]

### EC 3.4.21.32

Accepted name: brachyurin

**Reaction:** Hydrolysis of proteins, with broad specificity for peptide bonds. Native collagen is cleaved about 75%

of the length of the molecule from the N-terminus. Low activity on small molecule substrates of both

trypsin and chymotrypsin

Other name(s): Uca pugilator collagenolytic proteinase; crab protease I; crab protease II

**Comments:** From hepatopancreas of the fiddler crab, *Uca pugilator*. In peptidase family S1 (trypsin family).

Other serine endopeptidases that degrade collagen, but are not listed separately here, include a second endopeptidase from *Uca pugilator* [3158], digestive enzymes from other decapod crustacea

[1475, 1745], and an enzyme from the fungus Entomophthora coronata [1210].

**References:** [1210, 965, 3159, 3158, 1475, 1745]

[EC 3.4.21.32 created 1978]

[3.4.21.33 Deleted entry. Entomophthora collagenolytic proteinase]

[EC 3.4.21.33 created 1978, deleted 1992]

#### EC 3.4.21.34

Accepted name: plasma kallikrein

**Reaction:** Selective cleavage of some Arg—and Lys—bonds, including Lys—Arg and Arg—Ser in (human)

kininogen to release bradykinin

Other name(s): serum kallikrein; kininogenin; kallikrein I; kallikrein II; kininogenase; kallikrein; callicrein; glumorin;

padreatin; padutin; kallidinogenase; bradykininogenase; panceatic kallikrein; onokrein P; dilminal D;

depot-Padutin; urokallikrein; urinary kallikrein

**Comments:** Formed from plasma prokallikrein (Fletcher factor) by factor XIIa. Activates coagulation factors XII,

VII and plasminogen. Selective for Arg > Lys in P1, in small molecule substrates.

**References:** [1100, 1876, 2663, 2608, 2978]

[EC 3.4.21.34 created 1965 as EC 3.4.4.21, transferred 1972 to EC 3.4.21.8, part transferred 1981 to EC 3.4.21.34]

## EC 3.4.21.35

Accepted name: tissue kallikrein

**Reaction:** Preferential cleavage of Arg—bonds in small molecule substrates. Highly selective action to release

kallidin (lysyl-bradykinin) from kininogen involves hydrolysis of Met— or Leu—. The rat enzyme is unusual in liberating bradykinin directly from autologous kininogens by cleavage at two Arg— bonds

[5]

Other name(s): glandular kallikrein; pancreatic kallikrein; submandibular kallikrein; submaxillary kallikrein; kidney

kallikrein; urinary kallikrein; kallikrein; kallikrein; kininogenin; kininogenase; callicrein; glumorin; padreatin; padutin; kallidinogenase; bradykininogenase; depot-padutin; urokallikrein; dilminal

D; onokrein P

**Comments:** Formed from tissue prokallikrein by activation with trypsin. In peptidase family S1 (trypsin family).

A large number of tissue kallikrein-related sequences have been reported for rats [3195] and mice [723], though fewer seem to exist in other mammals. The few that have been isolated and tested on substrates include mouse  $\gamma$ -renin (EC 3.4.21.54), submandibular proteinase A [60, 208], epidermal growth-factor-binding protein, nerve growth factor  $\gamma$ -subunit, rat tonin [3,4,9], submaxillary proteinases A and B [1398], T-kininogenase [3230], kallikreins k7 and k8 [694] and human prostate-

specific antigen (γ-seminoprotein, [26])

**References:** [764, 60, 2259, 1009, 1397, 26, 723, 763, 844, 1398, 119, 226, 400, 889, 208, 3195, 694, 3230]

[EC 3.4.21.35 created 1965 as EC 3.4.4.21, transferred 1972 to EC 3.4.21.8, part transferred 1981 to EC 3.4.21.35]

## EC 3.4.21.36

Accepted name: pancreatic elastase

**Reaction:** Hydrolysis of proteins, including elastin. Preferential cleavage: Ala— **Other name(s):** pancreatopeptidase E; pancreatic elastase I; elastase; elaszym; serine elastase

**Comments:** Formed by activation of proelastase from mammalian pancreas by trypsin. In peptidase family S1

(trypsin family). Formerly included in EC 3.4.21.11

**References:** [2646, 1059, 1407, 214, 251]

[EC 3.4.21.36 created 1981 (EC 3.4.4.7 created 1961, transferred 1972 to EC 3.4.21.11 created 1972, part incorporated 1984)]

#### EC 3.4.21.37

Accepted name: leukocyte elastase

**Reaction:** Hydrolysis of proteins, including elastin. Preferential cleavage Val— ¿ Ala—

Other name(s): lysosomal elastase; neutrophil elastase; polymorphonuclear leukocyte elastase; elastase; elastase; elastase;

serine elastase; lysosomal elastase; granulocyte elastase

Comments: Differs from pancreatic elastase in specificity on synthetic substrates and in inhibitor sensitivity. In

peptidase family S1 (trypsin family). Formerly included in EC 3.4.21.11

**References:** [156, 1059, 2760, 251]

[EC 3.4.21.37 created 1981 (EC 3.4.4.7 created 1961, transferred 1972 to EC 3.4.21.11 created 1972, part incorporated 1984)]

#### EC 3.4.21.38

Accepted name: coagulation factor XIIa

Reaction: Selective cleavage of Arg—Ile bonds in factor VII to form factor VIIa and factor XI to form factor

XIa

Other name(s): Hageman factor (activated); blood-coagulation factor XIIf; activated  $\beta$  blood-coagulation factor XII;

prealbumin activator; Hageman factor β-fragment; Hageman factor fragment HFf; blood-coagulation

factor XIIaβ; prekallikrein activator; kallikreinogen activator

**Comments:** Also activates plasminogen and plasma prokallikrein. Formed from the proenzyme, factor XII, by

plasma kallikrein or factor XIIa. In peptidase family S1 (trypsin family)

**References:** [837, 437, 2342, 833, 2662]

[EC 3.4.21.38 created 1981]

## EC 3.4.21.39

**Accepted name:** chymase

**Reaction:** Preferential cleavage: Phe— ¿ Tyr— ¿ Trp— ¿ Leu—

Other name(s): mast cell protease I; skeletal muscle protease; skin chymotryptic proteinase; mast cell serine pro-

teinase, chymase; skeletal muscle (SK) protease

Comments: In mast cell granules. In peptidase family S1 (trypsin family)

**References:** [3212, 2321, 1327]

[EC 3.4.21.39 created 1981]

[3.4.21.40 Deleted entry. submandibular proteinase A]

[EC 3.4.21.40 created 1981, deleted 1992]

# EC 3.4.21.41

**Accepted name:** complement subcomponent  $C^{\overline{1r}}$ 

**Reaction:** Selective cleavage of Lys(or Arg)—Ile bond in complement subcomponent C1s to form  $C^{\overline{1s}}$  (EC

3.4.21.42)

**Other name(s):** activated complement C1r;  $C^{\overline{1r}}$  esterase; activated complement C1r

**Comments:** Activated from proenzyme  $C^{\overline{1r}}$  in plasma during activation of the complement system by the "classi-

cal" route. In peptidase family S1 (trypsin family)

**References:** [2664, 1662, 2000]

# [EC 3.4.21.41 created 1981]

EC 3.4.21.42

**Accepted name:** complement subcomponent C<sup>1s</sup>

Reaction: Cleavage of Arg—Ala bond in complement component C4 to form C4a and C4b, and Lys(or

Arg)—Lys bond in complement component C2 to form C2a and C2b: the "classical" pathway C3

convertase

**Other name(s):** C1 esterase; activated complement C1s; complement  $C^{1r}$ 

Comments: Activated from proenzyme C1s in plasma by complement subcomponent  $C^{1r}$ . In peptidase family S1

(trypsin family)

**References:** [2664, 1767, 2000, 2684]

[EC 3.4.21.42 created 1981]

EC 3.4.21.43

Accepted name: classical-complement-pathway C3/C5 convertase

**Reaction:** Selective cleavage of Arg—Ser bond in complement component C3 α-chain to form C3a and C3b,

and Arg + bond in complement component C5  $\alpha$ -chain to form C5a and C5b

Other name(s): C3 convertase;  $C^{\overline{42}}$ ; C4b,2a; C5 convertase;  $C^{\overline{423}}$ ; C4b,2a,3b; C42; C5 convertase; C423; C4b,2a,3b;

complement C.hivin.4.hivin2; complement C3 convertase

Comments: A complex of complement fragments C4b, C2a and C2b. C2a contains the active site, C2b the site for

C4b binding. C2a and C2b are formed by cleavage of proenzyme C2 by complement subcomponent  $C^{\overline{1s}}$ . Cleavage of C5 requires complement fragment C3b which binds C5 and renders it susceptible to cleavage by the C4b,2a complex. Includes former EC 3.4.21.44. Complement component C2a is in

peptidase family S1 (trypsin family)

**References:** [1420, 2000]

[EC 3.4.21.43 created 1981 (EC 3.4.21.44 created 1981, incorporated 1984)]

[3.4.21.44 Transferred entry. complement component C5 convertase. Now EC 3.4.21.43, classical-complement-pathway C3/C5 convertase]

[EC 3.4.21.44 created 1981, deleted 1984]

EC 3.4.21.45

**Accepted name:** complement factor I

**Reaction:** Inactivates complement subcomponents C3b, iC3b and C4b by proteolytic cleavage

Other name(s): complement component C3b inactivator; C3b inactivator; C3b/C4b inactivator; C3bINA; complement

C3b/C4b inactivator; complement C4b inactivator; conglutinogen-activating factor C; complement

C3b inactivator; factor I; complement C4bi

Comments: Cleavage of complement subcomponent C3b requires its binding to cofactor factor H or complement

receptor CR1; cleavage of iC3b requires complement receptor CR1; cleavage of C4b requires C4b-

binding protein. In peptidase family S1 (trypsin family)

**References:** [2023, 509, 2000]

[EC 3.4.21.45 created 1981]

EC 3.4.21.46

**Accepted name:** complement factor D

**Reaction:** Selective cleavage of Arg—Lys bond in complement factor B when in complex with complement

subcomponent C3b or with cobra venom factor

Other name(s): C3 proactivator convertase; properdin factor D esterase; factor D; factor D (complement)

**Comments:** A component of the alternative pathway of complement activation. This reaction is analogous to the

activation of complement component C2 by complement subcomponent  $C^{\overline{1s}}$ . In peptidase family S1

(trypsin family)

[2401, 2000] **References:** 

[EC 3.4.21.46 created 1981]

EC 3.4.21.47

alternative-complement-pathway C3/C5 convertase Accepted name:

Cleavage of Arg—Ser bond in complement component C3 α-chain to yield C3a and C3b, and Arg— **Reaction:** 

bond in complement component C5  $\alpha$ -chain to yield C5a and C5b

Other name(s): complement component C3/C5 convertase (alternative); proenzyme factor B; properdin factor B; C3

> proactivator; glycine-rich β-glycoprotein; heat-labile factor; C3 convertase; C3b,Bb,CVF,Bb,C5 convertase; (C3b)n,Bb; complement C 3(C 5) convertase (amplification); alternative complement pathway C3(C5) convertase; C5 convertase; CVF,Bb; (CVF)-dependent glycine-rich-β-glucoprotein; co-

bra venom factor-dependent C3 convertase

A bimolecular complex of complement fragment Bb with either C3b or cobra venom factor; Bb con-**Comments:** 

> tains the active site. Bb is formed by cleavage of proenzyme factor B by factor D. Cleavage of complement component C5 requires additional C3b which binds C5 and renders it susceptible to cleavage by C3b,Bb complex. C3b,Bb is stabilized in plasma by factor P. Complement factor B is in peptidase

family S1 (trypsin family)

**References:** [1421, 1980, 2000]

[EC 3.4.21.47 created 1981]

EC 3.4.21.48

Accepted name: cerevisin

> **Reaction:** Hydrolysis of proteins with broad specificity, and of Bz-Arg-OEt Ac-Tyr-OEt. Does not hydrolyse

> > peptide amides

Other name(s): yeast proteinase B; proteinase yscB; baker's yeast proteinase B; brewer's yeast proteinase; peptidase  $\beta$ **Comments:** From Saccharomyces cerevisiae (baker's yeast, brewer's yeast). In peptidase family S8 (subtilisin

family), but contains a Cys residue near the active site His, and is inhibited by mercurials. Proteinase

ycaB is a similar enzyme from the yeast Candida albicans [738]

[751, 1505, 738, 1944] **References:** 

[EC 3.4.21.48 created 1972 as EC 3.4.22.9, transferred 1981 to EC 3.4.21.48]

EC 3.4.21.49

Accepted name: hypodermin C

> Reaction: Hydrolysis of proteins including native collagen at —Ala bond leaving an N-terminal (75%) and a

C-terminal (25%) fragment

Other name(s): Hypoderma collagenase

**Comments:** From the larva of a warble fly, Hypoderma lineatum. Little action on small molecule substrates of

trypsin, chymotrypsin, elastase or microbial collagenases. In peptidase family S1 (trypsin family)

**References:** [1613, 1615, 1614]

[EC 3.4.21.49 created 1981]

EC 3.4.21.50

Accepted name: lysyl endopeptidase

> Reaction: Preferential cleavage: Lys—, including -Lys—Pro-

Other name(s): Achromobacter proteinase I (also see Comment); Achromobacter lyticus alkaline proteinase I; pro-

tease I; achromopeptidase; lysyl bond specific proteinase

Comments: From Achromobacter lyticus [2984]. Enzymes with similar specificity are produced by Lysobacter en-

zymogenes (Endoproteinase Lys-C; [1315]) and Pseudomonas aeruginosa (Ps-1; [692]). In peptidase

family S1 (trypsin family)

**References:** [1831, 1830, 1315, 692, 2157, 2984]

[EC 3.4.21.50 created 1983]

[3.4.21.51 Deleted entry. Leukocyte-membrane neutral endopeptidase]

[EC 3.4.21.51 created 1984, deleted 1992]

[3.4.21.52 Deleted entry. Cathepsin R]

[EC 3.4.21.52 created 1981 as EC 3.4.99.33, transferred 1984 to EC 3.4.21.52, deleted 1992]

EC 3.4.21.53

Accepted name: endopeptidase La

**Reaction:** Hydrolysis of proteins in presence of ATP

**Other name(s):** ATP-dependent serine proteinase; lon proteinase; protease La; proteinase La; ATP-dependent lon

proteinase; ATP-dependent protease La; *Escherichia coli* proteinase La; *Escherichia coli* serine proteinase La; gene lon protease; gene lon proteins; PIM1 protease; PIM1 proteinase; serine protease La Product of the *lon* gene in *Escherichia coli*. ATP hydrolysis is linked with peptide bond hydrolysis;

vanadate inhibits both reactions. Type example of peptidase family S16. A similar enzyme occurs in

animal mitochondria

**References:** [587, 1602, 432]

[EC 3.4.21.53 created 1986]

EC 3.4.21.54

**Accepted name:** γ-renin

**Comments:** 

**Reaction:** Cleavage of the Leu—Leu bond in synthetic tetradecapeptide renin substrate (horse), to produce an-

giotensin I, but not active on natural angiotensinogen, unlike renin (EC 3.4.23.15). Also hydrolyses

Bz-Arg-*p*-nitroanilide

**Comments:** A member of the tissue kallikrein family, from submandibular glands of male mice. In peptidase fam-

ily S1 (trypsin family)

**References:** [2298, 653]

[EC 3.4.21.54 created 1986]

EC 3.4.21.55

Accepted name: venombin AB

**Reaction:** Selective cleavage at Arg—bonds in fibrinogen to form fibrin and release fibrinopeptides A and B

Other name(s): gabonase; okinaxobin II; Bitis gabonica venom serine proteinase; afaâcytin

**Comments:** From the venom of the Gaboon viper *Bitis gabonica*. Activates Factor XIII. Not inhibited by an-

tithrombin III/heparin or hirudin, unlike EC 3.4.21.5, thrombin

**References:** [2286]

[EC 3.4.21.55 created 1989]

[3.4.21.56 Deleted entry. euphorbain. Now considered EC 3.4.21.25, cucumisin]

[EC 3.4.21.56 created 1972 as EC 3.4.99.7, transferred 1989 to EC 3.4.21.56, deleted 1992]

EC 3.4.21.57

Accepted name: leucyl endopeptidase

**Reaction:** Hydrolysis of proteins. Preferential cleavage: Leu—in small molecule substrates

Other name(s): plant Leu-proteinase; leucine-specific serine proteinase; leucine endopeptidase; spinach serine pro-

teinase (leucine specific); spinach leucine-specific serine proteinase; Leu-proteinase

**Comments:** From leaves of the spinach plant (*Spinacia oleracea*)

**References:** [19, 18]

[EC 3.4.21.57 created 1989]

[3.4.21.58 Deleted entry. prohormone serine proteinase]

[EC 3.4.21.58 created 1989, deleted 1992]

EC 3.4.21.59

**Accepted name:** tryptase

**Reaction:** Preferential cleavage: Arg—, Lys—, but with more restricted specificity than trypsin

Other name(s): mast cell tryptase; mast cell protease II; skin tryptase; lung tryptase; pituitary tryptase; mast cell neu-

tral proteinase; mast cell tryptase; mast cell neutral proteinase; mast cell serine proteinase II; mast cell

proteinase II; mast cell serine proteinase tryptase; rat mast cell protease II; tryptase M

Comments: Occurs as a tetrameric molecule with high affinity for heparin, in mast cell granules. In peptidase fam-

ily S1 (trypsin family). Not inhibited by  $\alpha_1$ -proteinase inhibitor or  $\alpha_2$ -macroglobulin

**References:** [2878, 1436, 506, 1065, 3050]

[EC 3.4.21.59 created 1992]

EC 3.4.21.60

Accepted name: scutelarin

**Reaction:** Selective cleavage of Arg—Thr and Arg—Ile in prothrombin to form thrombin and two inactive

fragments

Other name(s): taipan activator; Oxyuranus scutellatus prothrombin-activating proteinase

**Comments:** From the venom of the Taipan snake (Oxyuranus scutellatus). Converts prothrombin to thrombin.

Specificity is similar to that of Factor Xa (EC 3.4.21.6). However, unlike Factor Xa this enzyme can cleave its target in the absence of coagulation Factor Va. Activity is potentiated by phospholipid and  $Ca^{2+}$  which binds via  $\gamma$ -carboxyglutamic acid residues. Similar enzymes are known from the venom of other Australian elapid snakes, including *Pseudonaja textilis textilis*, *Oxyuranus microlepidotus* and

Demansia nuchalis affinis.

**References:** [3094, 2739]

[EC 3.4.21.60 created 1978 as EC 3.4.99.28, transferred 1992 to EC 3.4.21.60, modified 2010, modified 2011]

EC 3.4.21.61

Accepted name: kexin

**Reaction:** Cleavage of -Lys-Arg + and -Arg-Arg + bonds to process yeast  $\alpha$ -factor pheromone and killer toxin

precursors

Other name(s): yeast KEX2 protease; proteinase yscF; prohormone-processing endoprotease; paired-basic en-

dopeptidase; yeast cysteine proteinase F (misleading); paired-basic endopeptidase; andrenorphin-Gly-generating enzyme; endoproteinase Kex2p; gene KEX2 dibasic proteinase; Kex 2p proteinase; Kex2 endopeptidase; Kex2 endoprotease; Kex2 endoproteinase; Kex2 proteinase; Fex2-like precursor protein processing endoprotease; prohormone-processing KEX2 proteinase; prohormone-processing proteinase; proprotein convertase; protease KEX2; Kex2 proteinase; Kex2-

like endoproteinase

**Comments:** A Ca<sup>2+</sup>-activated peptidase of peptidase family S8, containing Cys near the active site His, and inhib-

ited by p-mercuribenzoate. Similar enzymes occur in mammals.

**References:** [1346, 10, 1940, 862, 1941]

#### EC 3.4.21.62

Accepted name: subtilisin

**Reaction:** Hydrolysis of proteins with broad specificity for peptide bonds, and a preference for a large uncharged

residue in P1. Hydrolyses peptide amides

Other name(s): alcalase; alcalase 0.6L; alcalase 2.5L; ALK-enzyme; bacillopeptidase A; bacillopeptidase B; Bacillus

subtilis alkaline proteinase bioprase; bioprase AL 15; bioprase APL 30; colistinase; (see also comments); subtilisin J; subtilisin S41; subtilisin Sendai; subtilisin GX; subtilisin E; subtilisin BL; genenase I; esperase; maxatase; alcalase; thermoase PC 10; protease XXVII; thermoase; superase; subtilisin DY; subtilopeptidase; SP 266; savinase 8.0L; savinase 4.0T; kazusase; protease VIII; opticlean; Bacillus subtilis alkaline proteinase; protin A 3L; savinase; savinase 16.0L; savinase 32.0 L EX; ori-

entase 10B; protease S

**Comments:** Subtilisin is a serine endopeptidase, type example of peptidase family S8. It contains no cysteine

residues (although these are found in homologous enzymes). Species variants include subtilisin BPN' (also subtilisin B, subtilopeptidase B, subtilopeptidase C, Nagarse, Nagarse proteinase, subtilisin Novo, bacterial proteinase Novo) and subtilisin Carlsberg (subtilisin A, subtilopeptidase A, alcalase Novo). Similar enzymes are produced by various *Bacillus subtilis* strains and other *Bacillus* species

[2215, 2275]

**References:** [2215, 1814, 2275, 2063, 1232, 2303]

[EC 3.4.21.62 created 1992 (EC 3.4.21.14 created 1961 as EC 3.4.4.16, transferred 1972 to EC 3.4.21.14, modified 1986, part incorporated 1992)]

#### EC 3.4.21.63

Accepted name: oryzin

**Reaction:** Hydrolysis of proteins with broad specificity, and of Bz-Arg-OEt > Ac-Tyr-OEt. Does not hydrolyse

peptide amides

Other name(s): Aspergillus alkaline proteinase; aspergillopeptidase B; API 21; aspergillopepsin B; aspergillopepsin F;

Aspergillus candidus alkaline proteinase; Aspergillus flavus alkaline proteinase; Aspergillus melleus semi-alkaline proteinase; Aspergillus oryzae alkaline proteinase; Aspergillus parasiticus alkaline proteinase; Aspergillus serine proteinase; Aspergillus sydowi alkaline proteinase; Aspergillus soya alkaline proteinase; Aspergillus melleus alkaline proteinase; Aspergillus sulphureus alkaline proteinase; prozyme; P 5380; kyorinase; seaprose S; semi-alkaline protease; sumizyme MP; prozyme 10; ono-

prose; onoprose SA; protease P; promelase

Comments: A peptidase of family S8 (subtilisin family), not containing cysteine, that is the predominant extracel-

lular alkaline endopeptidase of the mold Aspergillus oryzae. Identical or closely related enzymes are

produced by A. flavus and A. sojae [2,3,4]

**References:** [2035, 1089, 2998, 1969, 2735]

[EC 3.4.21.63 created 1992 (EC 3.4.21.14 created 1961 as EC 3.4.4.16, transferred 1972 to EC 3.4.21.14, modified 1986, part incorporated 1992)]

# EC 3.4.21.64

**Accepted name:** peptidase K

**Reaction:** Hydrolysis of keratin, and of other proteins with subtilisin-like specificity. Hydrolyses peptide amides **Other name(s):** *Tritirachium* alkaline proteinase; *Tritirachium album* serine proteinase; proteinase K; *Tritirachium* 

album proteinase K; endopeptidase K

Comments: From the mold *Tritirachium album* Limber. A peptidase of family S8 (subtilisin family) containing

two disulfide bridges and one free Cys near the active site His. Formerly included in EC 3.4.21.14

**References:** [677, 1972, 1530, 1310, 209]

[EC 3.4.21.64 created 1992 (EC 3.4.21.14 created 1961 as EC 3.4.4.16, transferred 1972 to EC 3.4.21.14, modified 1986, part incorporated 1992)]

### EC 3.4.21.65

Accepted name: thermomycolin

**Reaction:** Rather nonspecific hydrolysis of proteins. Preferential cleavage: Ala—, Tyr—, Phe— in small

molecule substrates

**Other name(s):** thermomycolase

Comments: A peptidase of family S8 (subtilisin family) from the thermophilic fungus Malbranchea pulchella

var. sulfurea containing Cys, but not inhibited by p-mercuribenzoate. Very thermostable. Formerly

included in EC 3.4.21.14

**References:** [884]

[EC 3.4.21.65 created 1992 (EC 3.4.21.14 created 1961 as EC 3.4.4.16, transferred 1972 to EC 3.4.21.14, modified 1986, part incorporated 1992)]

# EC 3.4.21.66

**Accepted name:** thermitase

**Reaction:** Hydrolysis of proteins, including collagen

**Other name(s):** thermophilic *Streptomyces* serine proteinase; *Thermoactinomyces vulgaris* serine proteinase

**Comments:** A peptidase of family S8 (subtilisin family) from *Thermoactinomyces vulgaris* containing a single

Cys, near the active site His, and inhibited by p-mercuribenzoate. The N-terminal extension of the polypeptide chain relative to subtilisin contributes to  $Ca^{2+}$ -binding and the high thermostability. The amino acid composition and properties of the thermostable enzyme from *Streptomyces rectus* var.

proteolyticus (formerly included in EC 3.4.21.14) are closely similar [1942, 264].

**References:** [1942, 264, 1474, 1883, 2899]

[EC 3.4.21.66 created 1992]

#### EC 3.4.21.67

Accepted name: endopeptidase So

**Reaction:** Hydrolysis of proteins, but not Bz-Tyr-OEt, Ac-Phe-β-naphthylester, or Bz-Arg-OEt **Other name(s):** *E. coli* cytoplasmic proteinase; proteinase So; *Escherichia coli* serine proteinase So

Comments: An Escherichia coli cytoplasmic endopeptidase formerly included in EC 3.4.21.14. Inhibited by Tos-

Phe-CH<sub>2</sub>Cl, but not by Tos-Lys-CH<sub>2</sub>Cl

**References:** [943, 447]

[EC 3.4.21.67 created 1992 (EC 3.4.21.14 created 1961 as EC 3.4.4.16, transferred 1972 to EC 3.4.21.14, modified 1986, part incorporated 1992)]

# EC 3.4.21.68

**Accepted name:** *t*-plasminogen activator

**Reaction:** Specific cleavage of Arg—Val bond in plasminogen to form plasmin

Other name(s): tissue plasminogen activator; plasminogen activator, tissue-type; tissue-type plasminogen activator;

tPA; t-PA

**Comments:** A peptidase of family S1 (trypsin family) from a wide variety of mammalian tissues, especially en-

dothelial cells. Secreted as a single chain precursor which is cleaved to a two-chain form by plasmin. Activity is considerably enhanced by fibrin. Formerly included in EC 3.4.21.31 and EC 3.4.99.26

**References:** [2255, 1738, 2261, 3059, 891, 467]

[EC 3.4.21.68 created 1972 as EC 3.4.99.26, transferred 1978 as EC 3.4.21.31, part transferred 1992 to EC 3.4.21.68]

#### EC 3.4.21.69

**Accepted name:** protein C (activated)

**Reaction:** Degradation of blood coagulation factors Va and VIIIa

Other name(s): blood-coagulation factor XIVa; activated blood coagulation factor XIV; activated protein C; autopro-

thrombin II-A; protein Ca; APC; GSAPC

**Comments:** A peptidase of family S1 (trypsin family), one of the γ-carboxyglutamic acid-containing coagulation

factors. Formed from protein C, the proenzyme that circulates in plasma, by the action of a complex of thrombin with thrombomodulin, or by serine endopeptidases present in several snake venoms

**References:** [716, 717]

[EC 3.4.21.69 created 1992]

EC 3.4.21.70

Accepted name: pancreatic endopeptidase E

**Reaction:** Preferential cleavage: Ala—. Does not hydrolyse elastin

Other name(s): cholesterol-binding proteinase; proteinase E; cholesterol-binding serine proteinase; pancreatic pro-

tease E; pancreatic proteinase E; cholesterol-binding pancreatic proteinase; CBPP; pancreas E pro-

teinase

Comments: A peptidase of family S1 (trypsin family) from pancreatic juice. Unlike elastases, has an acidic pI.

Binds cholesterol

**References:** [1792, 2626]

[EC 3.4.21.70 created 1992]

EC 3.4.21.71

Accepted name: pancreatic elastase II

**Reaction:** Preferential cleavage: Leu—, Met— and Phe—. Hydrolyses elastin

Other name(s): pancreatic elastase 2

**Comments:** A peptidase of family S1 (trypsin family) formed by activation of proelastase II from mammalian pan-

creas by trypsin. Usually, only one of the pancreatic elastases (see also EC 3.4.21.36) is expressed in a

given species; human pancreatic elastase is of type II

**References:** [784, 2643]

[EC 3.4.21.71 created 1992]

EC 3.4.21.72

**Accepted name:** IgA-specific serine endopeptidase

Reaction: Cleavage of immunoglobulin A molecules at certain Pro—bonds in the hinge region. No small

molecule substrates are known

Other name(s): IgA protease; IgA proteinase; IgA-specific proteinase; immunoglobulin A protease; immunoglobulin

A proteinase

Comments: Species variants differing slightly in specificity are secreted by Gram-negative bacteria Neisseria gon-

orrhoeae and Haemophilus influenzae. Type example of peptidase family S6. Some other bacterial endopeptidases with similar specificity are of metallo- type (see EC 3.4.24.13, IgA-specific metal-

loendopeptidase)

**References:** [2292, 116]

[EC 3.4.21.72 created 1992]

EC 3.4.21.73

Accepted name: u-plasminogen activator

**Reaction:** Specific cleavage of Arg—Val bond in plasminogen to form plasmin

Other name(s): urokinase; urinary plasminogen activator; cellular plasminogen activator; urokinase-type plasminogen

activator; double-chain urokinase-type plasminogen activator; two-chain urokinase-type plasminogen

activator; urokinase plasminogen activator; uPA; u-PA; abbokinase; urinary esterase A

Comments: Formed from the inactive precursor by action of plasmin or plasma kallikrein. Differs in structure

from t-plasminogen activator (EC 3.4.21.68), and does not bind to fibrin. In peptidase family S1

(trypsin family). Formerly included in EC 3.4.21.31 and EC 3.4.99.26

**References:** [1740, 1738, 2505, 467, 1686]

#### EC 3.4.21.74

Accepted name: venombin A

**Reaction:** Selective cleavage of Arg—bond in fibrinogen, to form fibrin, and release fibrinopeptide A. The

specificity of further degradation of fibrinogen varies with species origin of the enzyme

Other name(s): \(\alpha\)-fibrinogenase; habutobin; zinc metalloproteinase Cbfib1.1; zinc metalloproteinase Cbfib1.2; zinc

metalloproteinase Cbfib2; ancrod; (see also Comments)

**Comments:** A somewhat thrombin-like enzyme from venoms of snakes of the viper/rattlesnake group. Species

variants of the enzyme include ancrod from *Agkistrodon rhodostoma* (Malayan pit viper) (formerly EC 3.4.21.28) [2106], batroxobin from *Bothrops atrox* (South American pit viper) (formerly EC 3.4.21.29) [2774, 1269] and crotalase from *Crotalus adamanteus* (Eastern diamondback rattlesnake) (formerly EC 3.4.21.30) [1813, 2665]. In peptidase family S1 (trypsin family). Does not require acti-

vation by Ca<sup>2+</sup>

**References:** [2106, 2774, 1813, 2665, 1269]

[EC 3.4.21.74 created 1992 (EC 3.4.21.28, EC 3.4.21.29 and 3.4.21.30 all created 1978 and incorporated 1992)]

#### EC 3.4.21.75

Accepted name: furin

**Reaction:** Release of mature proteins from their proproteins by cleavage of -Arg-Xaa-Yaa-Arg- bonds, where

Xaa can by any amino acid and Yaa is Arg or Lys. Releases albumin, complement component C3 and

von Willebrand factor from their respective precursors

**Other name(s):** prohormone convertase; dibasic processing enzyme; PACE; paired basic amino acid cleaving enzyme;

paired basic amino acid converting enzyme; serine proteinase PACE; PC1; SPC3; proprotein conver-

tase

Comments: One of a group of peptidases in peptidase family S8 (subtilisin family) that is structurally and func-

tionally similar to kexin. All are activated by  $Ca^{2+}$ , contain Cys near the active site His, and are inhibited by p-mercuribenzoate. At least three related enzymes are recognized in mammals: PC2, PC3 and

PC4, which have somewhat different specificities

**References:** [564, 563, 1079, 2604, 2761]

[EC 3.4.21.75 created 1993]

# EC 3.4.21.76

Accepted name: myeloblastin

**Reaction:** Hydrolysis of proteins, including elastin, by preferential cleavage: -Ala+> -Val+

Other name(s): leukocyte proteinase 3; leukocyte proteinase 4; Wegener's granulomatosis autoantigen; proteinase PR-

3; proteinase-3; PMNL proteinase

**Comments:** From polymorphonuclear leukocyte granules. In peptidase family S1 (trypsin family). Not inhibited

by secretory leukocyte proteinase inhibitor

**References:** [1584, 2374, 314, 1367]

[EC 3.4.21.76 created 1993]

## EC 3.4.21.77

Accepted name: semenogelase

**Reaction:** Preferential cleavage: -Tyr—

**Other name(s):** prostate-specific antigen; α-seminoprotein; seminin; P-30 antigen; antigen (human clone HPSA-1

prostate-specific protein moiety reduced);  $\gamma$ -seminoglycoprotein (human protein moiety reduced);  $\gamma$ -SM; antigen PSA (human prostate-specific); human glandular kallikrein; antigen PSA (human clone

5P1 protein moiety reduced)

**Comments:** A peptidase of family S1 (trypsin family) from seminal plasma. Slowly inhibited by  $\alpha_1$ -

antichymotrypsin

**References:** [613, 443]

[EC 3.4.21.77 created 1993]

EC 3.4.21.78

Accepted name: granzyme A

**Reaction:** Hydrolysis of proteins, including fibronectin, type IV collagen and nucleolin. Preferential cleavage:

-Arg—, -Lys—>> -Phe— in small molecule substrates

Other name(s): CTLA3; HuTPS; T-cell associated protease 1; cytotoxic T lymphocyte serine protease; TSP-1; T-cell

derived serine proteinase

Comments: From cytotoxic T lymphocyte granules. In peptidase family S1 (trypsin family). The human enzyme

does not cleave Phe-

**References:** [2667, 895, 2138]

[EC 3.4.21.78 created 1993]

EC 3.4.21.79

**Accepted name:** granzyme B

**Reaction:** Preferential cleavage: -Asp+>> -Asn+> -Met+, -Ser+

**Other name(s):** CTLA1; CCPII; cytotoxic cell proteinase-1; granzyme G; granzyme H; CCP<sub>1</sub> proteinase

**Comments:** From cytotoxic T lymphocyte granules. In peptidase family S1 (trypsin family)

**References:** [2568, 2138, 2297]

[EC 3.4.21.79 created 1993]

EC 3.4.21.80

**Accepted name:** streptogrisin A

**Reaction:** Hydrolysis of proteins with specificity similar to chymotrypsin

Other name(s): Streptomyces griseus protease A; protease A; proteinase A; Streptomyces griseus proteinase A; Streptomyces A; Streptomyces proteinase A; Streptomyces A; Strept

tomyces griseus serine proteinase 3; Streptomyces griseus serine proteinase A

**Comments:** From *Streptomyces griseus*. A component of Pronase, in family S1 (trypsin family). Not inhibited by

Tos-Phe-CH<sub>2</sub>Cl or ovomucoid

**References:** [1331, 2656, 1306, 576, 1110]

[EC 3.4.21.80 created 1993]

EC 3.4.21.81

Accepted name: streptogrisin B

**Reaction:** Hydrolysis of proteins with trypsin-like specificity

**Other name(s):** Streptomyces griseus protease B; pronase B; serine proteinase B; Streptomyces griseus proteinase B;

Streptomyces griseus proteinase 1; Streptomyces griseus serine proteinase B

**Comments:** From *Streptomyces griseus*. A component of Pronase, in peptidase family S1 (trypsin family), distinct

from Streptomyces trypsin

**References:** [1350, 845, 2390, 1110, 975]

[EC 3.4.21.81 created 1993]

EC 3.4.21.82

Accepted name: glutamyl endopeptidase II

**Reaction:** Preferential cleavage: -Glu+>> -Asp+. Preference for Pro or Leu at P2 and Phe at P3. Cleavage

of -Glu-Asp- and -Glu-Pro- bonds is slow

Other name(s): GluSGP

**Comments:** From *Streptomyces griseus*. A peptidase of family S1 (trypsin family). Inhibited by [Leu<sup>18</sup> $\rightarrow$ Glu]-

modified turkey ovomucoid third domain

**References:** [3300, 1506, 2025, 2819, 284]

[EC 3.4.21.82 created 1993]

EC 3.4.21.83

**Accepted name:** oligopeptidase B

**Reaction:** Hydrolysis of -Arg—, -Lys— bonds in oligopeptides, even when P1' residue is proline

Other name(s): protease II; Escherichia coli alkaline proteinase II; protease II

Comments: Known from Escherichia coli. Inhibited by Tos-Lys-CH<sub>2</sub>Cl. In peptidase family S9 (prolyl oligopep-

tidase family)

**References:** [1378]

[EC 3.4.21.83 created 1993]

EC 3.4.21.84

**Accepted name:** limulus clotting factor  $\overline{C}$ 

**Reaction:** Selective cleavage of -Arg<sup>103</sup>—Ser- and -Ile<sup>124</sup>—Ile- bonds in limulus clotting factor B to form fac-

tor  $\overline{B}$ . Cleavage of -Pro-Arg—bonds in synthetic substrates

**Other name(s):** factor C; limulus factor C

**Comments:** From the hemocyte granules of the horseshoe crabs *Limulus* and *Tachypleus*. Factor C is activated

by Gram-negative bacterial lipopolysaccharides and chymotrypsin. Inhibited by antithrombin III. In

peptidase family S1 (trypsin family)

**References:** [2048, 2011, 2931]

[EC 3.4.21.84 created 1993]

EC 3.4.21.85

**Accepted name:** limulus clotting factor  $\overline{B}$ 

**Reaction:** Selective cleavage of -Arg<sup>98</sup>—Ile- bond in limulus proclotting enzyme to form active clotting enzyme **Comments:** From the hemocyte granules of the horseshoe crabs *Limulus* and *Tachypleus*. Factor B is activated by

limulus clotting factor  $\overline{C}$ . In peptidase family S1 (trypsin family)

**References:** [2046]

[EC 3.4.21.85 created 1993]

EC 3.4.21.86

**Accepted name:** limulus clotting enzyme

**Reaction:** Selective cleavage of -Arg<sup>18</sup>— and -Arg<sup>47</sup>— bonds in coagulogen to form coagulin and fragments

Other name(s): clotting enzyme

Comments: From the hemocyte granules of horseshoe crabs Limulus and Tachypleus. Proclotting enzyme is acti-

vated by limulus clotting factor . In peptidase family S1 (trypsin family)

**References:** [2010, 2931]

[EC 3.4.21.86 created 1993]

[3.4.21.87 Transferred entry. omptin. Now EC 3.4.23.49, omptin. The enzyme is not a serine protease, as thought previously, but an aspartate protease]

[EC 3.4.21.87 created 1993, deleted 2006]

EC 3.4.21.88

Accepted name: repressor LexA

**Reaction:** Hydrolysis of Ala<sup>84</sup>—Gly bond in repressor LexA

Other name(s): LexA repressor

Comments: RecA protein and single-stranded DNA are required for activity, which is attributed to a Ser/Lys dyad

[2690]. The LexA protein represses the SOS regulon, which regulates the genes involved in DNA repair. In the presence of single-stranded DNA, the RecA protein interacts with repressor LexA, causing it to undergo an autocatalytic cleavage which disrupts the DNA-binding part of the repressor, and inactivates it. The consequent derepression of the SOS regulon leads to DNA repair. This peptidase activity of LexA was previously attributed to the RecA protein. Type example of peptidase family S24

**References:** [1178, 2690, 1442, 1714]

[EC 3.4.21.88 created 1995]

EC 3.4.21.89

Accepted name: signal peptidase I

**Reaction:** Cleavage of hydrophobic, N-terminal signal or leader sequences

**Other name(s):** leader peptidase I; signal proteinase; *Escherichia coli* leader peptidase; eukaryotic signal peptidase;

eukaryotic signal proteinase; leader peptidase; leader peptide hydrolase; leader proteinase; signal peptidase; pilin leader peptidase; SPC; prokaryotic signal peptidase; prokaryotic leader peptidase; HOSP; prokaryotic signal proteinase; propeptidase; PuIO prepilin peptidase; signal peptide hydrolase; signal

peptide peptidase; signalase; bacterial leader peptidase 1; pilin leader peptidase

**Comments:** The enzyme is found in bacterial membranes and in chloroplast thylakoid membranes. Unaffected

by inhibitors of most serine peptidases, but site-directed mutagenesis implicates a Ser/Lys catalytic dyad in activity [227, 2973]. Hydrolyses a single bond -Ala—Ala- in M13 phage procoat protein, producing free signal peptide and coat protein. Formerly included in EC 3.4.99.36. Eukaryote signal peptidases that may have somewhat different specificity are known from the endoplasmic reticulum membrane [1728] and mitochondrial inner membrane [2120]. Type example of peptidase family S26

**References:** [227, 2120, 2973, 1728, 2972, 388, 1253]

[EC 3.4.21.89 created 1984 as EC 3.4.99.36, transferred 1995 to EC 3.4.21.89]

EC 3.4.21.90

Accepted name: togavirin

**Reaction:** Autocatalytic release of the core protein from the N-terminus of the togavirus structural polyprotein

by hydrolysis of a -Trp—Ser- bond

Other name(s): Sindbis virus protease; Sindbis virus core protein; NsP2 proteinase

Comments: Known from the Sindbis and Semliki forest togaviruses. Once released, the core protein does not re-

tain catalytic activity. Togavirin is the type example of peptidase family S3 and has a similar tertiary

structure to chymotrypsin [2943]

**References:** [1531, 2779, 2943]

[EC 3.4.21.90 created 1995]

EC 3.4.21.91

Accepted name: flavivirin

**Reaction:** Selective hydrolysis of -Xaa-Xaa—Yaa- bonds in which each of the Xaa can be either Arg or Lys and

Yaa can be either Ser or Ala

**Other name(s):** Yellow fever virus (flavivirus) protease; NS2B-3 proteinase

**Comments:** Known from classical flaviviruses (yellow fever, dengue fever). The functional viral peptidase is part

of the NS2B protein. Catalytic His, Asp and Ser residues are arranged as in chymotrypsin, but fla-

vivrin is the type example of peptidase family S7.

**References:** [389, 353, 1690]

### [EC 3.4.21.91 created 1995]

EC 3.4.21.92

**Accepted name:** endopeptidase Clp

**Reaction:** Hydrolysis of proteins to small peptides in the presence of ATP and  $Mg^{2+}$ .  $\alpha$ -Casein is the usual test

-Tyr—Trp bonds also occurs)

Other name(s): endopeptidase Ti; caseinolytic protease; protease Ti; ATP-dependent Clp protease; endopeptidase Ti;

caseinolytic protease; ClpP; Clp protease

**Comments:** An enzyme from bacteria that contains subunits of two types, ClpP, with peptidase activity, and ClpA,

with ATPase activity. The ClpAP complex, which displays ATP-dependent endopeptidase activity, has

the composition ( $ClpP_{14}ClpA_6$ )<sub>2</sub> [1425]. ClpP is the type example of peptidase family S14

**References:** [960, 1850, 1851, 1425]

[EC 3.4.21.92 created 1996]

EC 3.4.21.93

**Accepted name:** proprotein convertase 1

**Reaction:** Release of protein hormones, neuropeptides and renin from their precursors, generally by hydrolysis

of -Lys-Arg- bonds

Other name(s): prohormone convertase 3; neuroendocrine convertase 1; PC1

**Comments:** A Ca<sup>2+</sup>-dependent enzyme, maximally active at about pH 5.5. Substrates include *pro-*

opiomelanocortin, prorenin, proenkephalin, prodynorphin, prosomatostatin and proinsulin. Unlike prohormone convertase 2, does not hydrolyse proluteinizing-hormone-releasing-hormone. Unusually, processing of prodynorphin occurs at a bond in which P2 is Thr. Present in the regulated secretory pathway of neuroendocrine cells, commonly acting co-operatively with prohormone convertase 2. In

peptidase family S8 (subtilisin family)

**References:** [2607, 2693, 2761, 2605, 1312]

[EC 3.4.21.93 created 1996]

EC 3.4.21.94

**Accepted name:** proprotein convertase 2

**Reaction:** Release of protein hormones and neuropeptides from their precursors, generally by hydrolysis of -

Lys-Arg—bonds

Other name(s): neuroendocrine convertase 2; PC2

**Comments:** A Ca<sup>2+</sup>-dependent enzyme, maximally active at about pH 5.5. Specificity is broader than that of

prohormone convertase 1. Substrates include pro-opiomelanocortin, proenkephalin, prodynorphin, proglucagon, proinsulin and proluteinizing-hormone-releasing-hormone. Does not hydrolyse prorenin or prosomatostatin, however. Unusually, processing of prodynorphin occurs at a bond in which P2 is Thr. Present in the regulated secretory pathway of neuroendocrine cells, commonly acting co-

operatively with prohormone convertase 1. In peptidase family S8 (subtilisin family)

**References:** [2607, 2694, 2466, 2605]

[EC 3.4.21.94 created 1996]

EC 3.4.21.95

**Accepted name:** snake venom factor V activator

**Reaction:** Fully activates human clotting factor V by a single cleavage at the Trp-Tyr-Leu-Arg<sup>1545</sup>—Ser-Asn-

Asn-Gly bond. Cattle, but not rabbit, factor V is cleaved, and no other proteins of the clotting system are attacked. Esterase activity is observed on Bz-Arg-OEt and Tos-Arg-OMe, and amidase activity on

Phe-pipecolyl-Arg-NHPhNO<sub>2</sub>

Known from venom of Vipera russelli. Inhibited by di-isopropyl fluorophosphate, unlike the met-**Comments:** 

allopeptidase russellysin (EC 3.4.24.58) that is specific for factor X [1468]. In peptidase family S1

(trypsin family) [2930].

**References:** [1468, 2930]

[EC 3.4.21.95 created 1997]

EC 3.4.21.96

Accepted name: lactocepin

> **Reaction:** Endopeptidase activity with very broad specificity, although some subsite preferences have been

noted, e.g. large hydrophobic residues in the P1 and P4 positions, and Pro in the P2 position [1,2]. Best known for its action on caseins, although it has been shown to hydrolyse hemoglobin and oxi-

dized insulin B chain

CEP; extracellular lactococcal proteinase; lactococcal cell wall-associated proteinase; lactococcal cell Other name(s):

envelope-associated proteinase; lactococcal proteinase; PrtP

**Comments:** Associated with the cell envelope of Lactococcus lactis and attached via a C-terminal membrane an-

> chor sequence. Responsible for the hydrolysis of casein in milk and the provision of peptides essential to cell growth. Important in cheese making and the production of lactic casein, being required for rapid growth to high cell densities with concomitant production of adequate levels of lactic acid. Specificity differences between lactocepins from different starter strains may be partly responsible for

imparting different flavour qualities to cheese [2326]. In peptidase family S8 (subtilisin family)

[3069, 1952, 727, 2326] **References:** 

[EC 3.4.21.96 created 1997]

EC 3.4.21.97

Accepted name: assemblin

> Reaction: Cleaves -Ala—Ser- and -Ala—Ala- bonds in the scaffold protein

**Comments:** Involved in the breakdown of the scaffold protein during the late stages of assembly of the herpes-

virus virion. Inhibited by diisopropyl fluorophosphate. Type example of peptidase family S21. Catalytic residues are His, Ser, His, a combination not known for any other peptidase, and the protein fold also is unique. Known from herpes viruses of several types, cytomegalovirus, Epstein-Barr virus

and human herpesvirus 3

**References:** [415, 537]

[EC 3.4.21.97 created 2000]

EC 3.4.21.98

Accepted name: hepacivirin

> Reaction: Hydrolysis of four peptide bonds in the viral precursor polyprotein, commonly with Asp or Glu in the

> > P6 position, Cys or Thr in P1 and Ser or Ala in P1'

Other name(s): Cpro-2; hepatitis C virus NS3 serine proteinase; NS3-4A serine proteinase complex

Encoded by the genome of the viruses of the hepatitis C group, and contributes to the maturation of **Comments:** 

the precursor polyproteins. The enzyme is greatly activated by binding of the 54-residue NS4A 'cofactor' protein also derived from the viral polyprotein. Type example of peptidase family S29. The

crystallographic structure shows a chymotrypsin-like fold.

**References:** [1450, 2418]

[EC 3.4.21.98 created 2000]

EC 3.4.21.99

Accepted name: spermosin

> Reaction: Hydrolyses arginyl bonds, preferably with Pro in the P2 position

**Comments:** The enzyme from the ascidian (Prochordate) Halocynthia roretzi is localized in the sperm head, and

released during sperm activation. A proline-rich region is involved in binding to the vitelline coat of

the egg. Belongs in peptidase family S1 (trypsin family).

**References:** [2541, 2542, 2539, 2540]

[EC 3.4.21.99 created 2001]

EC 3.4.21.100

Accepted name: sedolisin

> Hydrolysis of the B chain of insulin at -Glu<sup>13</sup>—Ala-, -Leu<sup>15</sup>—Tyr- and -Phe<sup>25</sup>—Tyr-, and an-**Reaction:**

giotensin I at -Tyr<sup>4</sup>—Ile-. A good synthetic substrate is Lys-Pro-Ile-Glu-Phe—Phe(NO<sub>2</sub>)-Arg-Leu.

Pseudomonas sp. pepstatin-insensitive carboxyl proteinase; pseudomonapepsin; pseudomonalisin; Other name(s):

sedolysin

**Comments:** An enzyme secreted by *Pseudomonas* sp. No. 101. Optimum pH is 4. It is distinguished from xan-

thomonapepsin by its insensitivity to EPNP and from scytalidopepsin B by this property and by its unrelated amino-acid sequence. Inhibited by tyrostatin, a peptide aldehyde [2133]. Type example of

peptidase family S53.

[2135, 2133, 3203, 3204] **References:** 

[EC 3.4.21.100 created 1995 as EC 3.4.23.37, transferred 2001 to EC 3.4.21.100, modified 2003]

EC 3.4.21.101

Accepted name: xanthomonalisin Reaction: Cleavage of casein

Other name(s): Xanthomonas aspartic proteinase; xanthomonapepsin; sedolisin-B

**Comments:** Secreted by the bacterium *Xanthomonas* sp. Belongs in peptidase family S53.

**References:** [2134, 3204]

[EC 3.4.21.101 created 1995 as EC 3.4.23.33, transferred 2001 to EC 3.4.21.101, modified 2003]

EC 3.4.21.102

Accepted name: C-terminal processing peptidase

> **Reaction:** The enzyme shows specific recognition of a C-terminal tripeptide, Xaa-Yaa-Zaa, in which Xaa is

> > preferably Ala or Leu, Yaa is preferably Ala or Tyr, and Zaa is preferably Ala, but then cleaves at a variable distance from the C-terminus. A typical cleavage is -Ala-Ala-H-Arg-Ala-Ala-Lys-Glu-Asn-Tyr-Ala-Leu-Ala-Ala. In the plant chloroplast, the enzyme removes the C-terminal extension of the

D1 polypeptide of photosystem II

Other name(s): CtpA gene product (Synechocystis sp.); photosystem II D1 protein processing peptidase; protease Re;

tail-specific protease; Tsp protease

**Comments:** Proteolytic processing of the D1 protein of photosystem II is necessary to allow the light-driven as-

sembly of the tetranuclear manganese cluster, which is responsible for photosynthetic water oxidation. The recognition of the substrate is mediated by a PDZ domain, a small protein module that promotes protein-protein interactions by binding to internal or C-terminal sequences of their partner proteins.

Type example of peptidase family S41.

**References:** [1412, 184, 1679]

[EC 3.4.21.102 created 2001]

EC 3.4.21.103

Accepted name: physarolisin

Milk clotting activity. Preferential cleavage of  $Gly^8$ —Ser in B chain of insulin most rapidly, followed by  $Leu^{11}$ —Val,  $Cys(SO_3H)^{19}$ —Gly and  $Phe^{24}$ —Phe. No action on Ac-Phe-Tyr(I)<sub>2</sub>. **Reaction:** 

Other name(s): Dictyostelium discoideum aspartic proteinase; Dictyostelium discoideum aspartic proteinase E;

Physarum flavicomum aspartic proteinase; Physarum polycephalum acid proteinase; Physarum as-

partic proteinase; physaropepsin

**Comments:** Belongs in peptidase family S53. From the slime mold *Physarum polycephalum*. Is not inhibited by

pepstatin, but is blocked by methyl 2-diazoacetamidohexanoate. Closely similar enzymes are found in

Dictyostelium discoideum and P. flavicomum. Formerly included in EC 3.4.23.6.

**References:** [1113, 2003, 2115, 3204, 2093]

[EC 3.4.21.103 created 1992 as EC 3.4.23.27 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992), transferred 2003 to EC 3.4.21.103]

# EC 3.4.21.104

**Accepted name:** mannan-binding lectin-associated serine protease-2

**Reaction:** Selective cleavage after Arg<sup>223</sup> in complement component C2 (-Ser-Leu-Gly-Arg—Lys-Ile-Gln-Ile)

and after Arg<sup>76</sup> in complement component C4 (-Gly-Leu-Gln-Arg-Hala-Leu-Glu-Ile)

Other name(s): MASP-2; MASP2; MBP-associated serine protease-2; mannose-binding lectin-associated serine

protease-2; p100; mannan-binding lectin-associated serine peptidase 2

Comments: Mannan-binding lectin (MBL) recognizes patterns of neutral carbohydrates, such as mannose and N-

acetylglucosamine, on a wide range of microbial surfaces and is able to initiate activation of the lectin pathway of complement [2897]. This enzyme displays  $C^{\overline{1s}}$ -like esterolytic activity (*cf.* EC 3.4.21.42, complement subcomponent  $C^{\overline{1s}}$ ). It also cleaves C4 and C2 with efficiencies that are relatively higher

than those of EC 3.4.21.42 [2464]. Belongs in peptidase family S1A.

**References:** [1839, 2909, 2464, 45, 1055, 410, 2897]

[EC 3.4.21.104 created 2005]

#### EC 3.4.21.105

**Accepted name:** rhomboid protease

**Reaction:** Cleaves type-1 transmembrane domains using a catalytic dyad composed of serine and histidine that

are contributed by different transmembrane domains

Comments: These endopeptidases are multi-spanning membrane proteins. Their catalytic site is embedded within

the membrane and they cleave type-1 transmembrane domains. A catalytic dyad is involved in proteolysis rather than a catalytic triad, as was thought previously [1643]. They are important for embryo development in *Drosophila melanogaster*. Rhomboid is a key regulator of EGF receptor signalling and is responsible for cleaving Spitz, the main ligand of the *Drosophila* EGF receptor pathway. Belongs in peptidase family S54. Parasite-encoded rhomboid enzymes are also important for invasion of host cells by *Toxoplasma* and the malaria parasite. Rhomboids are widely conserved from bacteria to

archaea to humans [1509, 3020].

**References:** [3023, 306, 1128, 1311, 2658, 3019, 1129, 1875, 1509, 3018, 3022, 3021, 3020, 1643, 3125]

[EC 3.4.21.105 created 2005]

# EC 3.4.21.106

Accepted name: hepsin

**Reaction:** Cleavage after basic amino-acid residues, with Arg strongly preferred to Lys

Comments: This type-II membrane-associated serine peptidase has been implicated in cell growth and develop-

ment [3340, 2949]. The enzyme has been shown to activate blood coagulation factor VII by cleavage of the Arg<sup>152</sup>—Ile<sup>153</sup> peptide bound in BHK cells, thus indicating a possible role in the initiation of blood coagulation [1408]. There is no cleavage after aromatic or aliphatic residues [3340]. The occupancy of the S2 site is an absolute requirement for catalysis and a basic residue at that site is preferred to an aliphatic residue. The nature of the residue at S3 also affects hydrolysis, with Gln being much

more favourable than Ala [3340]. Belongs in peptidase family S1A.

**References:** [3340, 1408, 2949]

EC 3.4.21.107

**Accepted name:** peptidase Do

**Reaction:** Acts on substrates that are at least partially unfolded. The cleavage site P1 residue is normally be-

tween a pair of hydrophobic residues, such as Val—Val

Other name(s): DegP; DegP protease; HtrA; high temperature requirement protease A; HrtA heat shock protein; pro-

tease Do; Do protease

**Comments:** This serine endopeptidase is essential for the clearance of denatured or aggregated proteins from the

inner-membrane and periplasmic space in *Escherichia coli*. Natural substrates of the enzyme include colicin A lysis protein, pilin subunits and MalS from *E. coli* [1335]. The enzyme has weak peptidase activity with casein and other non-native substrates [1335]. The peptidase acts as a chaperone at low temperatures but switches to a peptidase (heat shock protein) at higher temperatures [1710, 1537]. Molecular chaperones and peptidases control the folded state of proteins by recognizing hydrophobic stretches of polypeptide that become exposed by misfolding or unfolding. They then bind these hydrophobic substrates to prevent aggregation or assist in protein refolding. If attempts at refolding fail, then irreversibly damaged proteins are degraded by peptidases such as this enzyme [1537]. Belongs in

peptidase family S1C.

**References:** [1710, 2611, 1335, 2820, 2226, 1537]

[EC 3.4.21.107 created 2006]

EC 3.4.21.108

**Accepted name:** HtrA2 peptidase

**Reaction:** Cleavage of non-polar aliphatic amino-acids at the P1 position, with a preference for Val, Ile and Met.

At the P2 and P3 positions, Arg is selected most strongly with a secondary preference for other hy-

drophilic residues

Other name(s): high temperature requirement protein A2; HtrA2; Omi stress-regulated endoprotease; serine pro-

teinase OMI; HtrA2 protease; OMI/HtrA2 protease; HtrA2/Omi; Omi/HtrA2

Comments: This enzyme is upregulated in mammalian cells in response to stress induced by both heat shock and

tunicamycin treatment [967]. It can induce apoptosis in a caspase-independent manner through its peptidase activity and in a caspase-dependent manner by disrupting the interaction between caspase

and the inhibitor of apoptosis (IAP) [1824]. Belongs in peptidase family S1C.

**References:** [2746, 2538, 1824, 967, 1671]

[EC 3.4.21.108 created 2006]

EC 3.4.21.109

Accepted name: matriptase

**Reaction:** Cleaves various synthetic substrates with Arg or Lys at the P1 position and prefers small side-chain

amino acids, such as Ala and Gly, at the P2 position

Other name(s): serine protease 14; membrane-type serine protease 1; MT-SP1; prostamin; serine protease TADG-

15; tumor-associated differentially-expressed gene 15 protein; ST14; breast cancer 80 kDa protease;

epithin; serine endopeptidase SNC19

**Comments:** This trypsin-like integral-membrane serine peptidase has been implicated in breast cancer invasion

and metastasis [1631, 1691]. The enzyme can activate hepatocyte growth factor/scattering factor (HGF/SF) by cleavage of the two-chain form at an Arg residue to give active  $\alpha$ - and  $\beta$ -HGF, but It does not activate plasminogen, which shares high homology with HGF [1631]. The enzyme can also activate urokinase plasminogen activator (uPA), which initiates the matrix-degrading peptidase cas-

cade [1631, 1691]. Belongs in peptidase family S1A.

**References:** [1631, 1691]

[EC 3.4.21.109 created 2006]

EC 3.4.21.110

**Accepted name:** C5a peptidase

**Reaction:** The primary cleavage site is at His<sup>67</sup>+Lys<sup>68</sup> in human C5a with a minor secondary cleavage site at

Ala<sup>58</sup>—Ser<sup>59</sup>

Other name(s): streptococcal C5a peptidase; ScpA; ScpB; SCPA

**Comments:** This enzyme is a surface-associated subtilisin-like serine peptidase with very specific substrate speci-

ficity. Virulent strains of streptococci, including *Streptococcus pyogenes*, can evade human detection and phagocytosis by destroying the complement chemotaxin C5a. Cleavage of human C5a by this enzyme reduces the ability of C5a to bind receptors on the surface of polymorphonuclear neutrophil leukocytes (PMNLs) and thereby abolishes its chemotactic properties [3167, 52]. Belongs in pepti-

dase family S8A.

**References:** [3167, 254, 461, 52, 2753, 2901]

[EC 3.4.21.110 created 2006]

EC 3.4.21.111

Accepted name: aqualysin 1

**Reaction:** Exhibits low specificity towards esters of amino acids with small hydrophobic or aromatic residues at

the P1 position

Other name(s): caldolysin

**Comments:** This enzyme from the extreme thermophile, *Thermus aquaticus*, is an alkaline serine peptidase. It has

three subsites, S1, S2, and S3, in the substrate binding site. The preferred amino acids at the S1 site are Ala and Phe, at the S2 site are Ala and norleucine and at the S3 site are Phe and Ile [2877]. These specificities are similar to those of EC 3.4.21.64 (peptidase K) and EC 3.4.21.62 (subtilisin BPN') [2877]. The enzyme displays broad specificity for cleavage of insulin B-chain and hydrolyses elastin substrates such as succinyl-(Ala) $_n$ - $_p$ -nitroanilide (n = 1,2,3) and some peptide esters [1843, 2877].

Belongs in peptidase family S8A.

**References:** [1843, 2876, 2877]

[EC 3.4.21.111 created 2006]

EC 3.4.21.112

**Accepted name:** site-1 protease

Reaction: Processes precursors containing basic and hydrophobic/aliphatic residues at P4 and P2, respectively,

with a relatively relaxed acceptance of amino acids at P1 and P3

Other name(s): mammalian subtilisin/kexin isozyme 1; membrane-bound transcription factor site-1 protease; propro-

tein convertase SKI-1; proprotein convertase SKI-1/S1PPS1; S1P endopeptidase; S1P protease; site-1 peptidase; site-1 protease; SKI-1; SREBP proteinase; SREBP S1 protease; SREBP-1 proteinase; SREBP-2 proteinase; sterol regulatory element-binding protein proteinase; sterol regulatory element-binding protein site 1 protease; sterol-regulated luminal protease; subtilase SKI-1; subtilase SKI-1

1/S1P; subtilisin/kexin-isozyme 1

**Comments:** Cleaves sterol regulatory element-binding proteins (SREBPs) and thereby initiates a process by which

the active fragments of the SREBPs translocate to the nucleus and activate genes controlling the synthesis and uptake of cholesterol and unsaturated fatty acids into the bloodstream [718]. The enzyme also processes pro-brain-derived neurotrophic factor and undergoes autocatalytic activation in the endoplasmic reticulum through sequential cleavages [1647]. The enzyme can also process the unfolded protein response stress factor ATF6 at an Arg-His-Lys-Lys—site [3281, 2606], and the envelope glycoprotein of the highly infectious Lassa virus [1647, 2606] and Crimean Congo hemorrhagic fever

virus at Arg-Arg-Lys-Lys+ [3067, 2606]. Belongs in peptidase family S8A.

**References:** [718, 422, 2951, 3281, 1647, 167, 3067, 2606]

[EC 3.4.21.112 created 2006]

EC 3.4.21.113

**Accepted name:** pestivirus NS3 polyprotein peptidase

**Reaction:** Leu is conserved at position P1 for all four cleavage sites. Alanine is found at position P1' of the

NS4A-NS4B cleavage site, whereas serine is found at position P1' of the NS3-NS4A, NS4B-NS5A

and NS5A-NS5B cleavage sites

Other name(s): border disease virus NS3 endopeptidase; BDV NS3 endopeptidase; bovine viral diarrhea virus NS3

endopeptidase; BVDV NS3 endopeptidase; classical swine fever virus NS3 endopeptidase; CSFV

NS3 endopeptidase; p80

**Comments:** The polyprotein of noncytopathogenic pestiviruses is cleaved co- and post-translationally into at least

11 proteins (N<sup>pro</sup>, C, E<sup>rns</sup>, E1, E2, p7, NS2-3, NS4A, NS4B, NS5A, and NS5B) [2893]. The genomes of cytopathogenic pestivirus strains express at least one additional protein, called NS3 (p80) [2893]. This enzyme, which resides in the N-terminal region of NS3 (nonstructural protein 3), is essential for generation of its own C-terminus and for processing of the downstream cleavage sites, leading to the release of the pestivirus nonstructural proteins NS4A, NS4B, NS5A and NS5B [3200, 2893]. Belongs

in peptidase family S31.

**References:** [3200, 2893, 3232, 2894]

[EC 3.4.21.113 created 2006]

EC 3.4.21.114

**Accepted name:** equine arterivirus serine peptidase

**Reaction:** Cleavage of (Glu/Gln)—(Gly/Ser/Ala) in arterivirus replicase translation products ORF1a and

ORF1ab

**Comments:** In the equine arterivirus (EAV), the replicase gene is translated into open reading frame 1a (ORF1a)

and ORF1ab polyproteins. This enzyme is the main viral proteinase and processes five cleavage sites in the ORF1a protein and three in the ORF1b-encoded part of the ORF1ab protein to yield nonstructural proteins (nsp5-nsp12) [162]. It combines the catalytic system of a chymotrypsin-like serine peptidase (His-Asp-Ser catalytic triad) with the substrate specificity of a 3C-like serine peptidase (Glu or Gln) at the P1 position and a small amino-acid residue (Gly, Ser or Ala) at the P1' position [2704]. Cleavage of ORF1ab by this enzyme is essential for viral replication [3046]. Belongs in peptidase

family S32.

**References:** [2704, 3046, 162]

[EC 3.4.21.114 created 2006]

EC 3.4.21.115

**Accepted name:** infectious pancreatic necrosis birnavirus Vp4 peptidase

**Reaction:** Cleaves the (Ser/Thr)-Xaa-Ala—(Ser/Ala)-Gly motif in the polyprotein NH<sub>2</sub>-pVP2-VP4-VP3-COOH

of infectious pancreatic necrosis virus at the pVP2-VP4 and VP4-VP3 junctions

Other name(s): infectious pancreatic necrosis virus protease; IPNV Vp4 protease; IPNV Vp4 peptidase; NS protease;

NS-associated protease; Vp4 protease

Comments: Infectious pancreatic necrosis virus (IPNV) is a birnavirus that causes an acute, contagious disease in

young salmonid fish [2266]. As with most viruses that infect eukaryotic cells, the proteolytic processing of viral precursor proteins is a crucial step in the life cycle of this virus [2266]. pVP2 is converted into VP2 by cleavage near the carboxy end of pVP2. This cleavage is most likely due to host-cell proteases rather than VP4 [2266, 622]. Differs from most serine peptidases in not having the catalytic

triad Ser-His-Asp [2266]. Belongs in peptidase family S50.

**References:** [1802, 2266, 622]

[EC 3.4.21.115 created 2006]

EC 3.4.21.116

**Accepted name:** SpoIVB peptidase

**Reaction:** Self-cleaves Val<sup>52</sup>—Asn<sup>53</sup>, Ala<sup>62</sup>—Phe<sup>63</sup> and Val<sup>74</sup>—Thr<sup>75</sup> at the N-terminus of SpoIVB

**Other name(s):** sporulation factor IV B protease

Comments: This enzyme plays a central role in a regulatory checkpoint (the  $\sigma^{K}$  checkpoint), which coordinates

gene expression during the later stages of spore formation in *Bacillus subtilis* [3092, 1156]. The enzyme activates proteolytic processing of a sporulation-specific sigma factor, pro- $\sigma^K$ , to its mature and active form,  $\sigma^K$ , by self-cleavage [3092, 1156]. The enzyme is also subject to secondary proteolysis, which presumably inactivates SpoIVB [1156]. The enzyme is also essential for the formation of heat-

resistant spores. Belongs in peptidase family S55.

**References:** [3092, 1155, 1156, 635]

[EC 3.4.21.116 created 2006]

EC 3.4.21.117

**Accepted name:** stratum corneum chymotryptic enzyme

**Reaction:** Cleavage of proteins with aromatic side chains in the P1 position

Other name(s): kallikrein 7; SCCE; KLK7; PRSS6; hK7

**Comments:** This enzyme has wide substrate specificity, being able to degrade heat-denatured bovine casein and

the  $\alpha$ -chain of native human fibrinogen. It cleaves the B chain of bovine insulin at Leu<sup>6</sup>—Cya<sup>7</sup>, Tyr<sup>16</sup>—Leu<sup>17</sup>, Phe<sup>25</sup>—Tyr<sup>26</sup> and Tyr<sup>26</sup>—Thr<sup>27</sup> [2687]. It is thought to play a role in the desquamation (skin-shedding) of the outer layer of skin, the stratum corneum, by degrading intercellular co-

hesive structures [2687, 680]. Belongs in peptidase family S1A.

**References:** [2687, 680, 1046, 3307, 3055]

[EC 3.4.21.117 created 2006]

EC 3.4.21.118

Accepted name: kallikrein 8

Reaction: Cleavage of amide substrates following the basic amino acids Arg or Lys at the P1 position, with a

preference for Arg over Lys

Other name(s): KLK8; PRSS19; human kallikrein 8; hK8; mK8; ovasin; tumor-associated differentially expressed

gene 14; TADG-14; NP; neuropsin

**Comments:** The enzyme is activated by removal of an N-terminal prepropertide [2633, 1466]. The highest ami-

dolytic activity is observed using Boc-Val-Pro-Arg+7-amido-4-methylcoumarin, which is a substrate of  $\alpha$ -thrombin [2633, 1466]. Substrates lacking basic amino acids in the P1 position are not cleaved [1466]. The enzyme degrades casein, fibronectin, gelatin, collagen type IV, fibrinogen, and high-molecular-mass kininogen [2362] and is associated with diseases such as ovarian cancer and

Alzheimer's disease [1466]. Belongs in peptidase family S1A.

**References:** [421, 2633, 2362, 1466]

[EC 3.4.21.118 created 2006]

EC 3.4.21.119

Accepted name: kallikrein 13

Reaction: Hydrolyses mouse Ren2 protein (a species of prorenin present in the submandibular gland) on the

carboxy side of the arginine residue at the Lys-Arg—pair in the N-terminus, to yield mature renin

Other name(s): KLK13; kallikrein mK13; mGK-13; mK13; mKLK13; prorenin converting enzyme 1; PRECE-1;

prorenin-converting enzyme; PRECE; proteinase P

**Comments:** The enzyme is specific for prorenin from the mouse submandibular gland, as prorenin from the mouse

kidney (Ren1) and human prorenin are not substrates [2052]. Site-directed mutagenesis studies have shown that the enzyme will also cleave prorenin when Lys-Arg is replaced by Arg-Arg or Gln-Arg but the rate of reaction is much slower when Lys-Lys is used. This enzyme is also able to process prointerleukin-1β (pro-IL-1β) in mouse submandibular gland to form IL-1β [3261]. Belongs in peptidase

family S1A.

**References:** [2052, 1455, 1439, 3261]

### [EC 3.4.21.119 created 2006]

EC 3.4.21.120

Accepted name: oviductin

**Reaction:** Preferential cleavage at Gly-Ser-Arg<sup>373</sup>— of glycoprotein gp43 in *Xenopus laevis* coelemic egg enve-

lope to yield gp41

Other name(s): oviductal protease

Comments: The egg envelope of the South African clawed frog (Xenopus laevis) is modified during transit of the

egg through the pars rectus oviduct, changing the egg envelope from an unfertilizable form to a fertilizable form. This process involves the conversion of glycoprotein gp43 to gp41 (ZPC) by the pars recta protease oviductin. It is thought that the enzymically active protease molecule comprises the N-terminal protease domain coupled to two C-terminal CUB domains, which are related to the mammalian spermadhesin molecules implicated in mediating sperm-envelope interactions [1703]. The enzyme is also found in the Japanese toad (*Bufo japonicus*) [1153]. Belongs in peptidase family S1.

**References:** [1053, 1703, 1153]

[EC 3.4.21.120 created 2007]

EC 3.4.21.121

Accepted name: Lys-Lys/Arg-Xaa endopeptidase

**Reaction:** Cleavage of -Lys-Lys— and -Lys-Arg— bonds.

Other name(s): ASP (Aeromonas sobria)-type peptidase; Aeromonas extracellular serine protease

**Comments:** The enzyme is a serine peptidase, which has been shown to cleave prothrombin and prekallikrein. It

hydrolyses the complement component C5 releasing complement component C5a.

**References:** [1489, 2100, 1488, 1241, 2099]

[EC 3.4.21.121 created 2013]

# EC 3.4.22 Cysteine endopeptidases

EC 3.4.22.1

Accepted name: cathepsin B

**Reaction:** Hydrolysis of proteins with broad specificity for peptide bonds. Preferentially cleaves -Arg-Arg—

bonds in small molecule substrates (thus differing from cathepsin L). In addition to being an endopep-

tidase, shows peptidyl-dipeptidase activity, liberating C-terminal dipeptides

**Other name(s):** cathepsin B1 (obsolete); cathepsin II

**Comments:** An intracellular (lysosomal) enzyme in peptidase family C1 (papain family)

**References:** [258, 159, 2304, 158, 1464]

[EC 3.4.22.1 created 1972]

EC 3.4.22.2

Accepted name: papain

**Reaction:** Hydrolysis of proteins with broad specificity for peptide bonds, but preference for an amino acid bear-

ing a large hydrophobic side chain at the P2 position. Does not accept Val in P1'

Other name(s): papayotin; summetrin; velardon; papaine; Papaya peptidase I

**Comments:** Type example of peptidase family C1 from latex of the papaya (*Carica papaya*) fruit. Inhibited by

compound E-64 and proteins of the cystatin family.

**References:** [1375, 1884]

[EC 3.4.22.2 created 1961 as EC 3.4.4.10, transferred 1972 to EC 3.4.22.2, modified 1976, modified 2000]

# EC 3.4.22.3

Accepted name: ficain

**Reaction:** Similar to that of papain

Other name(s): ficin; debricin; higueroxyl delabarre

**Comments:** The major proteolytic component of the latex of fig, *Ficus glabrata*. Cysteine endopeptidases with

similar properties are present in other members of the large genus Ficus. In peptidase family C1 (pa-

pain family).

**References:** [1684, 297]

[EC 3.4.22.3 created 1961 as EC 3.4.4.12, transferred 1972 to EC 3.4.22.3]

[3.4.22.4 Transferred entry. bromelain (stem). Now EC 3.4.22.32 (stem bromelain) and EC 3.4.22.33 (fruit bromelain)]

[EC 3.4.22.4 created 1972, deleted 1992 [EC 3.4.22.5 created 1972, incorporated 1978]]

[3.4.22.5] Transferred entry. bromelain (juice). Now EC 3.4.22.32 (stem bromelain) and EC 3.4.22.33 (fruit bromelain)]

[EC 3.4.22.5 created 1972, deleted 1978]

## EC 3.4.22.6

Accepted name: chymopapain

**Reaction:** Similar to that of papain

Other name(s): chymopapain A; chymopapain B; chymopapain S

Comments: The major endopeptidase of papaya (Carica papaya) latex. It has multiple chromatographic forms. In

peptidase family C1 (papain family).

**References:** [297, 1293, 339]

[EC 3.4.22.6 created 1961 as EC 3.4.4.11, transferred 1972 to EC 3.4.22.6]

# EC 3.4.22.7

Accepted name: asclepain

**Reaction:** Similar to that of papain

**Comments:** From the latex of milkweed, *Asclepias syriaca*. It has multiple forms, and is in peptidase family C1

(papain family)

**References:** [295]

[EC 3.4.22.7 created 1972]

## EC 3.4.22.8

**Accepted name:** clostripain

**Reaction:** Preferential cleavage: Arg—, including Arg—Pro, but not Lys-

Other name(s): clostridiopeptidase B; clostridium histolyticum proteinase B;  $\alpha$ -clostridipain; clostridiopeptidase Comments: From the bacterium *Clostridium histolyticum*. It requires Ca<sup>2+</sup> ions and is inhibited by EDTA. Type

example of peptidase family C11.

**References:** [1931, 922, 923]

[EC 3.4.22.8 created 1961 as EC 3.4.4.20, transferred 1972 to EC 3.4.22.8]

[3.4.22.9 Transferred entry. yeast proteinase B. Now EC 3.4.21.48, cerevisin]

[EC 3.4.22.9 created 1972, deleted 1981]

# EC 3.4.22.10

Accepted name: streptopain

**Reaction:** Preferential cleavage with hydrophobic residues at P2, P1 and P1'

Other name(s): Streptococcus peptidase A; streptococcal cysteine proteinase; Streptococcus protease

**Comments:** From the bacterium, group A *Streptococcus*. Formed from the proenzyme by limited proteolysis.

Type example of peptidase family C10.

**References:** [693, 1726, 2830, 1730]

[EC 3.4.22.10 created 1961 as EC 3.4.4.18, transferred 1972 to EC 3.4.22.10]

[3.4.22.11 Transferred entry. insulinase. Now EC 3.4.24.56, insulysin]

[EC 3.4.22.11 created 1976, deleted 1978 [transferred to EC 3.4.99.45, deleted 1993]]

[3.4.22.12 Transferred entry, \gamma-glutamyl hydrolase, Now EC 3.4.19.9, \gamma-glutamyl hydrolase]

[EC 3.4.22.12 created 1978, deleted 1992]

[3.4.22.13 Deleted entry. staphylococcal cysteine proteinase]

[EC 3.4.22.13 created 1978, modified 1981, deleted 1992]

### EC 3.4.22.14

Accepted name: actinidain

**Reaction:** Similar to that of papain

Other name(s): actinidin; Actinidia anionic protease; proteinase A<sub>2</sub> of Actinidia chinensis

**Comments:** From the kiwi fruit or Chinese gooseberry (*Actinidia chinensis*). In peptidase family C1 (papain fam-

ily)

**References:** [125, 1375, 126]

[EC 3.4.22.14 created 1978]

### EC 3.4.22.15

**Accepted name:** cathepsin L

**Reaction:** Similar to that of papain. As compared to cathepsin B, cathepsin L exhibits higher activity towards

protein substrates, but has little activity on Z-Arg-Arg-NHMec, and no peptidyl-dipeptidase activity

**Other name(s):** Aldrichina grahami cysteine proteinase

Comments: A lysosomal enzyme in peptidase family C1 (papain family) that is readily inhibited by the dia-

zomethane inhibitor Z-Phe-Phe-CHN<sub>2</sub> or the epoxide inhibitor E-64

**References:** [159, 158, 1341, 1464]

[EC 3.4.22.15 created 1978 (EC 3.4.99.19 created 1972, incorporated 1981)]

## EC 3.4.22.16

Accepted name: cathepsin H

**Reaction:** Hydrolysis of proteins, acting as an aminopeptidase (notably, cleaving Arg—bonds) as well as an

endopeptidase

Other name(s): cathepsin B3; benzoylarginine-naphthylamide (BANA) hydrolase (obsolete); cathepsin Ba, aleurain;

N-benzoylarginine- $\beta$ -naphthylamide hydrolase

**Comments:** Present in lysosomes of mammalian cells. In peptidase family C1 (papain family)

**References:** [159, 300, 830]

[EC 3.4.22.16 created 1981, modified 1989]

[3.4.22.17 Transferred entry. calpain. Now EC 3.4.22.53, calpain-2]

[EC 3.4.22.17 created 1981 [EC 3.4.24.5 created 1978, part incorporated 1989], deleted 2003]

[3.4.22.18 Transferred entry. prolyl endopeptidase (thiol-dependent). Now EC 3.4.21.26, prolyl oligopeptidase]

[EC 3.4.22.18 created 1981, deleted 1992]

[3.4.22.19 Transferred entry. endo-oligopeptidase. Now EC 3.4.24.15, thimet oligopeptidase]

[EC 3.4.22.19 created 1989, deleted 1992]

[3.4.22.20 Deleted entry. dinorphin-converting enzyme]

[EC 3.4.22.20 created 1989, deleted 1992]

[3.4.22.21 Transferred entry, yeast cysteine proteinase E. Now EC 3.4.25.1, proteasome endopeptidase complex]

[EC 3.4.22.21 created 1989, deleted 1992]

[3.4.22.22 Transferred entry. yeast cysteine proteinase D. Now EC 3.4.24.37, saccharolysin]

[EC 3.4.22.22 created 1989, deleted 1992]

[3.4.22.23 Transferred entry. yeast cysteine proteinase F. Now EC 3.4.21.61, kexin]

[EC 3.4.22.23 created 1989, deleted 1992]

EC 3.4.22.24

Accepted name: cathepsin T

**Reaction:** Interconversion of the three forms of tyrosine aminotransferase, EC 2.6.1.5

**Comments:** Degrades azocasein and denatured hemoglobin; the only native protein on which it has been shown to

act is tyrosine aminotransferase

**References:** [941, 940, 2288]

[EC 3.4.22.24 created 1990]

EC 3.4.22.25

Accepted name: glycyl endopeptidase

**Reaction:** Preferential cleavage: Gly—, in proteins and small molecule substrates

Other name(s): papaya peptidase B; papaya proteinase IV; glycine-specific proteinase; chymopapain; Papaya pro-

teinase 4; PPIV; chymopapain M

Comments: From the papaya plant, Carica papaya. Not inhibited by chicken cystatin, unlike most other homo-

logues of papain, but in peptidase family C1 (papain family)

**References:** [2301, 340, 2430, 342, 341]

[EC 3.4.22.25 created 1992]

EC 3.4.22.26

Accepted name: cancer procoagulant

**Reaction:** Specific cleavage of Arg—Ile bond in Factor X to form Factor Xa

**Comments:** Apparently produced only by malignant and fetal cells

**References:** [733, 734]

[EC 3.4.22.26 created 1992]

EC 3.4.22.27

Accepted name: cathepsin S

Reaction: Similar to cathepsin L, but with much less activity on Z-Phe-Arg—NHMec, and more activity on the

Z-Val-Val-Arg — compound

**Comments:** A lysosomal cysteine endopeptidase that is unusual amongst such enzymes for its stability to neutral

pH. In peptidase family C1 (papain family)

**References:** [3003, 304, 1463]

#### [EC 3.4.22.27 created 1992]

EC 3.4.22.28

Accepted name: picornain 3C

**Reaction:** Selective cleavage of Gln—Gly bond in the poliovirus polyprotein. In other picornavirus reactions

Glu may be substituted for Gln, and Ser or Thr for Gly

Other name(s): picornavirus endopeptidase 3C; poliovirus protease 3C; rhinovirus protease 3C; foot-and-mouth pro-

tease 3C; poliovirus proteinase 3C; rhinovirus proteinase 3C; coxsackievirus 3C proteinase; foot-and-mouth-disease virus proteinase 3C; 3C proteinase; 3C proteinase; cysteine proteinase 3C; hepatitis A

virus 3C proteinase; protease 3C; tomato ringspot nepovirus 3C-related protease

Comments: From entero-, rhino-, aphto- and cardioviruses. Larger than the homologous virus picornain 2A. Type

example of peptidase family C3

**References:** [1278, 179, 1531, 2081]

[EC 3.4.22.28 created 1992]

EC 3.4.22.29

Accepted name: picornain 2A

**Reaction:** Selective cleavage of Tyr—Gly bond in picornavirus polyprotein

Other name(s): picornavirus endopeptidase 2A; poliovirus protease 2A; rhinovirus protease 2A; 2A protease; 2A pro-

teinase; protease 2A; proteinase 2Apro; picornaviral 2A proteinase; Y-G proteinase 2A; poliovirus

proteinase 2A; poliovirus protease 2Apro; picornaviral 2A proteinase

Comments: From entero-, rhino-, aphto- and cardioviruses. Smaller than the homologous picornain 3C, which is

also in peptidase family C3 (picornain 3C family)

**References:** [179, 1508, 1531]

[EC 3.4.22.29 created 1992]

EC 3.4.22.30

Accepted name: caricain

Reaction: Hydrolysis of proteins with broad specificity for peptide bonds, similar to those of papain and chy-

mopapain

Other name(s): papaya peptidase A; papaya peptidase II; papaya proteinase ; papaya proteinase III; papaya proteinase

3; proteinase ω; papaya proteinase A; chymopapain S; Pp

**Comments:** From papaya plant, *Carica papaya*. In peptidase family C1 (papain family)

**References:** [2552, 2441, 2302, 296, 3350, 658]

[EC 3.4.22.30 created 1992]

EC 3.4.22.31

Accepted name: ananain

Reaction: Hydrolysis of proteins with broad specificity for peptide bonds. Best reported small molecule sub-

strate Bz-Phe-Val-Arg—NHMec, but broader specificity than fruit bromelain

**Other name(s):** stem bromelain; fruit bromelain

**Comments:** From stem of pineapple plant, *Ananas comosus*. Differs from stem and fruit bromelains in being in-

hibited by chicken cystatin. In peptidase family C1 (papain family)

**References:** [2468, 2469]

[EC 3.4.22.31 created 1992]

EC 3.4.22.32

Accepted name: stem bromelain

**Reaction:** Broad specificity for cleavage of proteins, but strong preference for Z-Arg-Arg—NHMec amongst

small molecule substrates

Other name(s): bromelain; pineapple stem bromelain

**Comments:** The most abundant of the cysteine endopeptidases of the stem of the pineapple plant, *Ananas como-*

sus. Distinct from the bromelain found in the pineapple fruit (EC 3.4.22.33). Scarcely inhibited by chicken cystatin and also very slowly inactivated by E-64. In peptidase family C1 (papain family).

**References:** [297, 2468, 2431, 2469]

[EC 3.4.22.32 created 1965 as EC 3.4.4.24, transferred 1972 to EC 3.4.22.4, part transferred 1992 to EC 3.4.22.32]

#### EC 3.4.22.33

Accepted name: fruit bromelain

**Reaction:** Hydrolysis of proteins with broad specificity for peptide bonds. Bz-Phe-Val-Arg—NHMec is a good

synthetic substrate, but there is no action on Z-Arg-Arg-NHMec (c.f. stem bromelain)

Other name(s): juice bromelain; ananase; bromelase; bromelin; extranase; juice bromelain; pinase; pineapple enzyme;

traumanase; fruit bromelain FA2

**Comments:** From the pineapple plant, *Ananas comosus*. Scarcely inhibited by chicken cystatin. Another cys-

teine endopeptidase, with similar action on small molecule substrates, pinguinain, is obtained from the related plant, *Bromelia pinguin*, but pinguinain differs from fruit bromelain in being inhibited by

chicken cystatin [2469].

**References:** [2535, 3241, 2211, 2469]

[EC 3.4.22.33 created 1965 as EC 3.4.4.24, transferred 1972 to EC 3.4.22.4, part transferred 1992 to EC 3.4.22.33]

### EC 3.4.22.34

Accepted name: legumain

**Reaction:** Hydrolysis of proteins and small molecule substrates at -Asn—Xaa- bonds

Other name(s): asparaginyl endopeptidase; citvac; proteinase B (ambiguous); hemoglobinase (ambiguous); PRSC1

 $gene\ product\ (Homo\ sapiens);\ vicilin\ peptidohydrolase;\ bean\ endopeptidase;\ vicilin\ peptidohydrolase$ 

Comments: Best known from legume seeds, the trematode Schistosoma mansoni and mammalian lysosomes. Not

inhibited by compound E-64. Type example of peptidase family C13

**References:** [1049, 531, 412]

[EC 3.4.22.34 created 1992, modified 2000]

### EC 3.4.22.35

**Accepted name:** histolysain

**Reaction:** Hydrolysis of proteins, including basement membrane collagen and azocasein. Preferential cleavage:

Arg-Arg—in small molecule substrates including Z-Arg-Arg—NHMec

Other name(s): histolysin; histolysin; Entamoeba histolytica cysteine proteinase; amebapain; Entamoeba histolytica

cysteine protease; Entamoeba histolytica neutral thiol proteinase

**Comments:** From the protozoan, *Entamoeba histolytica*. In peptidase family C1 (papain family)

**References:** [1757, 1748]

[EC 3.4.22.35 created 1992]

### EC 3.4.22.36

**Accepted name:** caspase-1

Reaction: Strict requirement for an Asp residue at position P1 and has a preferred cleavage sequence of Tyr-Val-

Ala-Asp+

**Other name(s):** interleukin 1β-converting enzyme; protease VII; protease A; interleukin 1β precursor proteinase; in-

terleukin 1 converting enzyme; interleukin 1 $\beta$ -converting endopeptidase; interleukin-1 $\beta$  convertase; interleukin-1 $\beta$  converting enzyme; interleukin-1 $\beta$  processe; prointerleukin 1 $\beta$  protease;

precursor interleukin- $1\beta$  converting enzyme; pro-interleukin  $1\beta$  proteinase; ICE

**Comments:** From mammalian monocytes. This enzyme is part of the family of inflammatory caspases, which

also includes caspase-4 (EC 3.4.22.57) and caspase-5 (EC 3.4.22.58) in humans and caspase-11 (EC 3.4.22.64), caspase-12, caspase-13 and caspase-14 in mice. Contains a caspase-recruitment domain (CARD) in its N-terminal prodomain, which plays a role in procaspase activation [1823, 393]. Cleaves pro-interleukin-1 $\beta$  (pro-IL-1 $\beta$ ) to form mature IL-1 $\beta$ , a potent mediator of inflammation. Also activates the proinflammatory cytokine, IL-18, which is also known as interferon- $\gamma$ -inducing factor [1823]. Inhibited by Ac-Tyr-Val-Ala-Asp-CHO. Caspase-11 plays a critical role in the activation of caspase-1 in mice, whereas caspase-4 enhances its activation in humans [393]. Belongs in pepti-

dase family C14.

**References:** [1191, 2919, 2918, 38, 1809, 1823, 393]

[EC 3.4.22.36 created 1993, modified 1997, modified 2007]

EC 3.4.22.37

Accepted name: gingipain R

**Reaction:** Hydrolysis of proteins and small molecule substrates, with a preference for Arg in P1

**Other name(s):** Arg-gingipain; gingipain-1; argingipain; Arg-gingivain-55 proteinase; Arg-gingivain-70 proteinase;

Arg-gingivain-75 proteinase; arginine-specific cysteine protease; arginine-specific gingipain; arginine-

specific gingivain; RGP-1; RGP

Comments: A secreted endopeptidase from the bacterium Porphyromonas gingivalis. Strongly activated by

glycine [420], and stabilized by Ca<sup>2+</sup>. Precursor molecule contains a hemagglutinin domain

[1465, 2250]. Misleadingly described in some literature as "trypsin-like", being a cysteine peptidase,

type example of family C25

**References:** [420, 1465, 2250]

[EC 3.4.22.37 created 1996]

EC 3.4.22.38

**Accepted name:** cathepsin K

Reaction: Broad proteolytic activity. With small-molecule substrates and inhibitors, the major determinant of

specificity is P2, which is preferably Leu, Met > Phe, and not Arg

Other name(s): cathepsin O and cathepsin X (both misleading, having been used for other enzymes); cathepsin O<sub>2</sub>

**Comments:** Prominently expressed in mammalian osteoclasts, and believed to play a role in bone resorption. In

peptidase family C1 (papain family)

**References:** [1248, 268, 301, 3331, 1872]

[EC 3.4.22.38 created 1997]

EC 3.4.22.39

Accepted name: adenain

**Reaction:** Cleaves proteins of the adenovirus and its host cell at two consensus sites: -Yaa-Xaa-Gly-Gly—Xaa-

and -Yaa-Xaa-Gly-Xaa—Gly- (in which Yaa is Met, Ile or Leu, and Xaa is any amino acid)

**Comments:** A cysteine endopeptidase from adenoviruses, the type example of peptidase family C5, with a protein

fold unlike that known for any other peptidase [617]. Activity is greatly stimulated by the binding to the enzyme of an 11-residue peptide from the adenovirus capsid protein pre-VI at a site separate from

the active site [3143]

**References:** [3143, 617, 3142]

[EC 3.4.22.39 created 2000]

EC 3.4.22.40

**Accepted name:** bleomycin hydrolase

**Reaction:** Inactivates bleomycin B2 (a cytotoxic glycometallopeptide) by hydrolysis of a carboxyamide bond of

β-aminoalanine, but also shows general aminopeptidase activity. The specificity varies somewhat with

source, but amino acid arylamides of Met, Leu and Ala are preferred [1]

Other name(s): aminopeptidase C (Lactococcus lactis) [4]

**Comments:** The molecule is a homohexamer in which the monomers have a papain-like tertiary structure (in pep-

tidase family C1). The active sites are on the walls of a central channel through the molecule, and access of substrate molecules to them is obstructed by this and by the C-terminus of each polypeptide chain [3333]. Bleomycin can scarcely be the natural substrate, and there are reports of limited endopeptidase activity. Known from bacteria as well as eukaryotic organisms. Hydrolase H from chicken muscle has many similarities to bleomycin hydrolase, but hydrolyses Ph-CO-Arg-2-

naphthylamine as well as aminopeptidase substrates [13].

**References:** [303, 13, 3333, 1929]

[EC 3.4.22.40 created 2000]

EC 3.4.22.41

**Accepted name:** cathepsin F

Reaction: The recombinant enzyme cleaves synthetic substrates with Phe and Leu (better than Val) in P2, with

high specificity constant  $(k_{cat}/K_m)$  comparable to that of cathepsin L

**Comments:** Cathepsin F is a lysosomal cysteine endopeptidase of family C1 (papain family), most active at pH

5.9. The enzyme is unstable at neutral pH values and is inhibited by compound E-64. Cathepsin F is

expressed in most tissues of human, mouse and rat. Human gene locus: 11q13.1-13.3

**References:** [2525, 2028, 3166, 3104]

[EC 3.4.22.41 created 2000]

EC 3.4.22.42

**Accepted name:** cathepsin O

Reaction: The recombinant human enzyme hydrolyses synthetic endopeptidase substrates including Z-Phe-Arg-

NHMec and Z-Arg-Arg-NHMec

**Comments:** Cathepsin O is a lysosomal cysteine peptidase of family C1 (papain family). The recombinant human

enzyme is catalytically active at pH 6.0 and is inhibited by compound E-64. Cathepsin O is ubiqui-

tously expressed in human tissues and the human gene locus is 4q31-32

**References:** [2523, 3057]

[EC 3.4.22.42 created 2000]

EC 3.4.22.43

**Accepted name:** cathepsin V

**Reaction:** The recombinant enzyme hydrolyses proteins (serum albumin, collagen) and synthetic substrates (Z-

Phe-Arg-NHMec > Z-Leu-Arg-NHMec > Z-Val-Arg-NHMec)

Other name(s): Cathepsin L2; cathepsin U

Comments: Cathepsin V is a human lysosomal cysteine endopeptidase of family C1 (papain family) that is maxi-

mally active at pH 5.7 and unstable at neutral pH. Compound E-64, leupeptin and chicken cystatin are inhibitors. Human cathepsin V shows expression restricted to thymus, testis, corneal epithelium and

some colon and breast carcinomas. Human gene locus: 9q22.2

**References:** [302, 14, 2524]

[EC 3.4.22.43 created 2000]

EC 3.4.22.44

Accepted name: nuclear-inclusion-a endopeptidase

Reaction: Hydrolyses glutaminyl bonds, and activity is further restricted by preferences for the amino acids in

P6 - P1' that vary with the species of potyvirus, e.g. Glu-Xaa-Xaa-Tyr-Xaa-Gln—(Ser or Gly) for the enzyme from tobacco etch virus. The natural substrate is the viral polyprotein, but other proteins and

oligopeptides containing the appropriate consensus sequence are also cleaved.

**Other name(s):** potyvirus NIa protease

**Comments:** The potyviruses cause diseases in plants, and inclusion bodies appear in the host cell nuclei; protein a

of the inclusion bodies is the endopeptidase. The enzyme finds practical use when encoded in vectors for the artificial expression of recombinant fusion proteins, since it can confer on them the capacity for autolytic cleavage. It is also reported that transgenic plants expressing the enzyme are resistant to

viral infection. Type example of peptidase family C4.

**References:** [752, 1443, 2844, 1446]

[EC 3.4.22.44 created 2000]

EC 3.4.22.45

Accepted name: helper-component proteinase

**Reaction:** Hydrolyses a Gly—Gly bond at its own C-terminus, commonly in the sequence -Tyr-Xaa-Val-

Gly—Gly, in the processing of the potyviral polyprotein

Other name(s): HC-Pro

**Comments:** Known from many potyviruses. The helper component-proteinase of the tobacco etch virus is a mul-

tifunctional protein with several known activities: the N-terminal region is required for aphid transmission and efficient genome amplification, the central region is required for long-distance movement in plants, and the C-terminal domain has cysteine endopeptidase activity. Type example of peptidase

family C6.

**References:** [1393, 3058]

[EC 3.4.22.45 created 2001]

EC 3.4.22.46

Accepted name: L-peptidase

**Reaction:** Autocatalytically cleaves itself from the polyprotein of the foot-and-mouth disease virus by hydrolysis

of a Lys—Gly bond, but then cleaves host cell initiation factor eIF-4G at bonds -Gly—Arg- and -

Lys-Arg-

**Comments:** Best known from foot-and-mouth disease virus, but occurs in other aphthoviruses and cardioviruses.

Destruction of initiation factor eIF-4G has the effect of shutting off host-cell protein synthesis while allowing synthesis of viral proteins to continue. The tertiary structure reveals a distant relationship to papain and, consistent with this, compound E-64 is inhibitory. Type example of peptidase family C28.

**References:** [2276, 997]

[EC 3.4.22.46 created 2001]

EC 3.4.22.47

Accepted name: gingipain K

**Reaction:** Endopeptidase with strict specificity for lysyl bonds

**Other name(s):** Lys-gingipain; PrtP proteinase

Comments: Activity is stimulated by glycine. Known from the bacterium Porphyromonas gingivalis and con-

tributes to the pathogenicity of the organism. In peptidase family  ${\tt C25}.$ 

**References:** [2284, 522]

[EC 3.4.22.47 created 2003]

EC 3.4.22.48

Accepted name: staphopain

Reaction: Broad endopeptidase action on proteins including elastin, but rather limited hydrolysis of small-

molecule substrates. Assays are conveniently made with hemoglobin, casein or Z-Phe-Arg-NHMec

as substrate

Other name(s): staphylopain

Comments: Known from species of Staphylococcus. Type example of peptidase family C47.

**References:** [1157, 2314, 657]

[EC 3.4.22.48 created 2003]

### EC 3.4.22.49

**Accepted name:** separase

Reaction: All bonds known to be hydrolysed by this endopeptidase have arginine in P1 and an acidic residue in

P4. P6 is often occupied by an acidic residue or by an hydroxy-amino-acid residue, the phosphoryla-

tion of which enhances cleavage

Other name(s): separin

Comments: In both budding yeast and human cells, cleavage of the cohesin subunit Scc1 by separase is required

for sister chromatid separation in mitosis. Budding yeast separase is also known to cleave the Rec8 subunit of a meiotic cohesin complex and the kinetochore protein Slk19. Type example of peptidase

family C50.

**References:** [3084]

[EC 3.4.22.49 created 2003]

#### EC 3.4.22.50

**Accepted name:** V-cath endopeptidase

**Reaction:** Endopeptidase of broad specificity, hydrolyzing substrates of both cathepsin L and cathepsin B **Other name(s):** AcNPV protease; BmNPV protease; NPV protease; baculovirus cathepsin; nucleopolyhedrosis virus

protease; viral cathepsin

**Comments:** In peptidase family C1. Contributes to the liquefaction of the tissues of the insect host in the late

stages of infection by the baculovirus.

**References:** [2688, 1082]

[EC 3.4.22.50 created 2003]

## EC 3.4.22.51

Accepted name: cruzipain

**Reaction:** Broad endopeptidase specificity similar to that of cathepsin L

Other name(s): congopain; cruzain; evansain; trypanopain

**Comments:** In peptidase family C1. Is located in the digestive vacuoles of the parasitic trypanosome and con-

tributes to the nutrition of the organism by digestion of host proteins.

**References:** [383]

[EC 3.4.22.51 created 2003]

## EC 3.4.22.52

Accepted name: calpain-1

**Reaction:** Broad endopeptidase specificity

**Other name(s):**  $\mu$ -calpain; calcium-activated neutral protease I

**Comments:** In peptidase family C2. Requires Ca<sup>2+</sup> at micromolar concentrations for activity. Cytosolic in animal

cells. The active enzyme molecule is a heterodimer in which the large subunit contains the peptidase

unit, and the small subunit is also a component of EC 3.4.22.53, calpain-2.

**References:** [671]

EC 3.4.22.53

Accepted name: calpain-2

**Reaction:** Broad endopeptidase specificity

**Other name(s):** calcium-activated neutral protease II; *m*-calpain; milli-calpain

**Comments:** Type example of peptidase family C2. Requires  $Ca^{2+}$  at millimolar concentrations for activity. Cy-

tosolic in animal cells. The active enzyme molecule is a heterodimer in which the large subunit con-

tains the peptidase unit, and the small subunit is also a component of EC 3.4.22.52, calpain-1.

**References:** [2783, 671]

[EC 3.4.22.53 created 1981 as EC 3.4.22.17, transferred 2003 to EC 3.4.22.53]

EC 3.4.22.54

Accepted name: calpain-3

**Reaction:** Broad endopeptidase activity

Other name(s): p94; calpain p94; CAPN3; muscle calpain; calpain 3; calcium-activated neutral proteinase 3; muscle-

specific calcium-activated neutral protease 3; CANP 3; calpain L3

**Comments:** This Ca<sup>2+</sup>-dependent enzyme is found in skeletal muscle and is genetically linked to limb girdle mus-

cular dystrophy type 2A [2727, 601]. The enzyme is activated by autoproteolytic cleavage of insertion sequence 1 (IS1), which allows substrates and inhibitors gain access to the active site [601]. Substrates include the protein itself [2412, 601] and connectin/titin [2728, 2198]. Belongs in peptidase

family C2.

**References:** [2727, 2728, 2412, 601, 2198]

[EC 3.4.22.54 created 2007]

EC 3.4.22.55

**Accepted name:** caspase-2

**Reaction:** Strict requirement for an Asp residue at P1, with Asp<sup>316</sup> being essential for proteolytic activity and

has a preferred cleavage sequence of Val-Asp-Val-Ala-Asp-

Other name(s): ICH-1; NEDD-2; caspase-2L; caspase-2S; neural precursor cell expressed developmentally down-

regulated protein 2; CASP-2; NEDD2 protein

**Comments:** Caspase-2 is an initiator caspase, as are caspase-8 (EC 3.4.22.61), caspase-9 (EC 3.4.22.62) and

caspase-10 (EC 3.4.22.63) [393]. Contains a caspase-recruitment domain (CARD) in its N-terminal prodomain, which plays a role in procaspase activation [393]. Two forms of caspase-2 with antagonistic effects exist: caspase-2L induces programmed cell death and caspase-2S suppresses cell death [2,3,5]. Caspase-2 is activated by caspase-3 (EC 3.4.22.56), or by a caspase-3-like protease. Activation involves cleavage of the N-terminal prodomain, followed by self-proteolysis between the large and small subunits of pro-caspase-2 and further proteolysis into smaller fragments [1663]. Proteolysis occurs at Asp residues and the preferred substrate for this enzyme is a pentapeptide rather than a tetrapeptide [3334]. Apart from itself, the enzyme can cleave golgin-16, which is present in the Golgi complex and has a cleavage site that is unique for caspase-2 [1796, 3334]. αΠ-Spectrin, a component of the membrane cytoskeleton, is a substrate of the large isoform of pro-caspase-2 (caspase-2L) but

not of the short isoform (caspase-2S). Belongs in peptidase family C14.

**References:** [1550, 3115, 1663, 1796, 3334, 393]

[EC 3.4.22.55 created 2007]

EC 3.4.22.56

**Accepted name:** caspase-3

**Reaction:** Strict requirement for an Asp residue at positions P1 and P4. It has a preferred cleavage sequence

of Asp-Xaa-Xaa-Asp— with a hydrophobic amino-acid residue at P2 and a hydrophilic amino-acid

residue at P3, although Val or Ala are also accepted at this position

Other name(s): CPP32; apopain; yama protein

**Comments:** Caspase-3 is an effector/executioner caspase, as are caspase-6 (EC 3.4.22.59) and caspase-7 (EC

3.4.22.60) [393]. These caspases are responsible for the proteolysis of the majority of cellular polypeptides [e.g. poly(ADP-ribose) polymerase (PARP)], which leads to the apoptotic phenotype [2079, 393]. Procaspase-3 can be activated by caspase-1 (EC 3.4.22.36), caspase-8 (EC 3.4.22.61), caspase-9 (EC 3.4.22.62) and caspase-10 (EC 3.4.22.63) as well as by the serine protease granzyme B [1533]. Caspase-3 can activate procaspase-2 (EC 3.4.22.55) [1663]. Activation occurs by interdomain cleavage followed by removal of the N-terminal prodomain [1820]. Although Asp-Glu-(Val/Ile)-Asp is thought to be the preferred cleavage sequence, the enzyme can accommodate different residues at P2 and P3 of the substrate [737]. Like caspase-2, a hydrophobic residue at P5 of caspase-3 leads to more efficient hydrolysis, e.g. (Val/Leu)-Asp-Val-Ala-Asp— is a better substrate than Asp-Val-Ala-Asp— . This is not the case for caspase-7 [737]. Belongs in peptidase family C14.

**References:** [1533, 1663, 2079, 737, 393, 1820]

[EC 3.4.22.56 created 2007]

EC 3.4.22.57

**Accepted name:** caspase-4

Reaction: Strict requirement for Asp at the P1 position. It has a preferred cleavage sequence of Tyr-Val-Ala-

Asp—but also cleaves at Asp-Glu-Val-Asp—

**Other name(s):** ICE<sub>rel</sub>II; ICErel-II; Ich-2; transcript X; TX; TX protease; caspase 4; CASP-4

**Comments:** This enzyme is part of the family of inflammatory caspases, which also includes caspase-1 (EC

3.4.22.36) and caspase-5 (EC 3.4.22.58) in humans and caspase-11 (EC 3.4.22.64), caspase-12, caspase-13 and caspase-14 in mice. Contains a caspase-recruitment domain (CARD) in its N-terminal prodomain, which plays a role in procaspase activation [3,5,6]. The enzyme is able to cleave itself and the p30 caspase-1 precursor, but, unlike caspase-1, it is very inefficient at generating mature interleukin-1 $\beta$  (IL-1 $\beta$ ) from pro-IL-1 $\beta$  [744, 741]. Both this enzyme and caspase-5 can cleave procaspase-3 to release the small subunit (p12) but not the large subunit (p17) [1369]. The caspase-1 inhibitor Ac-Tyr-Val-Ala-Asp-CHO can also inhibit this enzyme, but more slowly [741]. Belongs in

peptidase family C14.

**References:** [744, 1371, 1369, 741, 1823, 393]

[EC 3.4.22.57 created 2007]

EC 3.4.22.58

Accepted name: caspase-5

Reaction: Strict requirement for Asp at the P1 position. It has a preferred cleavage sequence of Tyr-Val-Ala-

Asp—but also cleaves at Asp-Glu-Val-Asp—

Other name(s): ICErel-III; Ich-3; ICH-3 protease; transcript Y; TY; CASP-5

Comments: This enzyme is part of the family of inflammatory caspases, which also includes caspase-1 (EC

3.4.22.36) and caspase-4 (EC 3.4.22.57) in humans and caspase-11 (EC 3.4.22.64), caspase-12, caspase-13 and caspase-14 in mice. Contains a caspase-recruitment domain (CARD) in its N-terminal prodomain, which plays a role in procaspase activation [3,5,6]. The enzyme is able to cleave itself and the p30 caspase-1 precursor, but is very inefficient at generating mature interleukin-1 $\beta$  (IL-1 $\beta$ ) from pro-IL-1 $\beta$  [743, 741]. Both this enzyme and caspase-4 can cleave pro-caspase-3 to release the small subunit (p12) but not the large subunit (p17) [1696]. Unlike caspase-4, this enzyme can be induced by

lipopolysaccharide [1696]. Belongs in peptidase family C14.

**References:** [743, 1369, 1696, 741, 1823, 393]

[EC 3.4.22.58 created 2007]

EC 3.4.22.59

**Accepted name:** caspase-6

Reaction: Strict requirement for Asp at position P1 and has a preferred cleavage sequence of Val-Glu-His-

Asp—

Other name(s): CASP-6; apoptotic protease Mch-2; Mch2

Comments: Caspase-6 is an effector/executioner caspase, as are caspase-3 (EC 3.4.22.56) and caspase-7 (EC

3.4.22.60) [393]. These caspases are responsible for the proteolysis of the majority of cellular polypeptides [e.g. poly(ADP-ribose) polymerase (PARP)], which leads to the apoptotic phenotype [393]. Caspase-6 can cleave its prodomain to produce mature caspase-6, which directly activates caspase-8 (EC 3.4.22.61) and leads to the release of cytochrome c from the mitochondria. The release of cytochrome c is an essential component of the intrinsic apoptosis pathway [500]. The enzyme can also cleave and inactivate lamins, the intermediate filament scaffold proteins of the nuclear envelope, leading to nuclear fragmentation in the final phases of apoptosis [2,4,5,6]. Belongs in peptidase

family C14.

**References:** [500, 393, 1384, 1629, 1769, 2836]

[EC 3.4.22.59 created 2007]

EC 3.4.22.60

**Accepted name:** caspase-7

**Reaction:** Strict requirement for an Asp residue at position P1 and has a preferred cleavage sequence of Asp-

Glu-Val-Asp—

**Other name(s):** CASP-7; ICE-like apoptotic protease 3; ICE-LAP3; apoptotic protease Mch-3; Mch3; CMH-1

Comments: Caspase-7 is an effector/executioner caspase, as are caspase-3 (EC 3.4.22.56) and caspase-6 (EC

3.4.22.59) [393]. These caspases are responsible for the proteolysis of the majority of cellular polypeptides [e.g. poly(ADP-ribose) polymerase (PARP)], which leads to the apoptotic phenotype [2079]. Although a hydrophobic residue at P5 of caspase-2 (EC 3.4.22.55) and caspase-3 leads to more efficient hydrolysis, the amino-acid residue at this location in caspase-7 has no effect [737]. Caspase-7 is activated by the initiator caspases [caspase-8 (EC 3.4.22.61), caspase-9 (EC 3.4.22.62) and caspase-10 (EC 3.4.22.63)]. Removal of the N-terminal prodomain occurs before cleavage in the

linker region between the large and small subunits [582]. Belongs in peptidase family C14.

**References:** [393, 2079, 737, 582]

[EC 3.4.22.60 created 2007]

EC 3.4.22.61

Accepted name: caspase-8

**Reaction:** Strict requirement for Asp at position P1 and has a preferred cleavage sequence of (Leu/Asp/Val)-

Glu-Thr-Asp—(Gly/Ser/Ala)

**Other name(s):** FLICE, FADD-like ICE; MACH; MORT<sub>1</sub>-associated CED-3 homolog; Mch5; mammalian Ced-3

homolog 5; CASP-8; ICE-like apoptotic protease 5; FADD-homologous ICE/CED-3-like protease;

apoptotic cysteine protease; apoptotic protease Mch-5; CAP4

Comments: Caspase-8 is an initiator caspase, as are caspase-2 (EC 3.4.22.55), caspase-9 (EC 3.4.22.62) and

caspase-10 (EC 3.4.22.63) [393]. Caspase-8 is the apical activator of the extrinsic (death receptor) apoptosis pathway, triggered by death receptor ligation [256]. It contains two tandem death effector domains (DEDs) in its N-terminal prodomain, and these play a role in procaspase activation [393]. This enzyme is linked to cell surface death receptors such as Fas [393, 774]. When Fas is aggregated by the Fas ligand, procaspase-8 is recruited to the death receptor where it is activated [393]. The enzyme has a preference for Glu at P3 and prefers small residues, such as Gly, Ser and Ala, at the P1' position. It has very broad P4 specificity, tolerating substrates with Asp, Val or Leu in this position [2,3,4]. Endogenous substrates for caspase-8 include procaspase-3, the pro-apoptotic Bcl-2 family member Bid, RIP, PAK2 and the caspase-8 activity modulator FLIP, [2511, 774]. Belongs in pepti-

dase family C14.

**References:** [393, 256, 2014, 2511, 774, 237, 249]

[EC 3.4.22.61 created 2007]

### EC 3.4.22.62

Accepted name: caspase-9

Reaction: Strict requirement for an Asp residue at position P1 and with a marked preference for His at position

P2. It has a preferred cleavage sequence of Leu-Gly-His-Asp—Xaa

Other name(s): CASP-9; ICE-like apoptotic protease 6; ICE-LAP6; apoptotic protease Mch-6; apoptotic protease

activating factor 3; APAF-3

Comments: Caspase-9 is an initiator caspase, as are caspase -2 (EC 3.4.22.55), caspase-8 (EC 3.4.22.61) and

caspase-10 (EC 3.4.22.63) [393]. Caspase-9 contains a caspase-recruitment domain (CARD) in its N-terminal prodomain, which plays a role in procaspase activation [393]. An alternatively spliced version of caspase-9 also exists, caspase-9S, that inhibits apoptosis, similar to the situation found with caspase-2 [393]. Phosphorylation of caspase-9 from some species by Akt, a serine-threonine protein kinase, inhibits caspase activity in vitro and caspase activation in vivo [393]. The activity of caspase-9 is increased dramatically upon association with the apoptosome but the enzyme can be activated without proteolytic cleavage [3284, 250]. Procaspase-3 is the enzyme's physiological substrate [3284].

Belongs in peptidase family C14.

**References:** [393, 3284, 250, 2512]

[EC 3.4.22.62 created 2007]

#### EC 3.4.22.63

Accepted name: caspase-10

**Reaction:** Strict requirement for Asp at position P1 and has a preferred cleavage sequence of Leu-Gln-Thr-

Asp—Gly

Other name(s): FLICE2, Mch4; CASP-10; ICE-like apoptotic protease 4; apoptotic protease Mch-4; FAS-associated

death domain protein interleukin-1β-converting enzyme 2

Comments: Caspase-10 is an initiator caspase, as are caspase-2 (EC 3.4.22.55), caspase-8 (EC 3.4.22.61) and

caspase-9 (EC 3.4.22.62) [393]. Like caspase-8, caspase-10 contains two tandem death effector domains (DEDs) in its N-terminal prodomain, and these play a role in procaspase activation [393]. The enzyme has many overlapping substrates in common with caspase-8, such as RIP (the cleavage of which impairs NF-kB survival signalling and starts the cell-death process) and PAK2 (associated with some of the morphological features of apoptosis, such as cell rounding and apoptotic body formation) [774]. Bid, a Bcl2 protein, can be cleaved by caspase-3 (EC 3.4.22.56), caspase-8 and caspase-10 at Lys-Gln-Thr-Asp—to yield the pro-apoptotic p15 fragment. The p15 fragment is N-myristoylated and enhances the release of cytochrome *c* from mitochondria (which, in turn, initiatiates the intrinsic apoptosis pathway). Bid can be further cleaved by caspase-10 and granzyme B but not by caspase-3 or caspase-8 at Ile-Glu-Thr-Asp—to yield a p13 fragment that is not N-myristoylated [774]. Belongs

in peptidase family C14.

**References:** [393, 774, 2631, 249]

[EC 3.4.22.63 created 2007]

# EC 3.4.22.64

Accepted name: caspase-11

**Reaction:** Strict requirement for Asp at the P1 position and has a preferred cleavage sequence of

(Ile/Leu/Val/Phe)-Gly-His-Asp—

Other name(s): CASP-11

Comments: This murine enzyme is part of the family of inflammatory caspases, which also includes caspase-1

(EC 3.4.22.36), caspase-4 (EC 3.4.22.57) and caspase-5 (EC 3.4.22.58) in humans and caspase-12, caspase-13 and caspase-14 in mice. Contains a caspase-recruitment domain (CARD) in its N-terminal prodomain, which plays a role in procaspase activation. Like caspase-5, but unlike caspase-4, this enzyme can be induced by lipopolysaccharide [1386]. This enzyme not only activates caspase-1, which is required for the maturation of proinflammatory cytokines such as interleukin-1 $\beta$  (IL-1 $\beta$ ) and IL-18, but it also activates caspase-3 (EC 3.4.22.56), which leads to cellular apoptosis under pathological

conditions [1386, 1209]. Belongs in peptidase family C14.

**References:** [1386, 1209, 3118, 703, 393]

EC 3.4.22.65

**Accepted name:** peptidase 1 (mite)

**Reaction:** Broad endopeptidase specificity

Other name(s): allergen Der f 1; allergen Der p 1; antigen Der p 1; antigen Eur m 1; antigen Pso o 1; major mite fecal

allergen Der p 1; Der p 1; Der f 1; Eur m 1; endopeptidase 1 (mite)

Comments: This enzyme, derived from the house dust mite, is a major component of the allergic immune re-

sponse [1366]. The substrate specificity of this enzyme is not altogether clear. It cleaves the low-affinity IgE receptor CD23 at Glu<sup>298</sup>—Ser<sup>299</sup> and Ser<sup>155</sup>—Ser<sup>156</sup> [1879]. It also cleaves the pulmonary structural proteins occludin and claudin at Leu—Leu, Asp—Leu and at Gly—Thr bonds [1879, 1366]. It can also cleave the α subunit of the interleukin-2 (IL-2) receptor (CD25) [2593]. Using a positional scanning combinatorial library, it was found that the major substrate-specificity determinant is for Ala in the P2 position [1060]. The enzyme shows only a slight preference for basic amino acids in the P1 and P3 positions and a preference for aliphatic amino acids such as Ile, Pro, Val,

Leu and norleucine in the P4 position [1060]. Belongs in peptidase family C1A.

**References:** [1879, 1366, 1060, 2593, 2592, 2850]

[EC 3.4.22.65 created 2007]

EC 3.4.22.66

Accepted name: calicivirin

**Reaction:** Endopeptidase with a preference for cleavage when the P1 position is occupied by Glu—and the P1'

position is occupied by Gly—

Other name(s): Camberwell virus processing peptidase; Chiba virus processing peptidase; Norwalk virus processing

peptidase; Southampton virus processing peptidase; Southampton virus; norovirus virus processing peptidase; calicivirus trypsin-like cysteine protease; calicivirus TCP; calicivirus 3C-like protease; cali-

civirus endopeptidase; rabbit hemorrhagic disease virus 3C endopeptidase

**Comments:** Viruses that are members of the Norovirus genus (Caliciviridae family) are a major cause of epidemic

acute viral gastroenteritis [1715]. The nonstructural proteins of these viruses are produced by proteolytic cleavage of a large precursor polyprotein, performed by a protease that is incorporated into the polyprotein []. Cleavage sites are apparently defined by features based on both sequence and structure since several sites in the polyprotein fulfilling the identified sequence requirements are not cleaved [1899]. The presence of acidic (Asp), basic (Arg), aromatic (Tyr) or aliphatic (Leu) amino acids at the P1' position results in only minor differences in cleavage efficiency, suggesting that steric or conformational constraints may play a role in determining specificity [1899]. Changes to the amino acid at the P2 position do not alter cleavage efficiency [1899, 3199]. Belongs in peptidase family C37.

**References:** [1899, 3199, 40, 1715, 1716]

[EC 3.4.22.66 created 2007]

EC 3.4.22.67

Accepted name: zingipain

**Reaction:** Preferential cleavage of peptides with a proline residue at the P2 position **Other name(s):** ginger protease; GP-I; GP-II; ginger protease II (*Zingiber officinale*); zingibain

**Comments:** This enzyme is found in ginger (*Zingiber officinale*) rhizome and is a member of the papain family.

GP-II contains two glycosylation sites. The enzyme is inhibited by some divalent metal ions, such as

 $Hg^{2+}$ ,  $Cu^{2+}$ ,  $Cd^{2+}$  and  $Zn^{2+}$  [2174]. Belongs in peptidase family C1.

**References:** [438, 2174, 439]

[EC 3.4.22.67 created 2007]

EC 3.4.22.68

**Accepted name:** Ulp1 peptidase

**Reaction:** Hydrolysis of the  $\alpha$ -linked peptide bond in the sequence Gly-Gly—Ala-Thr-Tyr at the C-terminal

end of the small ubiquitin-like modifier (SUMO) propeptide, Smt3, leading to the mature form of the protein. A second reaction involves the cleavage of an  $\epsilon$ -linked peptide bond between the C-terminal

glycine of the mature SUMO and the lysine  $\epsilon$ -amino group of the target protein

Other name(s): Smt3-protein conjugate proteinase; Ubl-specific protease 1; Ulp1; Ulp1 endopeptidase; Ulp1 protease

**Comments:** The enzyme from *Saccharomyces cerevisiae* can also recognize small ubiquitin-like modifier 1

(SUMO-1) from human as a substrate in both SUMO-processing ( $\alpha$ -linked peptide bonds) and SUMO-deconjugation ( $\epsilon$ -linked peptide bonds) reactions [1,2,3]. Ulp1 has several functions, including an essential role in chromosomal segregation and progression of the cell cycle through the G2/M

phase of the cell cycle. Belongs in peptidase family C48.

**References:** [1689, 1668, 2895, 1669, 1223, 1992]

[EC 3.4.22.68 created 2008, modified 2011]

EC 3.4.22.69

**Accepted name:** SARS coronavirus main proteinase

**Reaction:** TSAVLQ—SGFRK-NH<sub>2</sub> and SGVTFQ—GKFKK the two peptides corresponding to the two self-

cleavage sites of the SARS 3C-like proteinase are the two most reactive peptide substrates. The enzyme exhibits a strong preference for substrates containing Gln at P1 position and Leu at P2 position.

Other name(s): 3cLpro; 3C-like protease; coronavirus 3C-like protease; Mpro; SARS 3C-like protease; SARS coro-

navirus 3CL protease; SARS coronavirus main peptidase; SARS coronavirus main protease; SARS-CoV 3CLpro enzyme; SARS-CoV main protease; SARS-CoV Mpro; severe acute respiratory syn-

drome coronavirus main protease

Comments: SARS coronavirus main protease is the key enzyme in SARS coronavirus replicase polyprotein pro-

cessing. In peptidase family C30.

**References:** [938, 736, 24]

[EC 3.4.22.69 created 2009]

EC 3.4.22.70

**Accepted name:** sortase A

**Reaction:** The enzyme catalyses a cell wall sorting reaction in which a surface protein with a sorting signal con-

taining a LPXTG motif is cleaved between the Thr and Gly residue. The resulting threonine carboxyl

end of the protein is covalently attached to a pentaglycine cross-bridge of peptidoglycan.

**Other name(s):** SrtA; SrtA protein; SrtA sortase

**Comments:** In peptidase family C60. **References:** [2941, 3346, 2351]

[EC 3.4.22.70 created 2009]

EC 3.4.22.71

**Accepted name:** sortase B

**Reaction:** The enzyme catalyses a cell wall sorting reaction in which a surface protein with a sorting signal con-

taining a NPXTN motif is cleaved between the Thr and Asn residue. The resulting threonine carboxyl

end of the protein is covalently attached to a pentaglycine cross-bridge of peptidoglycan.

Other name(s): SrtB

**Comments:** In peptidase family C60.

**References:** [3347, 213, 470]

[EC 3.4.22.71 created 2009]

# EC 3.4.23 Aspartic endopeptidases

EC 3.4.23.1

**Accepted name:** pepsin A

**Reaction:** Preferential cleavage: hydrophobic, preferably aromatic, residues in P1 and P1' positions. Cleaves

Phe<sup>1</sup>—Val, Gln<sup>4</sup>—His, Glu<sup>13</sup>—Ala, Ala<sup>14</sup>—Leu, Leu<sup>15</sup>—Tyr, Tyr<sup>16</sup>—Leu, Gly<sup>23</sup>—Phe,

Phe<sup>24</sup> Phe and Phe<sup>25</sup> Tyr bonds in the B chain of insulin

Other name(s): pepsin; lactated pepsin; pepsin fortior; fundus-pepsin; elixir lactate of pepsin; P I; lactated pepsin

elixir; P II; pepsin R; pepsin D

**Comments:** The predominant endopeptidase in the gastric juice of vertebrates, formed from pepsinogen A by lim-

ited proteolysis. Human pepsin A occurs in five molecular forms. Pig pepsin D [1620, 1619] is un-

phosphorylated pepsin A. Type example of peptidase family A1.

**References:** [1620, 1619, 793, 1305, 826, 2881, 2300]

[EC 3.4.23.1 created 1961 as EC 3.4.4.1, transferred 1972 to EC 3.4.23.1, modified 1986, modified 1989]

EC 3.4.23.2

**Accepted name:** pepsin B

Reaction: Degradation of gelatin; little activity on hemoglobin. Specificity on B chain of insulin more restricted

than that of pepsin A; does not cleave at Phe<sup>1</sup>-Val, Gln<sup>4</sup>-His or Gly<sup>23</sup>-Phe

**Other name(s):** parapepsin I; pig gelatinase

**Comments:** Formed from pig pepsinogen B. In peptidase family A1 (pepsin A family)

**References:** [2482]

[EC 3.4.23.2 created 1961 as EC 3.4.4.2, transferred 1972 to EC 3.4.23.2, modified 1986]

EC 3.4.23.3

Accepted name: gastricsin

**Reaction:** More restricted specificity than pepsin A, but shows preferential cleavage at Tyr—bonds. High activ-

ity on hemoglobin

Other name(s): pepsin C; pig parapepsin II; parapepsin II

**Comments:** Formed from progastricsin, apparently in the gastric juice of most vertebrates. In addition to the fun-

dus, progastricsin is also secreted in antrum and proximal duodenum. Seminal plasma contains a zymogen that is immunologically identical with progastricsin [2402]. In peptidase family A1 (pepsin A

family).

**References:** [2482, 2880, 791, 792, 1819, 2402, 1088]

[EC 3.4.23.3 created 1965 as EC 3.4.4.22, transferred 1972 to EC 3.4.23.3, modified 1986]

EC 3.4.23.4

Accepted name: chymosin

**Reaction:** Broad specificity similar to that of pepsin A. Clots milk by cleavage of a single Ser-Phe<sup>105</sup>—Met-Ala

bond in κ-chain of casein

Other name(s): rennin (but this should be avoided since it leads to confusion with renin)

**Comments:** Neonatal gastric enzyme with high milk clotting and weak general proteolytic activity, formed from

prochymosin. Found among mammals with postnatal uptake of immunoglobulins. In peptidase family

A1(pepsin A family)

**References:** [790, 1063, 3070]

[EC 3.4.23.4 created 1961 as EC 3.4.4.3, transferred 1972 to EC 3.4.23.4, modified 1986]

EC 3.4.23.5

Accepted name: cathepsin D

**Reaction:** Specificity similar to, but narrower than, that of pepsin A. Does not cleave the Gln<sup>4</sup>-His bond in B

chain of insulin

Comments: Occurs intracellularly, in lysosomes. A zymogen form is known [478]. In peptidase family A1 (pepsin

A family).

**References:** [154, 2847, 748, 478]

[EC 3.4.23.5 created 1965 as EC 3.4.4.23, transferred 1972 to EC 3.4.23.5, modified 1986]

[3.4.23.6 Transferred entry. now EC 3.4.23.30 pycnoporopepsin]

[EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981, deleted 1992 [EC 3.4.23.7, EC 3.4.23.8, EC

 $3.4.23.9, EC\ 3.4.23.10, EC\ 3.4.99.1, EC\ 3.4.99.15\ and\ EC\ 3.4.99.25\ all\ created\ 1972\ and\ incorporated\ 1978]]$ 

[3.4.23.7 Transferred entry. Penicillium janthinellum acid proteinase. Now EC 3.4.23.20, penicillopepsin]

[EC 3.4.23.7 created 1972, modified 1981, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]

[3.4.23.8 Transferred entry. yeast proteinase A. Now EC 3.4.23.25, saccharopepsin]

[EC 3.4.23.8 created 1972, modified 1981, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]

[3.4.23.9 Transferred entry. Rhizopus acid proteinase. Now EC 3.4.23.21, rhizopuspepsin]

[EC 3.4.23.9 created 1972, modified 1981, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]

[3.4.23.10 Transferred entry. Endothia acid proteinase. Now EC 3.4.23.22, endothiapepsin]

[EC 3.4.23.10 created 1972, modified 1981, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]

[3.4.23.11 Deleted entry. thyroid aspartic proteinase]

[EC 3.4.23.11 created 1978, modified 1981, deleted 1992]

### EC 3.4.23.12

Accepted name: nepenthesin

**Reaction:** Similar to pepsin, but also cleaves on either side of Asp and at Lys—Arg

Other name(s): Nepenthes aspartic proteinase; Nepenthes acid proteinase; nepenthacin; nepenthasin; aspartyl en-

dopeptidase

**Comments:** From the insectivorous plants *Nepenthes* spp. (secretions) and *Drosera peltata* (ground-up leaves).

Aspartic endopeptidases are probably present in many other plants, including Lotus [2638] and

sorghum [879]. In peptidase family A1 (pepsin A family)

**References:** [44, 879, 2638, 43, 2839, 2929]

[EC 3.4.23.12 created 1972 as EC 3.4.99.4, transferred 1978 to EC 3.4.23.12, modified 1981]

[3.4.23.13 Deleted entry. Lotus aspartic proteinase]

[EC 3.4.23.13 created 1978, modified 1981, deleted 1992]

[3.4.23.14 Deleted entry. sorghum aspartic proteinase]

[EC 3.4.23.14 created 1978, modified 1981, deleted 1992]

### EC 3.4.23.15

Accepted name: renin

**Reaction:** Cleavage of Leu—bond in angiotensinogen to generate angiotensin I

Other name(s): angiotensin-forming enzyme; angiotensinogenase

**Comments:** Formed from prorenin in plasma and kidney. In peptidase family A1 (pepsin A family).

**References:** [1247, 2689, 1246, 2655]

#### EC 3.4.23.16

**Accepted name:** HIV-1 retropepsin

**Reaction:** Specific for a P1 residue that is hydrophobic, and P1' variable, but often Pro

Other name(s): human immunodeficiency virus type 1 protease; gag protease; HIV aspartyl protease; HIV proteinase;

retroproteinase; HIV-1 protease; HIV-2 protease

Comments: Present in human immunodeficiency virus type 1. Contributes to the maturation of the viral particle,

and is a target of antiviral drugs. Active enzyme is a dimer of identical 11-kDa subunits. Similar en-

zymes occur in other retroviruses [1560]. Type example of peptidase family A2

**References:** [1560, 667]

[EC 3.4.23.16 created 1992, modified 2000]

### EC 3.4.23.17

**Accepted name:** *pro*-opiomelanocortin converting enzyme

**Reaction:** Cleavage at paired basic residues in certain prohormones, either between them, or on the carboxyl

side

Other name(s): prohormone converting enzyme; pro-opiomelanocortin-converting enzyme; proopiomelanocortin pro-

teinase; PCE

**Comments:** A 70 kDa membrane-bound enzyme isolated from cattle pituitary secretory vesicle.

**References:** [1733, 1732, 721]

[EC 3.4.23.17 created 1989 as EC 3.4.99.38, transferred 1992 to EC 3.4.23.17]

#### EC 3.4.23.18

Accepted name: aspergillopepsin I

**Reaction:** Hydrolysis of proteins with broad specificity. Generally favours hydrophobic residues in P1 and P1',

but also accepts Lys in P1, which leads to activation of trypsinogen. Does not clot milk

Other name(s): Aspergillus acid protease; Aspergillus acid proteinase; Aspergillus aspartic proteinase; Aspergillus

awamori acid proteinase; Aspergillus carboxyl proteinase; (see also Comments); carboxyl proteinase; Aspergillus kawachii aspartic proteinase; Aspergillus saitoi acid proteinase; pepsin-type aspartic proteinase; Aspergillus niger acid proteinase; sumizyme AP; proctase P; denapsin; denapsin XP 271;

proctase

Comments: Found in a variety of Aspergillus species (imperfect fungi): Aspergillus awamori (awamorin, as-

pergillopepsin A: [2210]), *A. foetidus* (aspergillopepsin F: [2209]), *A. fumigatus* [2231], *A. kawachii* [3239], *A. niger* (proteinase B, proctase B: [1968, 397]), *A. oryzae* (trypsinogen kinase: [545, 1782]), *A. saitoi* (aspergillopeptidase A: [1782]), and *A. sojae* [2871, 1782]. In peptidase family A1 (pepsin A

family). Formerly included in EC 3.4.23.6

**References:** [1525, 1968, 545, 397, 2871, 2209, 2231, 2210, 3239, 1782]

[EC 3.4.23.18 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

#### EC 3.4.23.19

Accepted name: aspergillopepsin II

**Reaction:** Preferential cleavage in B chain of insulin: Asn<sup>3</sup>—Gln, Gly<sup>13</sup>—Ala, Tyr<sup>26</sup>—Thr **Other name(s):** proteinase A; *Aspergillus niger* var. macrosporus aspartic proteinase

**Comments:** Isolated from *Aspergillus niger* var. macrosporus, distinct from proteinase B (see aspergillopepsin I)

in specificity and insensitivity to pepstatin. In peptidase family A4 (scytalidopepsin B family). For-

merly included in EC 3.4.23.6

**References:** [397, 1226]

[EC 3.4.23.19 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

### EC 3.4.23.20

**Accepted name:** penicillopepsin

**Reaction:** Hydrolysis of proteins with broad specificity similar to that of pepsin A, preferring hydrophobic

residues at P1 and P1', but also cleaving Gly<sup>20</sup>—Glu in the B chain of insulin. Clots milk, and acti-

vates trypsinogen

Other name(s): peptidase A; Penicillium janthinellum aspartic proteinase; acid protease A; Penicillium citrinum acid

proteinase; *Penicillium cyclopium* acid proteinase; *Penicillium expansum* acid proteinase; *Penicillium janthinellum* acid proteinase; *Penicillium expansum* aspartic proteinase; *Penicillium caseicolum* aspartic proteinase; *Penicillium roqueforti* acid proteinase; *Penicillium* 

duponti aspartic proteinase; Penicillium citrinum aspartic proteinase

**Comments:** From the imperfect fungus *Penicillium janthinellum*. In peptidase family A1 (pepsin A family).

Closely related enzymes have been isolated from *P. roqueforti* [3321] and *P. duponti* [698].

**References:** [1779, 3321, 698, 1158, 1197]

[EC 3.4.23.20 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

#### EC 3.4.23.21

Accepted name: rhizopuspepsin

**Reaction:** Hydrolysis of proteins with broad specificity similar to that of pepsin A, preferring hydrophobic

residues at P1 and P1'. Clots milk and activates trypsinogen. Does not cleave Gln<sup>4</sup>-His, but does

cleave His<sup>10</sup>—Leu and Val<sup>12</sup>—Glu in B chain of insulin

Other name(s): Rhizopus aspartic proteinase; neurase; Rhizopus acid protease; Rhizopus acid proteinase

**Comments:** From the zygomycete fungus *Rhizopus chinensis*. A similar endopeptidase is found in *R. niveus* 

[1574]. In peptidase family A1 (pepsin A family).

**References:** [2987, 1574, 2175, 2796]

[EC 3.4.23.21 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

#### EC 3.4.23.22

Accepted name: endothiapepsin

**Reaction:** Hydrolysis of proteins with specificity similar to that of pepsin A; prefers hydrophobic residues at P1

and P1', but does not cleave Ala<sup>14</sup>-Leu in the B chain of insulin or Z-Glu-Tyr. Clots milk

Other name(s): Endothia aspartic proteinase; Endothia acid proteinase; Endothia parasitica acid proteinase; Endothia

parasitica aspartic proteinase

**Comments:** From the ascomycete *Endothia parasitica*. In peptidase family A1 (pepsin A family).

**References:** [3172, 3192, 143, 486]

[EC 3.4.23.22 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

### EC 3.4.23.23

Accepted name: mucorpepsin

**Reaction:** Hydrolysis of proteins, favouring hydrophobic residues at P1 and P1'. Clots milk. Does not accept

Lys at P1, and hence does not activate trypsinogen

Other name(s): Mucor rennin; Mucor aspartic proteinase; Mucor acid proteinase; Mucor acid protease; Mucor miehei

aspartic proteinase; Mucor miehei aspartic protease; Mucor aspartic proteinase; Mucor pusillus empo-

rase; Fromase 100; Mucor pusillus rennin; Fromase 46TL; Mucor miehei rennin

**Comments:** Isolated from the zygomycete fungi *Mucor pusillus* and *M. miehei*. The two species variants show

83% sequence identity and are immunologically crossreactive. In peptidase family A1 (pepsin A fam-

ily). Formerly included in EC 3.4.23.6

**References:** [73, 2214, 2765, 2178, 171]

[EC 3.4.23.23 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

### EC 3.4.23.24

Accepted name: candidapepsin

**Reaction:** Preferential cleavage at the carboxyl of hydrophobic amino acids, but fails to cleave Leu<sup>15</sup>-Tyr, Tyr<sup>16</sup>-

Leu and Phe<sup>24</sup>-Phe of insulin B chain. Activates trypsinogen, and degrades keratin

Other name(s): Candida albicans aspartic proteinase; Candida albicans carboxyl proteinase; Candida albicans secre-

tory acid proteinase; Candida olea acid proteinase; Candida aspartic proteinase; Candida olea aspar-

tic proteinase; Candida albicans aspartic proteinase

**Comments:** This endopeptidase from the imperfect yeast *Candida albicans* is inhibited by pepstatin, but not by

methyl 2-diazoacetamidohexanoate or 1,2-epoxy-3-(p-nitrophenoxy)propane. In peptidase family A1

(pepsin A family). Formerly included in EC 3.4.23.6

**References:** [2406, 2475, 2065, 1739]

[EC 3.4.23.24 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

### EC 3.4.23.25

Accepted name: saccharopepsin

**Reaction:** Hydrolysis of proteins with broad specificity for peptide bonds. Cleaves -Leu-Val-Tyr bond in

a synthetic substrate. Does not act on esters of Tyr or Arg

Other name(s): yeast endopeptidase A; Saccharomyces aspartic proteinase; aspartic proteinase yscA; proteinase A;

proteinase yscA; yeast proteinase A; Saccharomyces cerevisiae aspartic proteinase A; yeast proteinase

A; PRA

Comments: Located in the vacuole of the baker's yeast (Saccharomyces cerevisiae) cell. In peptidase family A1

(pepsin A family).

**References:** [1074, 1894, 47]

[EC 3.4.23.25 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

### EC 3.4.23.26

Accepted name: rhodotorulapepsin

**Reaction:** Specificity similar to that of pepsin A. Cleaves Z-Lys—Ala-Ala and activates trypsinogen **Other name(s):** Rhodotorula aspartic proteinase; *Cladosporium* acid proteinase; *Pae-*

cilomyces proteinase; *Cladosporium* aspartic proteinase; Paecilomyces proteinase; *Rhodotorula glutinis* aspartic proteinase; *Rhodotorula glutinis* acid proteinase; *Rhodotorula glutinis* aspartic proteinase

II; Rhodotorula acid proteinase

Comments: From the imperfect yeast *Rhodotorula glutinis*. Somewhat similar enzymes have been isolated from

the imperfect yeast-like organism Cladosporium sp. [2006, 2129] and the imperfect fungus Pae-

cilomyces varioti [2543, 2544].

**References:** [2543, 2544, 1368, 2006, 2130, 2129, 2838, 1782]

[EC 3.4.23.26 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

[3.4.23.27 Transferred entry. physaropepsin. Now EC 3.4.21.103, physarolisin]

[EC 3.4.23.27 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992), deleted 2003]

#### EC 3.4.23.28

Accepted name: acrocylindropepsin

**Reaction:** Preference for hydrophobic residues at P1 and P1'. Action on the B chain of insulin is generally simi-

lar to that of pepsin A, but it also cleaves Leu<sup>6</sup>—Cys(SO3H), Glu<sup>21</sup>—Arg and Asn<sup>3</sup>—Gln, although

not Gln<sup>4</sup>-His

Other name(s): Acrocylindrium proteinase; Acrocylindrium acid proteinase

**Comments:** From the imperfect fungus *Acrocylindrium* sp. Has a very low pH optimum on casein of 2.0. In pepti-

dase family A1 (pepsin A family).

**References:** [3007, 1218, 2838]

[EC 3.4.23.28 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

#### EC 3.4.23.29

Accepted name: polyporopepsin

Reaction: Milk clotting activity, broad specificity, but fails to cleave Leu<sup>15</sup>-Tyr or Tyr<sup>16</sup>-Leu of insulin B chain

Other name(s): Polyporus aspartic proteinase; *Irpex lacteus* aspartic proteinase; *Irpex lacteus* carboxyl proteinase B

From the basidiomycete *Polyporus tulipiferae* (formerly *Irpex lacteus*). In peptidase family A1

(pepsin A family)

**References:** [1485, 1487]

[EC 3.4.23.29 created 1992]

# EC 3.4.23.30

Accepted name: pycnoporopepsin

**Reaction:** Similar to pepsin A, but narrower, cleaving only three bonds in the B chain of insulin: Ala<sup>14</sup>—Leu,

Tyr<sup>16</sup>—Leu, and Phe<sup>24</sup>—Phe

Other name(s): proteinase Ia; Pycnoporus coccineus aspartic proteinase; Trametes acid proteinase

**Comments:** From the basidiomycete *Pycnoporus sanguineus*, formerly known as *P. coccineus* and *Trametes san-*

guinea. Formerly included in EC 3.4.23.6

**References:** [2939, 2987, 1219]

[EC 3.4.23.30 created 1992 (EC 3.4.23.6 created 1992 (EC 3.4.23.6 created 1961 as EC 3.4.4.17, transferred 1972 to EC 3.4.23.6, modified 1981 [EC 3.4.23.7, EC 3.4.23.8, EC 3.4.23.9, EC 3.4.23.10, EC 3.4.99.1, EC 3.4.99.15 and EC 3.4.99.25 all created 1972 and incorporated 1978], part incorporated 1992)]

### EC 3.4.23.31

Accepted name: scytalidopepsin A

**Reaction:** Hydrolysis of proteins with specificity similar to that of pepsin A, but also cleaves Cys(SO3H)<sup>7</sup>—Gly

and Leu<sup>17</sup>-Val in the B chain of insulin

Other name(s): Scytalidium aspartic proteinase A; Scytalidium lignicolum aspartic proteinase; Scytalidium lignicolum

aspartic proteinase A-2; *Scytalidium lignicolum* aspartic proteinase A-I; *Scytalidium lignicolum* aspartic proteinase C; *Scytalidium lignicolum* carboxyl proteinase; *Scytalidium lignicolum* acid proteinase

**Comments:** Isolated from the imperfect fungus Scytalidium lignicolum. Not inhibited by pepstatin-Ac, methyl 2-

diazoacetamidohexanoate or 1,2-epoxy-3-(p-nitrophenyl)propane. A related enzyme from the same

organism, proteinase C, is also insensitive to these inhibitors and has  $M_r = 406,000$  [2136]

**References:** [2131, 2132, 2136]

[EC 3.4.23.31 created 1992]

EC 3.4.23.32

Accepted name: scytalidopepsin B

Hydrolysis of proteins with broad specificity, cleaving Phe<sup>24</sup>—Phe, but not Leu<sup>15</sup>-Tyr and Phe<sup>25</sup>-Tyr **Reaction:** 

in the B chain of insulin

Other name(s): Scytalidium aspartic proteinase B; Ganoderma lucidum carboxyl proteinase; Ganoderma lucidum

aspartic proteinase; Scytalidium lignicolum aspartic proteinase B; SLB

A second endopeptidase from Scytalidium lignicolum (see scytalidopepsin A) that is insensitive to **Comments:** 

> pepstatin and methyl 2-diazoacetamidohexanoate. 1,2-Epoxy-3-(p-nitrophenoxy)propane reacts with Glu<sup>53</sup>, which replaces one of the aspartic residues at the active centre. One of the smallest aspartic endopeptidases active as the monomer, with  $M_r$  22,000. Similarly inhibitor-resistant endopeptidases are found in the basidiomycetes Lentinus edodes [2902] and Ganoderma lucidum [2903], and in Polyporus tulipiferae [1486], a second endopeptidase distinct from polyporopepsin, but these are of typi-

cal aspartic endopeptidase size,  $M_r$  about 36,000. Type example of peptidase family G1.

**References:** [2902, 1780, 2903, 1486, 2990]

[EC 3.4.23.32 created 1992]

[3.4.23.33 Transferred entry. xanthomonapepsin. Now EC 3.4.21.101, xanthomonalisin]

[EC 3.4.23.33 created 1992, deleted 2001]

EC 3.4.23.34

Accepted name: cathepsin E

> **Reaction:** Similar to cathepsin D, but slightly broader specificity

Other name(s): slow-moving proteinase; erythrocyte membrane aspartic proteinase; SMP; erythrocyte membrane as-

partic proteinase; EMAP; non-pepsin proteinase; cathepsin D-like acid proteinase; cathepsin E-like

acid proteinase; cathepsin D-type proteinase

**Comments:** Found in stomach, spleen, erythrocyte membrane; not lysosomal. Pro-cathepsin E is an 86 kDa

disulfide-linked dimer; activation or reduction produces monomer. In peptidase family A1 (pepsin

A family)

[1599, 3290, 1349, 108] **References:** 

[EC 3.4.23.34 created 1992]

EC 3.4.23.35

Accepted name: barrierpepsin

> Selective cleavage of -Leu<sup>6</sup>—Lys- bond in the pheromone  $\alpha$ -mating factor Reaction:

Other name(s): barrier proteinase; Bar proteinase

**Comments:** A secreted endopeptidase known from baker's yeast (Saccharomyces cerevisiae). In peptidase family

A1 (pepsin A family)

**References:** [1766, 1765]

[EC 3.4.23.35 created 1993]

EC 3.4.23.36

Accepted name: signal peptidase II **Reaction:** Release of signal peptides from bacterial membrane prolipoproteins including murein prolipoprotein.

Hydrolyses -Xaa-Yaa-Zaa—(S,diacylglyceryl)Cys-, in which Xaa is hydrophobic (preferably Leu),

and Yaa (Ala or Ser) and Zaa (Gly or Ala) have small, neutral sidechains

Other name(s): premurein-leader peptidase; prolipoprotein signal peptidase; leader peptidase II; premurein leader

proteinase; leader peptidase II

**Comments:** An 18-kDa enzyme present in bacterial inner membranes. Inhibited by pepstatin and the antibiotic

globomycin. Type example of peptidase family A8.

**References:** [596, 3332, 2519]

[EC 3.4.23.36 created 1984 as EC 3.4.99.35, transferred 1995 to EC 3.4.23.36]

[3.4.23.37 Transferred entry. pseudomonapepsin. Now EC 3.4.21.100, pseudomonalisin]

[EC 3.4.23.37 created 1995]

EC 3.4.23.38

Accepted name: plasmepsin I

**Reaction:** Hydrolysis of the -Phe<sup>33</sup>—Leu- bond in the  $\alpha$ -chain of hemoglobin, leading to denaturation of the

molecule

Other name(s): aspartic hemoglobinase I; PFAPG; malaria aspartic hemoglobinase

Comments: Known from the malaria organism, *Plasmodium*. About 37 kDa. In peptidase family A1 (pepsin A

family), closest to cathepsin D and renin in structure. Inhibited by pepstatin. Formerly included in EC

3.4.23.6

**References:** [944, 803, 933]

[EC 3.4.23.38 created 1995]

EC 3.4.23.39

Accepted name: plasmepsin II

**Reaction:** Hydrolysis of the bonds linking certain hydrophobic residues in hemoglobin or globin. Also cleaves

the small molecule substrates such as Ala-Leu-Glu-Arg-Thr-Phe—Phe(NO<sub>2</sub>)-Ser-Phe-Pro-Thr [3]

Other name(s): aspartic hemoglobinase II; PFAPD

Comments: Known from the malaria organism, *Plasmodium*. About 37 kDa. In peptidase family A1 (pepsin A

family), and is 73% identical in sequence to plasmepsin I. Inhibited by pepstatin. Formerly included

in EC 3.4.23.6

**References:** [532, 933, 1141]

[EC 3.4.23.39 created 1995]

EC 3.4.23.40

**Accepted name:** phytepsin

**Reaction:** Prefers hydrophobic residues Phe, Val, Ile, Leu, and Ala at P1 and P1', but also cleaves -Phe—Asp-

and -Asp—Asp- bonds in 2S albumin from plant seeds

Comments: Known particularly from barley grain, but present in other plants also. In peptidase family A1 (pepsin

A family), but structurally distinct in containing an internal region of about 100 amino acids not gen-

erally present in the family

**References:** [2478, 1423, 80, 1424]

[EC 3.4.23.40 created 1997]

EC 3.4.23.41

Accepted name: yapsin 1

**Reaction:** Hydrolyses various precursor proteins with Arg or Lys in P1, and commonly Arg or Lys also in P2.

The P3 amino acid is usually non-polar, but otherwise additional basic amino acids are favourable in

both non-prime and prime positions

**Other name(s):** yeast aspartic protease 3; Yap3 gene product (*Saccharomyces cerevisiae*)

**Comments:** In peptidase family A1 of pepsin, and weakly inhibited by pepstatin. Can partially substitute for kexin

in a deficient strain of yeast. The homologous product of the Mkc7 gene (Saccharomyces cerevisiae)

has similar catalytic activity and has been termed yapsin 2 [861]

**References:** [382, 861, 2194]

[EC 3.4.23.41 created 2000]

#### EC 3.4.23.42

Accepted name: thermopsin

**Reaction:** Similar in specificity to pepsin A preferring bulky hydrophobic amino acids in P1 and P1'

**Comments:** From the thermophilic archaeon *Sulfolobus acidocaldarius*. Maximally active at pH 2 and 90 °C.

Weakly inhibited by pepstatin but shows no sequence similarity to pepsin. Type example of peptidase

family A5.

**References:** [1695]

[EC 3.4.23.42 created 1992 as EC 3.4.99.43, transferred 2000 to EC 3.4.23.42]

#### EC 3.4.23.43

**Accepted name:** prepilin peptidase

**Reaction:** Typically cleaves a -Gly—Phe- bond to release an N-terminal, basic peptide of 5-8 residues from

type IV prepilin, and then N-methylates the new N-terminal amino group, the methyl donor being S-

adenosyl-L-methionine

**Comments:** Many species of bacteria carry pili on their cell surfaces. These are virulence determinants in

pathogenic strains, and are assembled biosynthetically from type IV prepilin subunits. Before assembly, the prepilin molecules require proteolytic processing, which is done by the prepilin peptidase. Prepilin peptidase and its homologues play a central role not only in type IV pilus biogenesis but also in transport of macromolecules across cell membranes. Although both peptide-bond hydrolysis and *N*-methylation are catalysed by the same molecule, the methylation can be inhibited without affecting peptidase activity, and it is believed that the enzyme has two separate catalytic sites. Type example of

peptidase family A24.

**References:** [1737, 1598]

[EC 3.4.23.43 created 2001]

### EC 3.4.23.44

Accepted name: nodavirus endopeptidase

**Reaction:** Hydrolysis of an asparaginyl bond involved in the maturation of the structural protein of the virus,

typically -Asn—Ala- or -Asn—Phe-

Other name(s): Black Beetle virus endopeptidase; Flock House virus endopeptidase

**Comments:** A single aspartic residue is critical for activity, and inhibition by EDTA indicates that a metal ion is

also important. The enzyme is known from several nodaviruses that are pathogens of insects. Type example of peptidase family A6, and structurally related to the tetravirus endopeptidase in family A21,

although in that family, the catalytic residue is thought to be Glu.

**References:** [3345, 1326]

[EC 3.4.23.44 created 2001]

#### EC 3.4.23.45

Accepted name: memapsin 1

Reaction: Broad endopeptidase specificity. Cleaves Glu-Val-Asn-Leu—Asp-Ala-Glu-Phe in the Swedish vari-

ant of Alzheimer's amyloid precursor protein

Other name(s): β-secretase; β-site Alzheimer's amyloid precursor protein cleaving enzyme 2 (BACE2); ASP1; Down

region aspartic protease

**Comments:** Can cleave  $\beta$ -amyloid precursor protein to form the amyloidogenic  $\beta$ -peptide that is implicated in

the pathology of Alzheimer's disease, but is not significantly expressed in human brain. In peptidase

family A1, but is atypical in containing a C-terminal membrane-spanning domain.

**References:** [3001]

[EC 3.4.23.45 created 2003]

EC 3.4.23.46

**Accepted name:** memapsin 2

Reaction: Broad endopeptidase specificity. Cleaves Glu-Val-Asn-Leu-Asp-Ala-Glu-Phe in the Swedish vari-

ant of Alzheimer's amyloid precursor protein

**Other name(s):** β-secretase; β-site Alzheimer's amyloid precursor protein cleaving enzyme 1 (BACE1)

Comments: Suggested to be the major " $\beta$ -secretase" responsible for the cleavage of the  $\beta$ -amyloid precursor pro-

tein to form the amyloidogenic  $\beta$ -peptide that is implicated in the pathology of Alzheimer's disease. In peptidase family A1 but is atypical in containing a C-terminal membrane-spanning domain.

**References:** [3002, 1168]

[EC 3.4.23.46 created 2003]

EC 3.4.23.47

**Accepted name:** HIV-2 retropepsin

**Reaction:** Endopeptidase for which the P1 residue is preferably hydrophobic

Comments: In peptidase family A2. Responsible for the post-translational processing of the human immunodefi-

ciency virus polyprotein.

**References:** [2957, 419]

[EC 3.4.23.47 created 2003]

EC 3.4.23.48

**Accepted name:** plasminogen activator Pla

**Reaction:** Converts human Glu-plasminogen to plasmin by cleaving the Arg<sup>560</sup>—Val peptide bond that is also

hydrolysed by the mammalian u-plasminogen activator and t-plasminogen activator. Also cleaves

arginyl bonds in other proteins

**Comments:** In peptidase family A26. From the bacterium *Yersinia pestis* that causes plague.

**References:** [1544]

[EC 3.4.23.48 created 2003]

EC 3.4.23.49

Accepted name: omptin

**Reaction:** Has a virtual requirement for Arg in the P1 position and a slightly less stringent preference for this

residue in the P1' position, which can also contain Lys, Gly or Val.

**Other name(s):** protease VII; protease A; gene *ompT* proteins; *ompT* protease; protein a; Pla; OmpT

**Comments:** A product of the *ompT* gene of *Escherichia coli*, and associated with the outer membrane. Omptin

shows a preference for cleavage between consecutive basic amino acids, but is capable of cleavage

when P1' is a non-basic residue [3049, 1862]. Belongs in peptidase family A26.

**References:** [989, 2794, 1045, 571, 3049, 1528, 1862]

[EC 3.4.23.49 created 1993 as EC 3.4.21.87, transferred 2006 to EC 3.4.23.49]

### EC 3.4.23.50

**Accepted name:** human endogenous retrovirus K endopeptidase

Reaction: Processing at the authentic HIV-1 PR recognition site and release of the mature p17 matrix and the

p24 capsid protein, as a result of the cleavage of the -SQNY—PIVQ- cleavage site.

Other name(s): human endogenous retrovirus K10 endopeptidase; endogenous retrovirus HERV-K10 putative

protease; human endogenous retrovirus K retropepsin; HERV K10 endopeptidase; HERV K10 retropepsin; HERV-K PR; HERV-K protease; HERV-K113 protease; human endogenous retrovirus

K113 protease; human retrovirus K10 retropepsin

**Comments:** In peptidase family A2.

**References:** [2953]

[EC 3.4.23.50 created 2009]

### EC 3.4.23.51

**Accepted name:** HycI peptidase

**Reaction:** This enzyme specifically removes a 32-amino acid peptide from the C-terminus of the precursor of

the large subunit of hydrogenase 3 in *Escherichia coli* by cleavage at the C-terminal side of Arg<sup>537</sup>.

Other name(s): HycI; HycE processing protein

**Comments:** The reaction requires nickel to be bound to the precursor of the large subunit of hydrogenase 3. The

endopeptidase uses the metal in the large subunit of [NiFe]-hydrogenases as a recognition motif

[2907]. In peptidase family A31.

**References:** [2907, 3257]

[EC 3.4.23.51 created 2009]

### EC 3.4.23.52

**Accepted name:** preflagellin peptidase

**Reaction:** Cleaves the signal peptide of 3 to 12 amino acids from the N-terminal of preflagellin, usually at Arg-

Gly— or Lys-Gly—, to release flagellin.

Other name(s): FlaK

**Comments:** An aspartic peptidase from Archaea but not bacteria. In peptidase family A24 (type IV prepilin pepti-

dase family).

**References:** [139, 2077, 1198]

[EC 3.4.23.52 created 2011]

### EC 3.4.24 Metalloendopeptidases

# EC 3.4.24.1

Accepted name: atrolysin A

**Reaction:** Cleavage of Asn<sup>3</sup>—Gln, His<sup>5</sup>—Leu, His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu and Tyr<sup>16</sup>—Leu in insulin B chain;

removes C-terminal Leu from small peptides

Other name(s): Crotalus atrox metalloendopeptidase a; hemorrhagic toxin a; Crotalus atrox α-proteinase; Crotalus

atrox proteinase; bothropasin

**Comments:** A hemorrhagic endopeptidase of 68 kDa, one of six hemorrhagic toxins in the venom of western di-

amondback rattlesnake. The 60 kDa hemorrhagic toxin 1 of *Crotalus ruber ruber* shows identical specificity [1966]. In peptidase family M12 (astacin family). Related metalloendopeptidases from rattlesnake venoms are EC 3.4.24.41 (atrolysin B), EC 3.4.24.42 (atrolysin C), EC 3.4.24.43 (atroxase), EC 3.4.24.44 (atrolysin E), EC 3.4.24.45 (atrolysin F), EC 3.4.24.46 (adamalysin), EC 3.4.24.47 (hor-

rilysin), and EC 3.4.24.48 (ruberlysin)

**References:** [224, 1966, 223, 222]

[EC 3.4.24.1 created 1972, modified 1986]

### [3.4.24.2 Deleted entry. Sepia proteinase]

[EC 3.4.24.2 created 1972, deleted 1992]

### EC 3.4.24.3

Accepted name: microbial collagenase

**Reaction:** Digestion of native collagen in the triple helical region at —Gly bonds. With synthetic peptides, a

preference is shown for Gly at P3 and P1', Pro and Ala at P2 and P2', and hydroxyproline, Ala or Arg

at P3'

Other name(s): Clostridium histolyticum collagenase; clostridiopeptidase A; collagenase A; collagenase I; Achro-

*mobacter* iophagus collagenase; collagenase; aspergillopeptidase C; nucleolysin; azocollase; metallocollagenase; soycollagestin; *Clostridium histolyticum* proteinase A; clostridiopeptidase II; MMP-8; clostridiopeptidase I; collagen peptidase; collagen protease; collagenase MMP-1; metalloproteinase-1; kollaza; matrix metalloproteinase-1; MMP-1; matrix metalloproteinase-8; matirx metalloproteinase-

18; interstitial collagenase

**Comments:** Six species of metalloendopeptidase acting on native collagen can be isolated from the medium of

Clostridium histolyticum. Class I has forms  $\alpha$  (68 kDa),  $\beta$  (115 kDa) and  $\gamma$  (79 kDa); class II has  $\delta$  (100 kDa),  $\epsilon$  (110 kDa) and  $\zeta$  (125 kDa). The two classes are immunologically crossreactive, but have significantly different sequences, and different specificities such that their actions on collagen are complementary. The enzymes also act as peptidyl-tripeptidases. Variants of the enzyme have been purified from Bacillus cereus [1783], Empedobacter collagenolyticum [1582], Pseudomonas marinoglutinosa [1041], and species of Vibrio, Vibrio B-30 (ATCC 21250) [1890] and V. alginolyticus (previously Achromobacter iophagus) [1101, 2944]. Also known from Streptomyces sp. [702]. The Vibrio

enzyme is the type example of peptidase family M9.

**References:** [1041, 1890, 1101, 1582, 259, 260, 3132, 2944, 702, 1783]

[EC 3.4.24.3 created 1961 as EC 3.4.4.19, transferred 1972 to EC 3.4.24.3 (EC 3.4.24.8 created 1978, incorporated 1992, EC 3.4.99.5 created 1972, incorporated 1978)]

[3.4.24.4 Transferred entry. now EC 3.4.24.40 serralysin]

[EC 3.4.24.4 created 1972 [EC 3.4.99.13 and EC 3.4.99.22 both created 1972, incorporated 1978], deleted 1992]

[3.4.24.5 Deleted entry. lens neutral proteinase. Now included with EC 3.4.22.53 (calpain-2) and EC 3.4.25.1 (proteasome endopeptidase complex)]

[EC 3.4.24.5 created 1978, deleted 1989]

# EC 3.4.24.6

Accepted name: leucolysin

**Reaction:** Cleavage of Phe<sup>1</sup>+Val, His<sup>5</sup>+Leu, Ala<sup>14</sup>+Leu, Gly<sup>20</sup>+Glu, Gly<sup>23</sup>+Phe and Phe<sup>24</sup>+Phe bonds

in insulin B chain as well as N-blocked dipeptides

Other name(s): Leucostoma neutral proteinase; Leucostoma peptidase A

**Comments:** From the venom of the western cottonmouth moccasin snake (*Agkistrodon piscivorus leucostoma*).

**References:** [3082, 2740]

[EC 3.4.24.6 created 1978]

### EC 3.4.24.7

Accepted name: interstitial collagenase

**Reaction:** Cleavage of the triple helix of collagen at about three-quarters of the length of the molecule from the

N-terminus, at Gly<sup>775</sup>—Ile in the  $\alpha$ 1(I) chain. Cleaves synthetic substrates and  $\alpha$ -macroglobulins at

bonds where P1' is a hydrophobic residue

**Other name(s):** vertebrate collagenase; matrix metalloproteinase 1

**Comments:** The enzyme takes its name from substrates of the interstitial collagen group - types I, II and III, all of

which are cleaved in the helical domain. However,  $\alpha$ -macroglobulins are cleaved much more rapidly. The enzyme is widely distributed in vertebrate animals. Type example of peptidase family M10

**References:** [945, 217, 766, 2729]

[EC 3.4.24.7 created 1978]

[3.4.24.8 Transferred entry. Achromobacter iophagus collagenase. Now EC 3.4.24.3, microbial collagenase]

[EC 3.4.24.8 created 1978, deleted 1992]

[3.4.24.9 Deleted entry. Trichophyton schoenleinii collagenase]

[EC 3.4.24.9 created 1978, deleted 1992]

[3.4.24.10 Deleted entry. Trichophyton mentagrophytes keratinase]

[EC 3.4.24.10 created 1972 as EC 3.4.99.12, transferred 1978 to EC 3.4.24.10, deleted 1992]

EC 3.4.24.11

Accepted name: neprilysin

Reaction: Preferential cleavage of polypeptides between hydrophobic residues, particularly with Phe or Tyr at

P1'

Other name(s): neutral endopeptidase; endopeptidase 24.11; kidney-brush-border neutral peptidase; enkephalinase

(misleading); endopeptidase-2; CALLA (common acute lymphoblastic leukemia-associated) antigens; CALLA antigen; endopeptidase; membrane metalloendopeptidase; kidney-brush-border neutral endopeptidase; kidney-brush-border neutral proteinase; endopeptidase-2; CALLA glycoprotein; CALLA; common acute lymphoblastic leukemia antigen; CALLA glycoproteins; common acute lymphoblastic leukemia-associated antigens; neutral metallendopeptidase; membrane metalloendopeptidase; NEP; neutral endopeptidase 24.11; CD10; neutral endopeptidase; acute lymphoblastic leukemia

antigen

**Comments:** A membrane-bound glycoprotein widely distributed in animal tissues. Inhibited by phosphoramidon

and thiorphan. Common acute lymphoblastic leukemia antigen (CALLA). Type example of peptidase

family M13

**References:** [1836, 1787, 1652, 711]

[EC 3.4.24.11 created 1978, modified 1989]

EC 3.4.24.12

Accepted name: envelysin

**Reaction:** Hydrolysis of proteins of the fertilization envelope and dimethylcasein

Other name(s): sea-urchin-hatching proteinase; hatching enzyme; chorionase; chorion-digesting proteinase; chy-

mostrypsin; sea urchin embryo hatching enzyme

**Comments:** A glycoprotein from various members of the class *Echinoidea*. Extracellular enzyme requiring Ca<sup>2+</sup>.

In peptidase family M10 (interstitial collagenase family)

**References:** [160, 1649, 1650, 2108]

[EC 3.4.24.12 created 1978]

EC 3.4.24.13

Accepted name: IgA-specific metalloendopeptidase

**Reaction:** Cleavage of Pro—Thr bond in the hinge region of the heavy chain of human IgA

Other name(s): immunoglobulin A<sub>1</sub> proteinase; IgA protease; IgA1-specific proteinase; IgA1 protease; IgA<sub>1</sub> pro-

teinase

**Comments:** A 190 kDa enzyme found in several pathogenic species of *Streptococcus* such as *sanguis* and *pneu-*

moniae. Type example of peptidase family M26. There is also an IgA-specific prolyl endopeptidase of

the serine-type (see EC 3.4.21.72, IgA-specific serine endopeptidase)

**References:** [1515, 920, 919]

### [EC 3.4.24.13 created 1984]

EC 3.4.24.14

**Accepted name:** procollagen *N*-endopeptidase

**Reaction:** Cleaves the *N*-propeptide of collagen chain  $\alpha 1(I)$  at Pro—Gln and of  $\alpha 1(II)$  and  $\alpha 2(I)$  at Ala—Gln **Other name(s):** procollagen N-terminal peptidase; procollagen aminopeptidase; aminoprocollagen peptidase;

aminoterminal procollagen peptidase; procollagen aminoterminal protease; procollagen N-terminal

proteinase; type I/II procollagen N-proteinase; type III procollagen

Comments: Removes the propeptides of type I and II collagens prior to fibril assembly. Does not act on type III

collagen. In peptidase family M12 (astacin family)

**References:** [1496, 1160]

[EC 3.4.24.14 created 1984]

EC 3.4.24.15

**Accepted name:** thimet oligopeptidase

**Reaction:** Preferential cleavage of bonds with hydrophobic residues at P1, P2 and P3' and a small residue at P1'

in substrates of 5-15 residues

Other name(s): Pz-peptidase; soluble metalloendopeptidase; endo-oligopeptidase A; tissue-endopeptidase degrading

collagenase-synthetic-substrate

**Comments:** Thiol compounds activate at low concentrations. Type example of peptidase family M3.

**References:** [452, 2200, 157, 2279, 2922]

[EC 3.4.24.15 created 1984 (EC 3.4.22.19 created 1989 and EC 3.4.99.31 created 1978 both incorporated 1992)]

EC 3.4.24.16

Accepted name: neurolysin

**Reaction:** Preferential cleavage in neurotensin: Pro<sup>10</sup>—Tyr

Other name(s): neurotensin endopeptidase; endopeptidase 24.16; endo-oligopeptidase B (proline-endopeptidase)

Comments: No absolute requirement for a prolyl bond: the enzyme acts on some peptides, such as dynorphin 1-8,

that do not contain proline, and does not act on some others that do. In peptidase family M3 (thimet

oligopeptidase family)

**References:** [408, 140, 407]

[EC 3.4.24.16 created 1989]

EC 3.4.24.17

**Accepted name:** stromelysin 1

**Reaction:** Preferential cleavage where P1', P2' and P3' are hydrophobic residues

Other name(s): matrix metalloproteinase 3; proteoglycanase; collagenase activating protein; procollagenase activator;

transin; MMP-3; neutral proteoglycanase; stromelysin; collagen-activating protein

Comments: An extracellular endopeptidase of vertebrate tissues homologous with interstitial collagenase. Di-

gests proteoglycan, fibronectin, collagen types III, IV, V, IX, and activates procollagenase. In pepti-

dase family M10 (interstitial collagenase family)

**References:** [433, 2181, 623, 700]

[EC 3.4.24.17 created 1990]

EC 3.4.24.18

Accepted name: meprin A

**Reaction:** Hydrolysis of protein and peptide substrates preferentially on carboxyl side of hydrophobic residues

Other name(s): endopeptidase-2; meprin-a; meprin; N-benzoyl-L-tyrosyl-p-aminobenzoic acid hydrolase; PABA-

peptide hydrolase; PPH

Comments: A membrane-bound metalloendopeptidase of rat and mouse kidney and intestinal brush borders, and

salivary ducts. Differences from neprilysin (EC 3.4.24.11 (astacin family). Formerly included in EC

3.4.24.11

**References:** [211, 337, 2763, 2764, 147]

[EC 3.4.24.18 created 1992]

EC 3.4.24.19

Accepted name: procollagen C-endopeptidase

**Reaction:** Cleavage of the C-terminal propeptide at Ala—Asp in type I and II procollagens and at Arg—Asp in

type III

Other name(s): procollagen C-terminal proteinase; carboxyprocollagen peptidase; procollagen C-terminal peptidase;

procollagen C-proteinase; procollagen C-terminal proteinase; procollagen carboxypeptidase; procolla-

gen carboxy-terminal proteinase; procollagen peptidase

**Comments:** A 100 kDa endopeptidase the activity of which is increased by Ca<sup>2+</sup> and by an enhancer glycoprotein.

In peptidase family M12 (astacin family)

**References:** [1161, 1426]

[EC 3.4.24.19 created 1992]

EC 3.4.24.20

Accepted name: peptidyl-Lys metalloendopeptidase

**Reaction:** Preferential cleavage in proteins: -Xaa—Lys- (in which Xaa may be Pro) **Other name(s):** Armillaria mellea neutral proteinase; peptidyllysine metalloproteinase

Comments: From the honey fungus Armillaria mellea. In peptidase family M35 (deuterolysin family).

**References:** [2483, 1661]

[EC 3.4.24.20 created 1978 as EC 3.4.99.32, transferred 1992 to EC 3.4.24.20 (EC 3.4.99.30 created 1978, incorporated 1992)]

EC 3.4.24.21

Accepted name: astacin

**Reaction:** Hydrolysis of peptide bonds in substrates containing five or more amino acids, preferentially with Ala

in P1', and Pro in P2'

**Other name(s):** Astacus proteinase; crayfish small-molecule proteinase

**Comments:** A 22.6 kDa digestive endopeptidase from the cardia of the crayfish *Astacus fluviatilis*. Type example

of peptidase family M12.

**References:** [1529, 2924, 2776, 2775]

[EC 3.4.24.21 created 1972 as EC 3.4.99.6, transferred 1992 to EC 3.4.24.21]

EC 3.4.24.22

**Accepted name:** stromelysin 2

**Reaction:** Similar to stromelysin 1, but action on collagen types III, IV and V is weak

**Other name(s):** matrix metalloproteinase 10; transin 2; proteoglycanase 2

**Comments:** In peptidase family M10 (interstitial collagenase family). Digests gelatin types I, III, IV, V, fibronectin

and proteoglycan

**References:** [282, 1994, 2080]

[EC 3.4.24.22 created 1992]

EC 3.4.24.23

Accepted name: matrilysin

**Reaction:** Cleavage of Ala<sup>14</sup>—Leu and Tyr<sup>16</sup>—Leu in B chain of insulin. No action on collagen types I, II, IV,

V. Cleaves gelatin chain  $\alpha 2(I)$ ;  $\alpha 1(I)$ 

Other name(s): matrin; uterine metalloendopeptidase; matrix metalloproteinase 7; putative (or punctuated)

metalloproteinase-1; matrix metalloproteinase pump 1; MMP 7; PUMP-1 proteinase; PUMP; met-

alloproteinase pump-1; putative metalloproteinase; MMP

**Comments:** Found in rat uterus; at 19 kDa, the smallest member of peptidase family M10 (interstitial collagenase

family). Similar in specificity to stromelysin, but more active on azocoll

**References:** [1994, 3205, 2341, 1938]

[EC 3.4.24.23 created 1992]

### EC 3.4.24.24

Accepted name: gelatinase A

Reaction: Cleavage of gelatin type I and collagen types IV, V, VII, X. Cleaves the collagen-like sequence Pro-

Gln-Gly—Ile-Ala-Gly-Gln

Other name(s): 72-kDa gelatinase; matrix metalloproteinase 2; type IV collagenase (ambiguous); 3/4 collagenase (ob-

solete); matrix metalloproteinase 5 (obsolete); 72 kDa gelatinase type A; collagenase IV (ambiguous); collagenase type IV (ambiguous); MMP 2; type IV collagen metalloproteinase (ambiguous); type IV

collagenase/gelatinase (ambiguous)

Comments: A secreted endopeptidase in peptidase family M10 (interstitial collagenase family), but possessing an

additional fibronectin-like domain

**References:** [2009, 468, 2182]

[EC 3.4.24.24 created 1992]

### EC 3.4.24.25

Accepted name: vibriolysin

**Reaction:** Preferential cleavage of bonds with bulky hydrophobic groups in P2 and P1'. Phe at P1' is the most

favoured residue, which distinguished this enzyme from thermolysin

**Other name(s):** Aeromonas proteolytica neutral proteinase; aeromonolysin

**Comments:** Thermostable enzyme from *Vibrio proteolyticus* (formerly *Aeromonas proteolytica*). Specificity re-

lated to, but distinct from, those of thermolysin and bacillolysin [1165]. A zinc metallopeptidase in

family M4 (thermolysin family). Formerly included in EC 3.4.24.4

**References:** [1165, 3189, 177, 3188, 544]

[EC 3.4.24.25 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.25, modified 1997]

### EC 3.4.24.26

Accepted name: pseudolysin

Reaction: Hydrolysis of proteins including elastin, collagen types III and IV, fibronectin and immunoglobulin

A, generally with bulky hydrophobic group at P1'. Insulin B chain cleavage pattern identical to that of

thermolysin, but specificity differs in other respects

Other name(s): Pseudomonas elastase; Pseudomonas aeruginosa neutral metalloproteinase

Comments: In peptidase family M4 (thermolysin family). From the pathogenic bacteria *Pseudomonas aeruginosa* 

and Legionella pneumophila, and causes tissue damage.

**References:** [1971, 2094, 652, 210, 231]

[EC 3.4.24.26 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.26]

#### EC 3.4.24.27

Accepted name: thermolysin

**Reaction:** Preferential cleavage: —Leu ; —Phe

**Other name(s):** Bacillus thermoproteolyticus neutral proteinase; thermoase; thermoase Y10; TLN

Comments: A thermostable extracellular metalloendopeptidase containing four calcium ions. Enzymes that

may be species variants of thermolysin are reported from *Micrococcus caseolyticus* [588] and *Aspergillus oryzae* [1970]. Type example of peptidase family M4. Closely related but distinct enzymes

are aeromonolysin, pseudolysin, bacillolysin, aureolysin and mycolysin

**References:** [2173, 1973, 1610, 588, 1970, 2923, 1846]

[EC 3.4.24.27 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.27]

EC 3.4.24.28

Accepted name: bacillolysin

**Reaction:** Similar, but not identical, to that of thermolysin

Other name(s): Bacillus metalloendopeptidase; Bacillus subtilis neutral proteinase; anilozyme P 10; Bacillus metallo-

proteinase; Bacillus neutral proteinase; megateriopeptidase

**Comments:** Variants of this enzyme have been found in species of *Bacillus* including *B. subtilis* [1973, 3259], *B.* 

amyloliquefaciens [3054], B. megaterium (megateriopeptidase, [1917]), B. mesentericus [2777], B. cereus [3,8,9] and B. stearothermophilus [2833]. In peptidase family M4 (thermolysin family). For-

merly included in EC 3.4.24.4

**References:** [1973, 1917, 749, 1165, 3054, 3259, 2833, 2653, 2249, 2777]

[EC 3.4.24.28 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.28]

EC 3.4.24.29

Accepted name: aureolysin

**Reaction:** Cleavage of insulin B chain with specificity similar to that of thermolysin, preferring hydrophobic P1'

residue. Activates the glutamyl endopeptidase (EC 3.4.21.19) of Staphylococcus aureus

Other name(s): Staphylococcus aureus neutral proteinase; Staphylococcus aureus neutral protease

Comments: A metalloenzyme from S. aureus earlier confused with staphylokinase (a non-enzymic activator of

plasminogen).

**References:** [79, 2494, 647, 2315]

[EC 3.4.24.29 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.29]

EC 3.4.24.30

Accepted name: coccolysin

**Reaction:** Preferential cleavage: —Leu, —Phe, —Tyr, —Ala

Other name(s): Streptococcus thermophilus intracellular proteinase; EM 19000

Comments: A 30 kDa endopeptidase found intracellularly in S. thermophilus [589] and S. diacetilactis [590] and

in the medium of S. faecalis [2699, 1784]. In peptidase family M4 (thermolysin family). Formerly

included in EC 3.4.24.4

**References:** [589, 590, 2699, 1784]

[EC 3.4.24.30 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.30]

EC 3.4.24.31

Accepted name: mycolysin

**Reaction:** Preferential cleavage of bonds with hydrophobic residues in P1'

Other name(s): pronase component; Streptomyces griseus neutral proteinase; actinase E; SGNPI

Comments: From Streptomyces griseus, S. naraensis, and S. cacaoi. Specificity similar to that of thermolysin,

but much more sensitive to inhibition by mercaptoacetyl-Phe-Leu. Little structural similarity to other bacterial metalloendopeptidases. Type example of peptidase family M5. Formerly included in EC

3.4.24.4

**References:** [1973, 1146, 246, 395]

### [EC 3.4.24.31 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.31]

EC 3.4.24.32

**Accepted name:** β-lytic metalloendopeptidase

Reaction: Cleavage of *N*-acetylmuramoyl—Ala, and of the insulin B chain at Gly<sup>23</sup>—Phe ¿ Val<sup>18</sup>—Cya

Other name(s): *Myxobacter* β-lytic proteinase; achromopeptidase component; β-lytic metalloproteinase; β-lytic pro-

tease; *Myxobacter*ium sorangium β-lytic proteinase; *Myxobacter*495 β-lytic proteinase

Comments: From Achromobacter lyticus and Lysobacter enzymogenes. Digests bacterial cell walls. Type example

of peptidase family M23.

**References:** [3171, 3170, 1666]

[EC 3.4.24.32 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.32]

EC 3.4.24.33

Accepted name: peptidyl-Asp metalloendopeptidase

**Reaction:** Cleavage of Xaa—Asp, Xaa—Glu and Xaa—cysteic acid bonds

**Other name(s):** endoproteinase Asp-N; peptidyl-Asp metalloproteinase

Comments: A metalloenzyme isolated from *Pseudomonas fragi*. Useful in protein sequencing applications be-

cause of its limited specificity. In peptidase family M72.

**References:** [2313, 649, 1250]

[EC 3.4.24.33 created 1992]

EC 3.4.24.34

**Accepted name:** neutrophil collagenase

**Reaction:** Cleavage of interstitial collagens in the triple helical domain. Unlike EC 3.4.24.7, interstitial collage-

nase, this enzyme cleaves type III collagen more slowly than type I

Other name(s): matrix metalloproteinase 8; PMNL collagenase; MMP-8

**Comments:** Similar to interstitial collagenase in specificity, but the product of a different gene and highly glyco-

sylated. Stored in the specific granules of neutrophil leukocytes. In peptidase family M10 (interstitial

collagenase family). Formerly included in EC 3.4.24.7

**References:** [1072, 1073, 1480]

[EC 3.4.24.34 created 1992]

EC 3.4.24.35

Accepted name: gelatinase B

**Reaction:** Cleavage of gelatin types I and V and collagen types IV and V

**Other name(s):** 92-kDa gelatinase; matrix metalloproteinase 9; type V collagenase; 92-kDa type IV collagenase;

macrophage gelatinase; 95 kDa type IV collagenase/gelatinase; collagenase IV (ambiguous); collagenase type IV (ambiguous); gelatinase MMP 9; MMP 9; type IV collagen metalloproteinase (am-

biguous)

Comments: Similar to gelatinase A, but possesses a further domain . In peptidase family M10 (interstitial collage-

nase family)

**References:** [1136, 3185, 1778]

[EC 3.4.24.35 created 1992]

EC 3.4.24.36

Accepted name: leishmanolysin

**Reaction:** Preference for hydrophobic residues at P1 and P1' and basic residues at P2' and P3'. A model non-

apeptide is cleaved at -Ala-Tyr-Leu-Lys-Lys-

Other name(s): promastigote surface endopeptidase; glycoprotein gp63; Leishmania metalloproteinase; surface acid

proteinase; promastigote surface protease

**Comments:** A membrane-bound glycoprotein found on the promastigote of various species of *Leishmania* proto-

zoans. Contains consensus sequence for a zinc-binding site; Z-Tyr-Leu-NHOH is a strong inhibitor. The enzyme can activate its proenzyme by cleavage of the Val Val bond. An acid pH optimum is

found with certain protein substrates. Type example of peptidase family M8

**References:** [343, 271, 405, 272]

[EC 3.4.24.36 created 1992]

EC 3.4.24.37

Accepted name: saccharolysin

**Reaction:** Cleavage of Pro—Phe and Ala—Ala bonds

Other name(s): proteinase yscD; yeast cysteine proteinase D (Misleading); Saccharomyces cerevisiae proteinase yscD

Comments: An 83 kDa cytoplasmic thiol-dependent metalloendopeptidase from Saccharomyces cerevisiae. In

peptidase family M3 (thimet oligopeptidase family).

**References:** [9, 875]

[EC 3.4.24.37 created 1989 as EC 3.4.22.22, transferred 1992 to EC 3.4.24.37]

EC 3.4.24.38

Accepted name: gametolysin

**Reaction:** Cleavage of the proline- and hydroxyproline-rich proteins of the *Chlamydomonas* cell wall; also

cleaves azocasein, gelatin and Leu-Trp-Met-Arg-Phe-Ala

Other name(s): autolysin, Chlamydomonas cell wall degrading protease; lysin; Chlamydomonas reinhardtii metallo-

proteinase; gamete lytic enzyme; gamete autolysin

Comments: A glycoprotein found in the periplasmic space of Chlamydomonas reinhardtii gametes in a 62 kDa

inactive form; decreased to 60 kDa upon activation. A zinc enzyme, inhibited by phosphoramidon, but

also thiol activated. Type example of peptidase family M11

**References:** [1294, 323, 1837]

[EC 3.4.24.38 created 1992, modified 2000]

EC 3.4.24.39

Accepted name: deuterolysin

**Reaction:** Cleavage of the proline- and hydroxyproline-rich proteins of the *Chlamydomonas* cell wall; also

cleaves azocasein, gelatin and Leu-Trp-Met-Arg-Phe-Ala

Other name(s): Penicillium roqueforti protease II; microbial neutral proteinase II; acid metalloproteinase; neutral pro-

teinase II; Penicillium roqueforti metalloproteinase

Comments: Proteolytic activity found in Penicillium roqueforti [984], P. caseicolum [984], Aspergillus sojae

[2609] and *A. oryzae* [2033, 3026]. Optimum pH of 5 for digesting various proteins. Strong action on protamine and histones. Insensitive to phosphoramidon. About 20 kDa. A distinct *Aspergillus sojae* endopeptidase is larger and has a neutral pH optimum. Type example of peptidase family M35.

Formerly included in EC 3.4.24.4

**References:** [2033, 983, 2609, 984, 3026]

[EC 3.4.24.39 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.39]

EC 3.4.24.40

Accepted name: serralysin

**Reaction:** Preferential cleavage of bonds with hydrophobic residues in P1'

Other name(s): Pseudomonas aeruginosa alkaline proteinase; Escherichia freundii proteinase; Serratia marcescens

extracellular proteinase; Serratia marcescens metalloproteinase; Pseudomonas aeruginosa alk. pro-

tease; Serratia marcescens metalloprotease

A 50 kDa extracellular endopeptidase from Pseudomonas aeruginosa [1,2,6], Escherichia freundii **Comments:** 

> [2038], Serratia marcescens [4,5,6] and Erwinia chrysanthemi [527]. There is broad specificity in cleavage of the insulin B chain, with some species variations. The pH optimum for digesting various proteins is about 9 - 10. In peptidase family M10 (interstitial collagenase family). Formerly included

in EC 3.4.24.4

**References:** [1973, 1974, 2038, 565, 627, 2037, 527, 2189]

[EC 3.4.24.40 created 1972 as EC 3.4.24.4, part transferred 1992 to EC 3.4.24.40]

EC 3.4.24.41

Accepted name: atrolysin B

Cleavage of  $\mathrm{His}^5$ —Leu,  $\mathrm{His}^{10}$ —Leu,  $\mathrm{Ala}^{14}$ —Leu,  $\mathrm{Tyr}^{16}$ —Leu and  $\mathrm{Gly}^{23}$ —Phe of insulin B chain; identical to the cleavage of insulin B chain by atrolysin C. Also cleaves —Ser bonds in glucagon **Reaction:** 

Crotalus atrox metalloendopeptidase b; hemorrhagic toxin b; Ht-b Other name(s):

**Comments:** From the venom of the western diamondback rattlesnake (*Crotalus atrox*). In peptidase family M12

(astacin family)

**References:** [224, 223]

[EC 3.4.24.41 created 1992]

EC 3.4.24.42

Accepted name: atrolysin C

> Cleavage of His<sup>5</sup>—Leu, His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu, Tyr<sup>16</sup>—Leu and Gly<sup>23</sup>—Phe bonds in B chain of **Reaction:**

> > insulin. With small molecule substrates prefers hydrophobic residue at P2' and small residue such as

Other name(s): Crotalus atrox metalloendopeptidase c; hemorrhagic toxin c and d

A 24 kDa hemorrhagic endopeptidase from the venom of the western diamondback rattlesnake (Cro-**Comments:** 

talus atrox) that digests type IV collagen, and exists as two forms, c and d. Phosphoramidon inhibits in the 0.1 mM range. In peptidase family M12 (astacin family). Hemorrhagic toxin-2 of C. ruber ru-

ber has the same  $M_r$  and specificity and is a homologue [1966, 2864].

[224, 799, 221, 1966, 2618, 2864] **References:** 

[EC 3.4.24.42 created 1992]

EC 3.4.24.43

Accepted name:

Cleavage of His<sup>5</sup>—Leu, Ser<sup>9</sup>—His, His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu and Tyr<sup>16</sup>—Leu of insulin B chain **Reaction:** A nonhemorrhagic endopeptidase from the venom of the western diamondback rattlesnake (Crotalus **Comments:** 

atrox) that cleaves fibrinogen. In peptidase family M12 (astacin family)

**References:** [3193]

[EC 3.4.24.43 created 1992]

EC 3.4.24.44

atrolysin E Accepted name:

> Cleavage of Asn<sup>3</sup>—Gln, Ser<sup>9</sup>—His and Ala<sup>14</sup>—Leu bonds in insulin B chain and Tyr<sup>14</sup>—Gln and **Reaction:**

> > Thr<sup>8</sup>—Ser in A chain. Cleaves type IV collagen at Ala<sup>73</sup>—Gln in α1(IV) and at Gly<sup>7</sup>—Leu in

 $\alpha 2(IV)$ 

Crotalus atrox metalloendopeptidase e; hemorrhagic toxin e Other name(s):

A 25.7 kDa hemorrhagic endopeptidase from the venom of the western diamondback rattlesnake **Comments:** 

(Crotalus atrox) that digests basement membrane components, including the triple helix of type IV collagen. Such action is believed to contribute to the hemorrhagic property by weakening capillary

walls. In peptidase family M12 (astacin family)

**References:** [224, 220, 136]

[EC 3.4.24.44 created 1992]

EC 3.4.24.45

**Accepted name:** atrolysin F

**Reaction:** Cleavage of Val<sup>2</sup>—Asn, Gln<sup>4</sup>—His, Leu<sup>6</sup>—Cys, His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu and Tyr<sup>16</sup>—Leu bonds

in insulin B chain

Other name(s): Crotalus atrox metalloendopeptidase; hemorrhagic toxin f; Crotalus atrox metalloendopeptidase f

**Comments:** A 64 kDa hemorrhagic endopeptidase from the venom of the western diamondback rattlesnake

(Crotalus atrox) that digests the γ chain of fibrinogen. Immunologically distinct from EC 3.4.24.1,

atrolysin A.

**References:** [2085]

[EC 3.4.24.45 created 1992]

EC 3.4.24.46

Accepted name: adamalysin

**Reaction:** Cleavage of Phe<sup>1</sup>+Val, His<sup>5</sup>+Leu, His<sup>10</sup>+Leu, Ala<sup>14</sup>+Leu, Leu<sup>15</sup>+Tyr, and Tyr<sup>16</sup>+Leu of

insulin B chain

Other name(s): Crotalus adamanteus metalloendopeptidase; proteinase I and II; Crotalus adamanteus venom pro-

teinase II; adamalysin II

**Comments:** From the venom of the eastern diamondback rattlesnake (*Crotalus adamanteus*). Two isoenzymes of

approx. 24 kDa that inactivate  $\alpha_1$ -proteinase inhibitor by a single cleavage. In peptidase family M12

(astacin family)

**References:** [1565]

[EC 3.4.24.46 created 1992]

EC 3.4.24.47

Accepted name: horrilysin

**Reaction:** Cleavage of only the single bond Ala<sup>14</sup>—Leu in the insulin B chain, Ser<sup>12</sup>—Leu in the A chain, and

Ile—Gly, Pro—Ala, and Ser—Trp in melittin

Other name(s): Crotalus horridus metalloendopeptidase; hemorrhagic proteinase IV; Crotalus horridus horridus

venom hemorrhagic proteinase

**Comments:** A 56 kDa hemorrhagic endopeptidase from the venom of the timber rattlesnake (*Crotalus horridus* 

horridus) that cleaves basement membrane, hide powder and fibrinogen.

**References:** [455, 456]

[EC 3.4.24.47 created 1992]

EC 3.4.24.48

Accepted name: ruberlysin

**Reaction:** Cleavage of His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu, Tyr<sup>16</sup>—Leu and Gly<sup>23</sup>—Phe bonds in the B chain of insulin;

His—Pro, Pro—Phe, and Trp—Ser of angiotensin I; and Gly—Phe of Met enkephalin

**Other name(s):** Crotalus ruber metalloendopeptidase II; hemorrhagic toxin II

**Comments:** A 25 kDa hemorrhagic endopeptidase from the venom of the red rattlesnake (*Crotalus ruber ruber*)

that cleaves fibrinogen. In peptidase family M12 (astacin family)

**References:** [1966, 2864]

[EC 3.4.24.48 created 1992]

### EC 3.4.24.49

Accepted name: bothropasin

**Reaction:** Cleavage of His<sup>5</sup>—Leu, His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu, Tyr<sup>16</sup>—Leu and Phe<sup>24</sup>—Phe in insulin B chain

Other name(s): Bothrops jararaca venom metalloproteinase

**Comments:** Caseinolytic endopeptidase of jararaca snake (*Bothrops jararaca*) venom; 48 kDa. In peptidase family

M12

References: [1797]

[EC 3.4.24.49 created 1992]

#### EC 3.4.24.50

Accepted name: bothrolysin

**Reaction:** Cleavage of Gln<sup>4</sup>—His, Ser<sup>9</sup>—His and Ala<sup>14</sup>—Leu of insulin B chain and Pro—Phe of angiotensin

I

**Other name(s):** Bothrops metalloendopeptidase J; J protease

Comments: A 22.5 kDa endopeptidase from the venom of the jararaca snake (Bothrops jararaca), insensitive to

phosphoramidon at 0.5 mM. In peptidase family M12 (astacin family)

**References:** [2884]

[EC 3.4.24.50 created 1992]

### EC 3.4.24.51

Accepted name: ophiolysin

**Reaction:** Cleavage of Asn<sup>3</sup>—Gln, Gln<sup>4</sup>—His, His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu, and Tyr<sup>16</sup>—Leu in insulin B chain

**Other name(s):** Ophiophagus metalloendopeptidase

**Comments:** A 70 kDa endopeptidase from the venom of the king cobra (*Ophiophagus hannah*)

**References:** [3246]

[EC 3.4.24.51 created 1992]

### EC 3.4.24.52

Accepted name: trimerelysin I

**Reaction:** Cleavage of only two bonds His<sup>10</sup>—Leu and Ala<sup>14</sup>—Leu in the insulin B chain

Other name(s): Trimeresurus metalloendopeptidase I; hemorrhagic proteinase HR1A; hemorrhagic metalloproteinase

HR1A; metalloproteinase HR1A

Comments: A 60 kDa hemorrhagic endopeptidase of pI 4.4 from the venom of the habu snake (Trimeresurus

flavoviridis). In peptidase family M12 (astacin family)

**References:** [2197, 3247, 2863]

[EC 3.4.24.52 created 1992]

### EC 3.4.24.53

Accepted name: trimerelysin II

**Reaction:** Cleavage of Asn<sup>3</sup>—Gln, His<sup>10</sup>—Leu and Ala<sup>14</sup>—Leu in the insulin B chain, and the bond Z-Gly-

Pro—Leu-Gly-Pro in a small molecule substrate of microbial collagenase

Other name(s): Trimeresurus metalloendopeptidase II; proteinase H<sub>2</sub>; H<sub>2</sub>-proteinase

Comments: A 24 kDa nonhemorrhagic endopeptidase from the venom of the habu snake (Trimeresurus

flavoviridis). In peptidase family M12 (astacin family)

**References:** [2845, 2861]

[EC 3.4.24.53 created 1992]

### EC 3.4.24.54

**Accepted name:** mucrolysin

**Reaction:** Cleavage of Ser<sup>9</sup>—His, His<sup>10</sup>—Leu, Ala<sup>14</sup>—Leu, Leu<sup>15</sup>—Tyr and Tyr<sup>16</sup>—Leu bonds in insulin B

chain

Other name(s): Trimeresurus metalloendopeptidase A; mucrotoxin A

Comments: A 94 kDa hemorrhagic and fibrinogenolytic endopeptidase from the Chinese habu snake (Trimeresu-

rus mucrosquamatus) venom. In peptidase family M12 (astacin family)

**References:** [2793, 1467]

[EC 3.4.24.54 created 1992]

#### EC 3.4.24.55

Accepted name: pitrilysin

**Reaction:** Preferential cleavage of -Tyr<sup>16</sup>— Leu- and -Phe<sup>25</sup>— Tyr-bonds of oxidized insulin B chain. Also acts

on other substrates of less than 7 kDa such as insulin and glucagon

Other name(s): Escherichia coli protease III; protease Pi; proteinase Pi; PTR; Escherichia coli metalloproteinase Pi

**Comments:** From the periplasmic space of *Escherichia coli*. Inhibited by EDTA and 1,10-phenanthroline; not

thiol-dependent. Type example of peptidase family M16

**References:** [768, 20, 183, 618, 49]

[EC 3.4.24.55 created 1992 as EC 3.4.99.44, transferred 1993 to EC 3.4.24.55 (EC 3.4.99.45 created 1992, incorporated 1993)]

### EC 3.4.24.56

Accepted name: insulysin

**Reaction:** Degradation of insulin, glucagon and other polypeptides. No action on proteins

Other name(s): insulin-degrading enzyme; insulin protease; insulin proteinase; insulin-degrading neutral

proteinase; insulin-specific protease; insulin-glucagon protease; metalloinsulinase; IDE

Comments: A 110 kDa cytosolic enzyme, known from mammals and the fruit fly, Drosophila melanogaster. In-

hibited by bacitracin, chelating agents EDTA and 1,10-phenanthroline, and by thiol-blocking reagents such as *N*-ethylmaleimide, but not by phosphoramidon. In peptidase family M16 (pitrilysin family).

**References:** [660, 21, 661, 1562, 618]

[EC 3.4.24.56 created 1972 as EC 3.4.99.10, transferred 1976 EC 3.4.22.11, transferred 1978 to EC 3.4.99.45, transferred 1993 to to EC 3.4.24.56 (EC 3.4.99.46 created 1992, incorporated 2000)]

### EC 3.4.24.57

**Accepted name:** *O*-sialoglycoprotein endopeptidase

**Reaction:** Hydrolysis of *O*-sialoglycoproteins; cleaves -Arg<sup>31</sup> +Asp- bond in glycophorin A. Does not cleave

unglycosylated proteins, desialylated glycoproteins or glycoproteins that are only N-glycosylated

Other name(s): glycoprotease; glycophorin A proteinase; glycoproteinase; sialoglycoprotease; sialoglycoproteinase

Comments: An enzyme secreted by the bacterium *Pasteurella haemolytica*. Inhibited by EDTA (100 mM) and

1,10-phenanthroline. Type example of peptidase family M22

**References:** [3, 4, 2808]

[EC 3.4.24.57 created 1993]

### EC 3.4.24.58

Accepted name: russellysin

**Reaction:** Specifically activates several components of the blood clotting system, including coagulation factor X,

coagulation factor IX and protein C by cleavage of -Arg-bonds. Has no action on insulin B chain

Other name(s): Russell's viper venom factor X activator, RVV-X; blood-coagulation factor X activating enzyme; met-

alloproteinase RVV-x; Vipera russelli proteinase; Russell's viper blood coagulation factor X activator;

**RVV-V** 

Comments: This enzyme from the venom of Russell's viper (Vipera russelli) of 79 kDa comprises a heavy (59

kDa) and a heterogeneous light (18-21 kDa) chain. Contains Ca<sup>2+</sup> and Zn<sup>2+</sup>. The heavy chain contains the zinc-binding endopeptidase domain and a disintegrin. In peptidase family M12 (astacin fam-

ily)

**References:** [864, 1702, 2862]

[EC 3.4.24.58 created 1993]

EC 3.4.24.59

Accepted name: mitochondrial intermediate peptidase

Reaction: Release of an N-terminal octapeptide as second stage of processing of some proteins imported into the

mitochondrion

Other name(s): mitochondrial intermediate precursor-processing proteinase; MIP

**Comments:** A homologue of thimet oligopeptidase. Natural substrates are precursor proteins that have already

been processed by mitochondrial processing peptidase. In peptidase family M3 (thimet oligopeptidase

family)

**References:** [1256, 1257]

[EC 3.4.24.59 created 1993]

EC 3.4.24.60

Accepted name: dactylysin

**Reaction:** Hydrolysis of peptides of at least six residues, with bulky hydrophobic residues in the P1' position.

Shows a preference for hydrophobic doublets such as -Phe—Phe- and -Phe—Leu- in somatostatin-

(1-14)-peptide and dynorphin A-(1-6)-peptide, respectively

Other name(s): peptide hormone inactivating endopeptidase; PHIE

Comments: An endopeptidase of 100 kDa secreted from the skin of the amphibian, Xenopus laevis (Dactylêtre

du Cap). Resembles neprilysin in insensitivity to 1  $\mu$ M captopril, but differs from it in being insensitive to thiorphan (1  $\mu$ M) and unable to digest [Met<sup>5</sup>]enkephalin, [Leu<sup>5</sup>]enkephalin, oxytocin, and substance P-(7-11)-peptide. A similar endopeptidase is found in human neuroblastoma cells [577]

**References:** [378, 577, 1343]

[EC 3.4.24.60 created 1995]

EC 3.4.24.61

Accepted name: nardilysin

**Reaction:** Hydrolysis of polypeptides, preferably at -Xaa—Arg-Lys-, and less commonly at -Arg-Xaa-,

in which Xaa is not Arg or Lys

**Other name(s):** *N*-arginine dibasic convertase; NRD-convertase

Comments: Enzyme of 133 kDa from rat brain and testis. A homologue of pitrilysin containing the His-Phe-Leu-

Glu-His zinc-binding sequence, and a highly acidic stretch of 71 residues. Unusually for a metal-loendopeptidase, inhibited by bestatin, amastatin and *N*-ethylmaleimide. In peptidase family M16

(pitrilysin family)

**References:** [953, 932, 425, 2280]

[EC 3.4.24.61 created 1995]

EC 3.4.24.62

Accepted name: magnolysin

**Reaction:** Hydrolysis of polypeptides with Arg or Lys in P1 and P2, e.g. to hydrolyse *pro*-oxytocin at -Lys-

Arg—Ala-Val-. The specificity further depends on the organization of a β-turn-α-helix of nine or

more residues containing the paired basic amino acids near the centre [3]

Other name(s): bovine neurosecretory granule protease cleaving pro-oxytocin/neurophysin; pro-oxytocin/neurophysin

convertase; prooxyphysin proteinase; pro-oxytocin convertase

Comments: An endopeptidase of 58 kDa known from bovine pituitary neurosecretory granules and bovine and

human corpus luteum [2293, 998]. Inhibited by EDTA [457]

**References:** [457, 503, 276, 2293, 998]

[EC 3.4.24.62 created 1995]

EC 3.4.24.63

Accepted name: meprin B

**Reaction:** Hydrolysis of proteins, including azocasein, and peptides. Hydrolysis of -His<sup>5</sup>—Leu-, -Leu<sup>6</sup>—Cys-,

-Ala<sup>14</sup>—Leu- and -Cys<sup>19</sup>—Gly- bonds in insulin B chain

Other name(s): meprin-b

**Comments:** A brush border membrane-bound metalloendopeptidase known from the intestine of all mouse strains

that have been tested, and the kidney of certain inbred strains. A tetramer of meprin  $\beta$  subunits (in contrast to meprin A, which contains both  $\alpha$  and  $\beta$  subunits). Occurs in the kidney as a proenzyme that can be activated by trypsin. Meprin B is inhibited by both EDTA and 1,10-phenanthroline, but

not by phosphoramidon, captopril or thiorphan. In peptidase family M12 (astacin family)

**References:** [1522, 957, 1325, 3210]

[EC 3.4.24.63 created 1995]

EC 3.4.24.64

**Accepted name:** mitochondrial processing peptidase

**Reaction:** Release of N-terminal targetting peptides from precursor proteins imported into the mitochondrion,

typically with Arg in position P2

Other name(s): processing enhancing peptidase (for one of two subunits); mitochondrial protein precursor-processing

proteinase; matrix peptidase; matrix processing peptidase; matrix processing proteinase; mitochon-

drial protein precursor-processing proteinase; MPP

Comments: Known from the mitochondrial matrix of fungi and mammals. Formed from two subunits, both ho-

mologous with pitrilysin [2382], and the products of the MAS1 and MAS2 genes in yeast. In peptidase

family M16 (pitrilysin family).

**References:** [1319, 3202, 2382, 1365, 320]

[EC 3.4.24.64 created 1989/90 as EC 3.4.99.41, transferred 1995 to EC 3.4.24.64]

EC 3.4.24.65

Accepted name: macrophage elastase

**Reaction:** Hydrolysis of soluble and insoluble elastin [1]. Specific cleavages are also produced at -Ala<sup>14</sup> Leu-

and -Tyr<sup>16</sup>—Leu- in the B chain of insulin [2]

Other name(s): metalloelastase; human macrophage metalloelastase (HME)

**Comments:** This enzyme is synthesized as a proenzyme of 53 kDa that is converted to an active form of 22 kDa.

cDNA sequences have been obtained for the mouse [2623] and human [2624] enzymes. In peptidase

family M10 (interstitial collagenase family)

**References:** [135, 1430, 2623, 2624]

[EC 3.4.24.65 created 1995]

EC 3.4.24.66

Accepted name: choriolysin L

Reaction: Hydrolysis of the inner layer of fish egg envelope. Also hydrolysis of casein and small molecule sub-

strates such as succinyl-Leu-Leu-Val-Tyr—7-(4-methyl)coumarylamide

**Other name(s):** teleost hatching enzyme (component); low choriolytic enzyme (LCE)

**Comments:** Known from the teleost fish *Oryzias latipes* (medaka). Efficient dissolution of the egg membrane re-

quires concerted action with choriolysin H. A 24 kDa peptidase family M12 (astacin family)

**References:** [3272, 3273, 3275, 3277]

[EC 3.4.24.66 created 1995]

EC 3.4.24.67

Accepted name: choriolysin H

Reaction: Hydrolysis of the inner layer of fish egg envelope. Also hydrolysis of casein and small molecule sub-

strates such as succinyl-Leu-Leu-Val-Tyr—7-(4-methyl)coumarylamide

**Other name(s):** teleost hatching enzyme (component); high choriolytic enzyme (HCE)

Comments: Known from the teleost fish Oryzias latipes (medaka). Efficient dissolution of the egg membrane re-

quires concerted action with choriolysin L. A 25.5 kDa peptidase in family M12 (astacin family)

**References:** [3243, 3274, 3276, 3277, 1625]

[EC 3.4.24.67 created 1995]

EC 3.4.24.68

Accepted name: tentoxilysin

**Reaction:** Hydrolysis of -Gln<sup>76</sup>—Phe- bond in synaptobrevin (also known as neuronal vesicle-associated mem-

brane protein, VAMP)

Other name(s): tetanus neurotoxin

Comments: Zinc enzyme produced by Clostridium tetani. Proenzyme of 150 kDa is processed to disulfide-linked

subunits of 100 and 50 kDa, the latter being responsible for the endopeptidase activity. Weakly inhibited by captopril, and phosphoramidon. The clostridial neurotoxins disable the neuroexocytosis apparatus, and have been described as the most toxic substances known. Tentoxilysin acts at the spinal inhibitory interneurons, blocking the release of various neurotransmitters to produce spastic paralysis.

Type example of peptidase family M27 (tentoxilysin family)

**References:** [832, 2557, 2561, 1954, 2559]

[EC 3.4.24.68 created 1995]

EC 3.4.24.69

Accepted name: bontoxilysin

Reaction: Limited hydrolysis of proteins of the neuroexocytosis apparatus, synaptobrevin (also known as neu-

ronal vesicle-associated membrane protein, VAMP), synaptosome-associated protein of 25 kDa

(SNAP25) or syntaxin. No detected action on small molecule substrates

Other name(s): botulinum neurotoxin; BoNT

**Comments:** This zinc enzyme, produced by *Clostridium botulinum*, occurs as forms A-G that differ in specificity

of action on the proteins of the neuroexocytosis apparatus [2560, 2562, 2563, 2558, 1954]. The 150-kDa proenzymes of bontoxilysin are processed to disulfide-linked subunits of 100 and 50 kDa, the latter being responsible for the endopeptidase activities. Weakly inhibited by captopril, and phosphoramidon. Toxicity is due to action at the neuromuscular junctions that blocks release of acetylcholine, causing flaccid paralysis, in contrast to the spastic paralysis caused by tentoxilysin. In peptidase fam-

ily M27 (tentoxilysin family)

**References:** [2560, 2562, 2563, 2558, 1954, 2559]

[EC 3.4.24.69 created 1995]

EC 3.4.24.70

Accepted name: oligopeptidase A

**Reaction:** Hydrolysis of oligopeptides, with broad specificity. Gly or Ala commonly occur as P1 or P1' residues,

but more distant residues are also important, as is shown by the fact that Z-Gly-Pro-Gly-Gly-Pro-

Ala is cleaved, but not Z-(Gly)<sub>5</sub> [4]

Other name(s): 68000-M signal peptide hydrolase

Comments: Known from Escherichia coli and Salmonella typhimurium. A zinc metallopeptidase, in peptidase

family M3 (thimet oligopeptidase family), but differs from thimet oligopeptidase in lack of thiol-

activation

**References:** [2118, 476, 475, 474]

[EC 3.4.24.70 created 1996]

EC 3.4.24.71

Accepted name: endothelin-converting enzyme 1

**Reaction:** Hydrolysis of the -Trp<sup>21</sup>—Val- bond in big endothelin to form endothelin 1

**Other name(s):** endothelin-converting enzyme; ECE-1

**Comments:** A phosphoramidon-sensitive metalloendopeptidase in peptidase family M13 (neprilysin family). An

integral membrane protein predominantly of endothelial cells, which generates the potent vasocon-

strictor endothelin 1 from its inactive precursor

**References:** [2840, 2632, 3231]

[EC 3.4.24.71 created 1996]

EC 3.4.24.72

Accepted name: fibrolase

**Reaction:** Hydrolysis of -Ala<sup>14</sup>—Leu- in insulin B chain and -Lys<sup>413</sup>—Leu- in A $\alpha$ -chain of fibrinogen **Other name(s):** fibrinolytic proteinase; *Agkistrodon contortrix contortrix* metalloproteinase; *Agkistrodon contortrix* 

contortrix venom metalloproteinase

**Comments:** A 23-kDa, non-hemorrhagic enzyme from the venom of the southern copperhead snake (Agkistrodon

contortix contortix). In peptidase family M12 (astacin family)

**References:** [12, 996, 2372, 1731, 2411]

[EC 3.4.24.72 created 1996]

EC 3.4.24.73

Accepted name: jararhagin

**Reaction:** Hydrolysis of -His<sup>10</sup>—Leu-, -Ala<sup>14</sup>—Leu-, -Tyr<sup>16</sup>—Leu-and -Phe<sup>24</sup>—Phe- bonds in insulin B chain

**Other name(s):** HF2-proteinase; JF1

**Comments:** Hemorrhagic endopeptidase from the venom of the jararaca snake (*Bothrops jararaca*). The 52-kDa

enzyme contains a disintegrin domain [2224]. In peptidase family M12 (astacin family)

**References:** [1798, 90, 2224]

[EC 3.4.24.73 created 1996]

EC 3.4.24.74

**Accepted name:** fragilysin

**Reaction:** Broad proteolytic specificity, bonds hydrolysed including -Gly—Leu-, -Met—Leu-, -Phe—Leu-,

-Cys—Leu-, Leu—Gly

Other name(s): Bacteroides fragilis (entero)toxin

Comments: Thought to be a cause of diarrhoea in animals and humans. Hydrolyses extracellular matrix proteins,

and disrupts tight junctions of intestinal epithelial cells. Also degrades intracellular, cytoskeletal pro-

teins actin, myosin and others. In peptidase family M10 (interstitial collagenase family)

**References:** [1951, 2124, 632, 1516, 1476]

[EC 3.4.24.74 created 1997]

EC 3.4.24.75

Accepted name: lysostaphin

**Reaction:** Hydrolysis of the -Gly—Gly- bond in the pentaglycine inter-peptide link joining staphylococcal cell

wall peptidoglycans

Other name(s): glycyl-glycine endopeptidase

Comments: A zinc-dependent, 25-kDa endopeptidase from Staphylococcus simulans. Lyses cells of S. aureus, in

particular, by its action on the cross-bridges of the cell wall. Type example of peptidase family M23.

**References:** [2392, 109, 2920]

[EC 3.4.24.75 created 1997]

EC 3.4.24.76

Accepted name: flavastacin

**Reaction:** Hydrolyses polypeptides on the amino-side of Asp in -Xaa—Asp-. Acts very slowly on -Xaa—Glu **Comments:** A zinc metalloendopeptidase in peptidase family M12 (astacin family), secreted by the bacterium

Flavobacterium meningosepticum. The specificity is similar to that of EC 3.4.24.33, peptidyl-Asp metalloendopeptidase from Pseudomonas fragi but the two are reported to be structurally dissimilar

**References:** [2889]

[EC 3.4.24.76 created 2000]

EC 3.4.24.77

Accepted name: snapalysin

**Reaction:** Hydrolyses proteins with a preference for Tyr or Phe in the P1' position. Has no action on amino-acid

*p*-nitroanilides

Other name(s): small neutral protease; SnpA gene product (Streptomyces lividans)

**Comments:** Type example of peptidase family M7.

**References:** [1570, 336, 1569]

[EC 3.4.24.77 created 2001]

EC 3.4.24.78

Accepted name: gpr endopeptidase

**Reaction:** Endopeptidase action with P4 Glu or Asp, P1 preferably Glu > Asp, P1' hydrophobic and P2' Ala

**Other name(s):** germination proteinase

Comments: Initiates the degradation of small, acid-soluble proteins during spore germination in Bacillus mega-

terium. Type example of peptidase family M63.

**References:** [2309]

[EC 3.4.24.78 created 2003]

EC 3.4.24.79

Accepted name: pappalysin-1

**Reaction:** Cleavage of the Met<sup>135</sup>—Lys bond in insulin-like growth factor binding protein (IGFBP)-4, and the

Ser<sup>143</sup>—Lys bond in IGFBP-5

Other name(s): insulin-like growth factor binding protein-4 protease; pregnancy-associated plasma protein-A

**Comments:** A 400-kDa disulfide-linked dimer. Circulates in human pregnancy mainly as a complex with the pro-

form of eosinophil major basic protein, which acts as an inhibitor of the peptidase. The rate of hydrolysis of IGFBP-4 is increased about 20-fold by the presence of insulin-like growth factor (IGF),

whereas that of IGFBP-5 is decreased about two-fold. In peptidase family M43.

**References:** [1611, 409]

[EC 3.4.24.79 created 2003]

EC 3.4.24.80

**Accepted name:** membrane-type matrix metalloproteinase-1

**Reaction:** Endopeptidase activity. Activates progelatinase A by cleavage of the propeptide at Asn<sup>37</sup>—Leu.

Other bonds hydrolysed include Gly<sup>35</sup>—Ile in the propeptide of collagenase 3, and Asn<sup>341</sup>—Phe,

Asp<sup>441</sup>—Leu and Gln<sup>354</sup>—Thr in the aggrecan interglobular domain

Other name(s): matrix metalloproteinase 14

Comments: In peptidase family M10, but, unlike most members of the family, is membrane-anchored. Believed to

play an important role in the activation of progelatinase A at cell surfaces.

**References:** [1277]

[EC 3.4.24.80 created 2003]

EC 3.4.24.81

**Accepted name:** ADAM10 endopeptidase

**Reaction:** Endopeptidase of broad specificity

**Other name(s):** Kuzbanian protein; myelin-associated disintegrin metalloproteinase

**Comments:** In peptidase family M12. Partially responsible for the "α-secretase" activity in brain that degrades the

potentially harmful β-amyloid peptide. Work with ADAM10-deficient mice supports a role in Notch

signalling.

**References:** [1010]

[EC 3.4.24.81 created 2003]

EC 3.4.24.82

**Accepted name:** ADAMTS-4 endopeptidase

**Reaction:** Glutamyl endopeptidase; bonds cleaved include -Thr-Glu-Gly-Glu<sup>373</sup>—Ala-Arg-Gly-Ser- in the in-

terglobular domain of mammalian aggrecan

Other name(s): aggrecanase-1

Comments: In peptidase family M12. Thought to be biologically significant for the degradation of the aggrecan

component of cartilage matrix.

**References:** [3165]

[EC 3.4.24.82 created 2003]

EC 3.4.24.83

Accepted name: anthrax lethal factor endopeptidase

**Reaction:** Preferred amino acids around the cleavage site can be denoted BBBBxHx—H, in which B denotes

Arg or Lys, H denotes a hydrophobic amino acid, and x is any amino acid. The only known protein

substrates are mitogen-activated protein (MAP) kinase kinases

Other name(s): lethal toxin

**Comments:** From the bacterium *Bacilus anthracis* that causes anthrax. One of three proteins that are collectively

termed anthrax toxin. Cleaves several MAP kinase kinases near their N-termini, preventing them from

phosphorylating the downstream mitogen-activated protein kinases. In peptidase family M34.

**References:** [2232]

[EC 3.4.24.83 created 2003]

EC 3.4.24.84

**Accepted name:** Ste24 endopeptidase

**Reaction:** The peptide bond hydrolysed can be designated -C—aaX in which C is an S-isoprenylated cysteine

residue, a is usually aliphatic and X is the C-terminal residue of the substrate protein, and may be any

of several amino acids

**Comments:** Type example of peptidase family M48. One of two enzymes that can catalyse this processing step

for mating a-factor in yeast. Subsequently, the S-isoprenylated cysteine residue that forms the new

C-terminus is methyl-esterified and forms a hydrophobic membrane-anchor.

**References:** [2865]

[EC 3.4.24.84 created 2003]

EC 3.4.24.85

**Accepted name:** S2P endopeptidase

**Reaction:** Cleaves several transcription factors that are type-2 transmembrane proteins within membrane-

spanning domains. Known substrates include sterol regulatory element-binding protein (SREBP) -1, SREBP-2 and forms of the transcriptional activator ATF6. SREBP-2 is cleaved at the site

DRSR<u>ILL</u><sup>483</sup>—CVLTFLCLSFNPLTSLLQWGGA, in which the membrane-spanning segment is underlined. The residues NP (bold), 11 residues distal to the site of cleavage in the membrane-spanning domain, are important for cleavage by S2P endopeptidase. Replacement of either of these residues

does not prevent cleavage, but there is no cleavage if both of these residues are replaced.

**Comments:** Type example of peptidase family M50. The transcription factors SREBP-1 and -2 are synthesized as

precursor proteins that are attached to the membranes of the endoplasmic reticulum and two cleavages are needed to release the active factor so that it can move to the nucleus. This enzyme cleaves the

second of these, and is thus the "site 2 protease", S2P.

**References:** [310]

[EC 3.4.24.85 created 2003]

EC 3.4.24.86

**Accepted name:** ADAM 17 endopeptidase

Reaction: Narrow endopeptidase specificity. Cleaves Pro-Leu-Ala-Gln-Ala-Val-Arg-Ser-Ser in the

membrane-bound, 26-kDa form of tumour necrosis factor α (TNFα). Similarly cleaves other

membrane-anchored, cell-surface proteins to "shed" the extracellular domains

Other name(s): tumor necrosis factor  $\alpha$ -converting enzyme; TACE

**Comments:** In peptidase family M12. *In vivo*, the cleavage of tumour necrosis factor  $\alpha$  precursor releases the solu-

ble, 17-kDa TNF $\alpha$ , which induces inflammation.

**References:** [228]

[EC 3.4.24.86 created 2003]

EC 3.4.24.87

Accepted name: ADAMTS13 endopeptidase

**Reaction:** The enzyme cleaves the von Willebrand factor at bond Tyr<sup>842</sup>—Met<sup>843</sup> within the A2 domain **Other name(s):** ADAMTS VWF cleaving metalloprotease; ADAMTS-13; ADAMTS13; vWF-cleaving protease;

VWF-CP; vWF-degrading protease; Upshaw factor; von Willebrand factor cleaving protease;

ADAMTS13 peptidase

**Comments:** In peptidase family M12.

**References:** [841, 634]

[EC 3.4.24.87 created 2009]

[3.4.24.88 Transferred entry. desampylase. Transferred to EC 3.4.19.15 desampylase]

[EC 3.4.24.88 created 2015, deleted 2016]

EC 3.4.24.89

**Accepted name:** Pro-Pro endopeptidase

**Reaction:** The enzyme catalyses the hydrolytic cleavage of peptide bonds between two proline residues

Other name(s): metalloprotease CD2830

**Comments:** This metalloprotease, which is secreted by the bacterium *Peptoclostridium difficile*, contains zinc.

**References:** [351, 1116, 1115]

[EC 3.4.24.89 created 2015]

# EC 3.4.25 Threonine endopeptidases

EC 3.4.25.1

**Accepted name:** proteasome endopeptidase complex

**Reaction:** Cleavage of peptide bonds with very broad specificity

Other name(s): ingensin; macropain; multicatalytic endopeptidase complex; prosome; multicatalytic proteinase (com-

plex); MCP; proteasome; large multicatalytic protease; multicatalytic proteinase; proteasome or-

ganelle; alkaline protease; 26S protease; tricorn proteinase; tricorn protease

Comments: A 20-S protein composed of 28 subunits arranged in four rings of seven. The outer rings are com-

posed of  $\alpha$  subunits, but the  $\beta$  subunits forming the inner rings are responsible for peptidase activity. In eukaryotic organisms there are up to seven different types of  $\beta$  subunits, three of which may carry the N-terminal threonine residues that are the nucleophiles in catalysis, and show different specificities. The molecule is barrel-shaped, and the active sites are on the inner surfaces. Terminal apertures restrict access of substrates to the active sites. There is evidence that catalytic subunits are replaced by others under some conditions so as to alter the specificity of proteolysis, perhaps optimizing it for the formation of antigenic peptides. A complex of the 20-S proteasome endopeptidase complex with a 19-S regulatory unit is the 26-S proteasome that degrades ubiquitin-protein conjugates. Type example of peptidase family T1.

**References:** [2600, 499, 991, 605]

[EC 3.4.25.1 created 1978 as EC 3.4.24.5, part transferred 1989 to EC 3.4.22.21, transferred 1992 to EC 3.4.99.46, transferred 2000 to EC 3.4.25.1]

EC 3.4.25.2

**Accepted name:** HslU—HslV peptidase

**Reaction:** ATP-dependent cleavage of peptide bonds with broad specificity.

Other name(s): HslUV; HslV-HslU; HslV peptidase; ATP-dependent HslV-HslU proteinase; caseinolytic protease

X; caseinolytic proteinase X; ClpXP ATP-dependent protease; ClpXP protease; ClpXP serine proteinase; *Escherichia coli* ClpXP serine proteinase; HslUV proteinase; HslUV proteinase; HslUV proteinase; HslVU pro-

tease; HslVU proteinase; protease HslVU; proteinase HslUV

Comments: The HslU subunit of the HslU—HslV complex functions as an ATP dependent 'unfoldase'. The bind-

ing of ATP and its subsequent hydrolysis by HslU are essential for unfolding of protein substrates subsequently hydrolysed by HslV [3291]. HslU recognizes the N-terminal part of its protein substrates and unfolds these before they are guided to HslV for hydrolysis [333]. In peptidase family T1.

**References:** [3112, 2092, 2364, 3292, 3291, 1382, 333]

[EC 3.4.25.2 created 2009, modified 2010]

### EC 3.4.99 Endopeptidases of unknown catalytic mechanism (sub-subclass is currently empty)

[3.4.99.1] Transferred entry, acrocylindricum proteinase, Now EC 3.4.23.28, acrocylindropepsin]

[EC 3.4.99.1 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]

[3.4.99.2 Deleted entry. agavain]

[EC 3.4.99.2 created 1972, deleted 1992]

[3.4.99.3	Deleted entry. angiotensinase]
	[EC 3.4.99.3 created 1972, deleted 1992]
[3.4.99.4	Transferred entry. aspartylendopeptidase. Now EC 3.4.23.12, nepenthesin]
	[EC 3.4.99.4 created 1972, deleted 1978]
[3.4.99.5	Transferred entry. Clostridium histolyticum collagenase 2. Now EC 3.4.24.3, microbial collagenase]
	[EC 3.4.99.5 created 1972, deleted 1978]
[3.4.99.6	Transferred entry. crayfish low-molecular-weight proteinase. Now EC 3.4.24.21, astacin]
	[EC 3.4.99.6 created 1972, deleted 1992]
[3.4.99.7	Deleted entry. euphorbain]
	[EC 3.4.99.7 created 1972, deleted 1989]
[3.4.99.8	Deleted entry. Gliocladium proteinase]
	[EC 3.4.99.8 created 1972, deleted 1992]
[3.4.99.9	Deleted entry. hurain. Now considered EC 3.4.21.25, cucumisin]
	[EC 3.4.99.9 created 1972, deleted 1992]
[3.4.99.10	Transferred entry. insulinase. Now EC 3.4.24.56, insulysin]
[EC 3.4	.99.10 created 1972, transferred 1976 to EC 3.4.22.11, transferred 1978 to EC 3.4.99.45, transferred 1993 to EC 3.4.24.56]
[3.4.99.11	Deleted entry. Streptomyces alkalophilic keratinase]
	[EC 3.4.99.11 created 1965 as EC 3.4.4.25, transferred 1972 to EC 3.4.99.11, deleted 1992]
[3.4.99.12	Deleted entry. Trichophyton mentagrophytes keratinase]
	[EC 3.4.99.12 created 1972, deleted 1978 [transferred to EC 3.4.24.10, deleted 1992]]
[3.4.99.13 dase]	Transferred entry. β-lytic proteinase (Mycobacterium sorangium). Now EC 3.4.24.32, β-lytic metalloendopepti-
	[EC 3.4.99.13 created 1972, deleted 1978 [transferred to EC 3.4.24.4, deleted 1992]]
[3.4.99.14	Deleted entry. mexicanain]
	[EC 3.4.99.14 created 1972, deleted 1992]
[3.4.99.15	Deleted entry. Paecilomyces proteinase]
	[EC 3.4.99.15 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]
[3.4.99.16	Deleted entry. Penicillium notatum extracellular proteinase]
	[EC 3.4.99.16 created 1972, deleted 1992]
[3.4.99.17	Deleted entry. peptidoglycan endopeptidase]
	[EC 3.4.99.17 created 1972, deleted 1992]
[3.4.99.18	Deleted entry. pinguinain]
	[EC 3.4.99.18 created 1972, deleted 1992]
[3.4.99.19	Transferred entry. renin. Now EC 3.4.23.15, renin]
	[EC 3.4.99.19 created 1972, deleted 1981]

[EC 3.4.99.20 created 1972, deleted 1992]   [3.4.99.21   Deleted entry: solamain. Now considered EC 3.4.21.25, cucumisin]	[3.4.99.20	Deleted entry. Scopulariopsis proteinase]
[EC 3.4.99.21 created 1972, deleted 1992] [3.4.99.22 Transferred entry: staphylokinase. Now EC 3.4.24.29, aureolysin] [EC 3.4.99.23 Deleted entry: tabernamontanain. Now considered EC 3.4.21.25, cucumisin] [EC 3.4.99.23 Created 1972, modified 1976, deleted 1978 [transferred to EC 3.4.24.4, deleted 1992]] [3.4.99.24 Deleted entry: tabernamontanain. Now considered EC 3.4.21.25, cucumisin] [EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]] [3.4.99.26 Transferred entry: transfers acid proteinase. Now EC 3.4.2.3.21, rhizopusepsin] [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]] [3.4.99.27 Deleted entry: urokinase. Now EC 3.4.21.68, ε-plasminogen activator] [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]] [3.4.99.27 Deleted entry: Echis carinatus prothrombin-activating proteinase. [EC 3.4.99.27 created 1978, deleted 1992] [3.4.99.28 Transferred entry: Oxyurarus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin] [EC 3.4.99.28 created 1978, deleted 1992] [3.4.99.29 Deleted entry: Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase] [EC 3.4.99.20 created 1978, deleted 1992] [3.4.99.30 Transferred entry: sissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992] [3.4.99.32 Transferred entry: Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase] [EC 3.4.99.32 created 1978, deleted 1992] [3.4.99.33 Deleted entry: cathepsin R] [EC 3.4.99.33 created 1978, deleted 1992] [3.4.99.34 Deleted entry: promurein-leeder peptidase. Now EC 3.4.23.36, signal peptidase II] [EC 3.4.99.35 created 1981, deleted 1994] [EC 3.4.99.35 created 1984, deleted 1994] [EC 3.4.99.35 created 1984, deleted 1995] [EC 3.4.99.35 created 1984, deleted 1995]		[EC 3.4.99.20 created 1972, deleted 1992]
	[3.4.99.21	Deleted entry. solanain. Now considered EC 3.4.21.25, cucumisin]
[EC 3.4.99.22 created 1972, modified 1976, deleted 1978 [transferred to EC 3.4.24.4, deleted 1992]     3.4.99.23   Deleted entry. tabernamontanain. Now considered EC 3.4.21.25, cucumisin]     (EC 3.4.99.23 created 1972, deleted 1992)     3.4.99.24   Deleted entry. Tenebrio G-proteinase]     (EC 3.4.99.24 created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]]     3.4.99.25   Transferred entry. trametes acid proteinase. Now EC 3.4.23.21, rhizopuspepsin]     (EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]     3.4.99.26   Transferred entry. urokinase. Now EC 3.4.21.68, t-plasminogen activator]     (EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]     3.4.99.27   Deleted entry. Echis carinains prothrombin-activating proteinase]     (EC 3.4.99.28 created 1978, deleted 1992)     3.4.99.28   Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]     (EC 3.4.99.28 created 1978, deleted 1992)     3.4.99.30   Deleted entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]     (EC 3.4.99.30 created 1978, deleted 1992)     3.4.99.31   Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase     (EC 3.4.99.31 created 1978, deleted 1992)     3.4.99.32   Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase     (EC 3.4.99.32 created 1978, deleted 1992)     3.4.99.33   Deleted entry: cathepsin R     (EC 3.4.99.33 created 1981, deleted 1992)     3.4.99.34   Deleted entry: mytilidase     (EC 3.4.99.34 created 1981, deleted 1992)     3.4.99.35   Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II     (EC 3.4.99.35 created 1984, deleted 1995)     3.4.99.36   Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase II		[EC 3.4.99.21 created 1972, deleted 1992]
[EC 3.4.99.23   created 1972, deleted 1992    [EC 3.4.99.24   created 1972, deleted 1992    [EC 3.4.99.24   created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]   [EC 3.4.99.24   created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]   [EC 3.4.99.25   created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]   [EC 3.4.99.25   created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]   [EC 3.4.99.26   created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]   [EC 3.4.99.26   created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]   [EC 3.4.99.26   created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]   [EC 3.4.99.27   created 1978, deleted 1992]   [EC 3.4.99.27   created 1978, deleted 1992]   [EC 3.4.99.28   created 1978, deleted 1992]   [EC 3.4.99.28   created 1978, deleted 1992]   [EC 3.4.99.29   created 1978, deleted 1992]   [EC 3.4.99.29   created 1978, deleted 1992]   [EC 3.4.99.30   created 1984, deleted	[3.4.99.22	Transferred entry. staphylokinase. Now EC 3.4.24.29, aureolysin]
[EC 3.4.99.23 created 1972, deleted 1992] [3.4.99.24 Deleted entry. Temebrio \(\alpha\)-proteinase] [EC 3.4.99.24 created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]] [3.4.99.25 Transferred entry. trametes acid proteinase. Now EC 3.4.23.21, rhizopuspepsin] [EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]] [3.4.99.26 Transferred entry. urokinase. Now EC 3.4.21.68, t-plasminogen activator] [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]] [3.4.99.27 Deleted entry. Echis carinatus prothrombin-activating proteinase] [EC 3.4.99.27 created 1978, deleted 1992] [3.4.99.28 Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin] [EC 3.4.99.28 created 1978, deleted 1992] [3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I] [EC 3.4.99.29 created 1978, deleted 1992] [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase] [EC 3.4.99.30 created 1978, deleted 1992] [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetalligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992] [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992] [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.34 created 1984 [transferred to EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.34 created 1984, deleted 1992] [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [EC 3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]  [EC 3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase II]		[EC 3.4.99.22 created 1972, modified 1976, deleted 1978 [transferred to EC 3.4.24.4, deleted 1992]]
13.4.99.24   Deleted entry. Tenebrio α-proteinase   [EC 3.4.99.24 created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]]   13.4.99.25   Transferred entry. trametes acid proteinase. Now EC 3.4.23.21, rhizopuspepsin]   [EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]   13.4.99.26   Transferred entry. urokinase. Now EC 3.4.21.68, t-plasminogen activator]   [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]   13.4.99.27   Deleted entry. Echis carinatus prothrombin-activating proteinase   [EC 3.4.99.27 created 1978, deleted 1992]   13.4.99.28   Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]   [EC 3.4.99.28 created 1978, deleted 1992]   13.4.99.30   Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]   [EC 3.4.99.30 created 1978, deleted 1992]   13.4.99.31   Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase   [EC 3.4.99.31 created 1978, deleted 1992]   13.4.99.32   Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]   [EC 3.4.99.32 created 1978, deleted 1992]   13.4.99.33   Deleted entry. cathepsin R]   [EC 3.4.99.34 created 1978, deleted 1992]   13.4.99.34   Deleted entry. mytilidase]   [EC 3.4.99.34 created 1984 [transferred to EC 3.4.21.52, deleted 1992]   13.4.99.35   Transferred entry. premurein-leader peptidase. Now EC 3.4.2.3.36, signal peptidase II]   [EC 3.4.99.35 created 1984, deleted 1995]   13.4.99.36   Transferred entry. teader peptidase. Now EC 3.4.21.89, signal peptidase II]   [EC 3.4.99.35 created 1984, deleted 1995]   13.4.99.36   13.4.99.36   13.4.99.36   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13.4.99.38   13	[3.4.99.23	Deleted entry. tabernamontanain. Now considered EC 3.4.21.25, cucumisin]
[EC 3.4.99.24 created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]]  [3.4.99.25 Transferred entry. trametes acid proteinase. Now EC 3.4.23.21, rhizopuspepsin]  [EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]  [3.4.99.26 Transferred entry. urokinase. Now EC 3.4.21.68, t-plasminogen activator]  [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]  [3.4.99.27 Deleted entry. Echis carinatus prothrombin-activating proteinase]  [EC 3.4.99.28 treated 1978, deleted 1992]  [3.4.99.28 Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I]  [EC 3.4.99.29 created 1978, deleted 1992]  [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry: tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.34 created 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.35 Transferred entry. mytilidase]  [EC 3.4.99.34 created 1984, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [EC 3.4.99.36 created 1984, deleted 1995]		[EC 3.4.99.23 created 1972, deleted 1992]
[EC 3.4.99.25 Transferred entry. trametes acid proteinase. Now EC 3.4.23.21, rhizopuspepsin]  [EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]  [3.4.99.26 Transferred entry. urokinase. Now EC 3.4.21.68, t-plasminogen activator]  [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]  [3.4.99.27 Deleted entry: Echis carinatus prothrombin-activating proteinase]  [EC 3.4.99.27 created 1978, deleted 1992]  [3.4.99.28 Transferred entry: Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.28 created 1978, deleted 1992]  [3.4.99.29 Deleted entry: Myxobacter AL-1 proteinase I]  [EC 3.4.99.29 created 1978, deleted 1992]  [3.4.99.30 Transferred entry: Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry: tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetolligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry: Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry: cathepsin R]  [EC 3.4.99.34 created 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.35 Transferred entry: mytilidase]  [EC 3.4.99.34 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.35 Transferred entry: premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]	[3.4.99.24	Deleted entry. Tenebrio α-proteinase]
[EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]  [3.4.99.26 Transferred entry: unokinase. Now EC 3.4.21.68, t-plasminogen activator]  [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]  [3.4.99.27 Deleted entry: Echis carinatus prothrombin-activating proteinase]  [EC 3.4.99.27 created 1978, deleted 1992]  [3.4.99.28 Transferred entry: Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.28 created 1978, deleted 1992]  [3.4.99.29 Deleted entry: Myxobacter AL-1 proteinase I]  [EC 3.4.99.29 created 1978, deleted 1992]  [3.4.99.30 Transferred entry: Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry: tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry: Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry: cathepsin R]  [EC 3.4.99.34 created 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.35 Transferred entry: mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry: premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]		[EC 3.4.99.24 created 1972, deleted 1978 [transferred to EC 3.4.21.18, deleted 1992]]
[3.4.99.26 Transferred entry. urokinase. Now EC 3.4.21.68, t-plasminogen activator]  [EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]  [3.4.99.27 Deleted entry. Echis carinatus prothrombin-activating proteinase]  [EC 3.4.99.27 created 1978, deleted 1992]  [3.4.99.28 Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.28 created 1978, deleted 1992]  [3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I]  [EC 3.4.99.29 created 1978, deleted 1992]  [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.34 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.35 Transferred entry. mytilidase]  [EC 3.4.99.35 created 1981, deleted 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]	[3.4.99.25	Transferred entry. trametes acid proteinase. Now EC 3.4.23.21, rhizopuspepsin]
[EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]  [3.4.99.27 Deleted entry. Echis carinatus prothrombin-activating proteinase]  [EC 3.4.99.27 created 1978, deleted 1992]  [3.4.99.28 Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.28 created 1978, deleted 1992]  [3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I]  [EC 3.4.99.29 created 1978, deleted 1992]  [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.34 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.35 Transferred entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.25 created 1972, deleted 1978 [transferred to EC 3.4.23.6, deleted 1992]]
[3.4.99.27 Deleted entry. Echis carinatus prothrombin-activating proteinase]  [EC 3.4.99.27 created 1978, deleted 1992] [3.4.99.28 Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.28 created 1978, deleted 1992] [3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I]  [EC 3.4.99.29 created 1978, deleted 1992] [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992] [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992] [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992] [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]] [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.35 created 1981, deleted 1991] [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.26	Transferred entry. urokinase. Now EC 3.4.21.68, t-plasminogen activator]
[EC 3.4.99.27 created 1978, deleted 1992]  [3.4.99.28 Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.28 created 1978, deleted 1992]  [3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I]  [EC 3.4.99.29 created 1978, deleted 1992]  [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry: tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry: Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry: cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry: mytilidase]  [EC 3.4.99.35 Transferred entry: premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 Created 1984, deleted 1995]  [Transferred entry: leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.26 created 1972, deleted 1978 [transferred to EC 3.4.21.31, deleted 1992]]
[EC 3.4.99.28 Transferred entry: Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]  [EC 3.4.99.28 created 1978, deleted 1992]  [3.4.99.30 Deleted entry: Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry: Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry: tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry: Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry: cathepsin R]  [EC 3.4.99.34 created 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.35 Transferred entry: premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry: leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.27	Deleted entry. Echis carinatus prothrombin-activating proteinase]
[EC 3.4.99.28 created 1978, deleted 1992] [3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I] [EC 3.4.99.29 created 1978, deleted 1992] [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase] [EC 3.4.99.30 created 1978, deleted 1992] [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase] [EC 3.4.99.31 created 1978, deleted 1992] [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase] [EC 3.4.99.32 created 1978, deleted 1992] [3.4.99.33 Deleted entry. cathepsin R] [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]] [3.4.99.34 Deleted entry. mytilidase] [EC 3.4.99.35 created 1981, deleted 1992] [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II] [EC 3.4.99.35 created 1984, deleted 1995] [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.27 created 1978, deleted 1992]
[3.4.99.29 Deleted entry. Myxobacter AL-1 proteinase I]  [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimetoligopeptidase  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.35 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.28	Transferred entry. Oxyuranus scutellatus prothrombin-activating proteinase. Now EC 3.4.21.60, scutelarin]
[EC 3.4.99.29 created 1978, deleted 1992] [3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992] [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992] [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992] [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.34 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]] [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.28 created 1978, deleted 1992]
[EC 3.4.99.30 Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.29	Deleted entry. Myxobacter AL-1 proteinase I]
[EC 3.4.99.30 created 1978, deleted 1992]  [3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.29 created 1978, deleted 1992]
[3.4.99.31 Transferred entry. tissue endopeptidase degrading collagenase synthetic substrate. Now EC 3.4.24.15, thimet oligopeptidase]  [EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.30	Transferred entry. Myxobacter AL-1 proteinase II. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]
[EC 3.4.99.31 created 1978, deleted 1992]  [3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.30 created 1978, deleted 1992]
[3.4.99.32 Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]  [EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	-	
[EC 3.4.99.32 created 1978, deleted 1992]  [3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.31 created 1978, deleted 1992]
[3.4.99.33 Deleted entry. cathepsin R]  [EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.32	Transferred entry. Armillaria mellea neutral proteinase. Now EC 3.4.24.20, peptidyl-Lys metalloendopeptidase]
[EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]  [3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.32 created 1978, deleted 1992]
[3.4.99.34 Deleted entry. mytilidase]  [EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.33	Deleted entry. cathepsin R]
[EC 3.4.99.34 created 1981, deleted 1992]  [3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.33 created 1981, deleted 1984 [transferred to EC 3.4.21.52, deleted 1992]]
[3.4.99.35 Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]  [EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.34	Deleted entry. mytilidase]
[EC 3.4.99.35 created 1984, deleted 1995]  [3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]		[EC 3.4.99.34 created 1981, deleted 1992]
[3.4.99.36 Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]	[3.4.99.35	Transferred entry. premurein-leader peptidase. Now EC 3.4.23.36, signal peptidase II]
		[EC 3.4.99.35 created 1984, deleted 1995]
[EC 3.4.99.36 created 1984, deleted 1995]	[3.4.99.36	Transferred entry. leader peptidase. Now EC 3.4.21.89, signal peptidase I]
		[EC 3.4.99.36 created 1984, deleted 1995]

[3.4.99.37	Deleted entry. RecA peptidase]
	[EC 3.4.99.37 created 1989, deleted 1992]
[3.4.99.38 verting enzym	Transferred entry. pro-opiomelanotropin-converting proteinase. Now EC 3.4.23.17, pro-opiomelanocortin con- te]
	[EC 3.4.99.38 created 1989, deleted 1992]
[3.4.99.39	Deleted entry. pseudomurein endopeptidase]
	[EC 3.4.99.39 created 1989, deleted 1992]
[3.4.99.40	Deleted entry. Pro-gonadoliberin proteinase]
	[EC 3.4.99.40 created 1989, deleted 1992]
[3.4.99.41	Transferred entry. mitochondrial processing peptidase. Now EC 3.4.24.64, mitochondrial processing peptidase]
	[EC 3.4.99.41 created 1989/90, deleted 1995]
[3.4.99.42	Deleted entry. leucyllysine endopeptidase]
	[EC 3.4.99.42 created 1990, deleted 1992]
[3.4.99.43	Transferred entry. thermopsin. Now EC 3.4.23.42, thermopsin]
	[EC 3.4.99.43 created 1992, deleted 2000]
[3.4.99.44	Transferred entry. pitrilysin. Now EC 3.4.24.55, pitrilysin]
	[EC 3.4.99.44 created 1992, deleted 1993]
[3.4.99.45	Transferred entry. insulinase. Now EC 3.4.24.56, insulysin]
	[EC 3.4.99.45 created 1992, deleted 1993]
[3.4.99.46	Transferred entry. multicatalytic endopeptidase complex. Now EC 3.4.25.1, proteasome endopeptidase complex]
	[EC 3.4.99.46 created 1992, deleted 2000]

# EC 3.5 Acting on carbon-nitrogen bonds, other than peptide bonds

This subclass contains those enzymes that hydrolyse amides, amidines and other C-N bonds. Sub-subclasses are based on the substrate: linear amides (EC 3.5.1), cyclic amides (EC 3.5.2), linear amidines (EC 3.5.3), cyclic amidines (EC 3.5.4), nitriles (EC 3.5.5) and other compounds (EC 3.5.99).

## EC 3.5.1 In linear amides

EC 3.5.1.1

Accepted name: asparaginase

**Reaction:** L-asparagine +  $H_2O$  = L-aspartate +  $NH_3$ 

Other name(s): asparaginase II; L-asparaginase; colaspase; elspar; leunase; crasnitin; α-asparaginase

**Systematic name:** L-asparagine amidohydrolase

**References:** [1031, 1154, 2797]

[EC 3.5.1.1 created 1961]

EC 3.5.1.2

Accepted name: glutaminase

**Reaction:** L-glutamine +  $H_2O$  = L-glutamate +  $NH_3$ 

Other name(s): glutaminase I; L-glutaminase; glutamine aminohydrolase

Systematic name: L-glutamine amidohydrolase

**References:** [1553, 2435]

[EC 3.5.1.2 created 1961]

EC 3.5.1.3

**Accepted name:** ω-amidase

**Reaction:** a monoamide of a dicarboxylate +  $H_2O$  = a dicarboxylate +  $NH_3$ 

Other name(s):  $\alpha$ -keto acid- $\omega$ -amidase

**Systematic name:** ω-amidodicarboxylate amidohydrolase

**Comments:** Acts on glutaramate, succinamate and their 2-oxo derivatives.

**References:** [1881, 1882]

[EC 3.5.1.3 created 1961]

EC 3.5.1.4

Accepted name: amidase

**Reaction:** a monocarboxylic acid amide  $+ H_2O = a$  monocarboxylate  $+ NH_3$ 

Other name(s): acylamidase; acylase (misleading); amidohydrolase (ambiguous); deaminase (ambiguous); fatty acy-

lamidase; *N*-acetylaminohydrolase (ambiguous)

**Systematic name:** acylamide amidohydrolase

**References:** [280, 281]

[EC 3.5.1.4 created 1961, modified 2011]

EC 3.5.1.5

Accepted name: urease

**Reaction:** urea +  $H_2O = CO_2 + 2 NH_3$ **Systematic name:** urea amidohydrolase

**Comments:** A nickel protein. **References:** [620, 2799, 3053]

[EC 3.5.1.5 created 1961]

EC 3.5.1.6

**Accepted name:** β-ureidopropionase

**Reaction:** 3-ureidopropanoate +  $H_2O = \beta$ -alanine +  $CO_2 + NH_3$ 

Other name(s): *N*-carbamoyl-β-alanine amidohydrolase Systematic name: 3-ureidopropanoate amidohydrolase

**Comments:** The animal enzyme also acts on  $\beta$ -ureidoisobutyrate.

**References:** [361, 368, 2961]

[EC 3.5.1.6 created 1961]

EC 3.5.1.7

Accepted name: ureidosuccinase

**Reaction:** N-carbamoyl-L-aspartate +  $H_2O$  = L-aspartate +  $CO_2$  +  $NH_3$ 

**Systematic name:** *N*-carbamoyl-L-aspartate amidohydrolase

**References:** [1683]

## [EC 3.5.1.7 created 1961]

EC 3.5.1.8

Accepted name: formylaspartate deformylase

**Reaction:** N-formyl-L-aspartate +  $H_2O$  = formate + L-aspartate **Other name(s):** formylaspartic formylase (formylase I, formylase II)

**Systematic name:** *N*-formyl-L-aspartate amidohydrolase

**References:** [2163]

[EC 3.5.1.8 created 1961]

EC 3.5.1.9

Accepted name: arylformamidase

**Reaction:** N-formyl-L-kynurenine +  $H_2O$  = formate + L-kynurenine

Other name(s): kynurenine formamidase; formylkynureninase; formylkynurenine formamidase; forma

dase I; formamidase II

**Systematic name:** aryl-formylamine amidohydrolase

**Comments:** Also acts on other aromatic formylamines.

**References:** [1086, 1298, 1878]

[EC 3.5.1.9 created 1961]

EC 3.5.1.10

Accepted name: formyltetrahydrofolate deformylase

**Reaction:** 10-formyltetrahydrofolate +  $H_2O$  = formate + tetrahydrofolate

**Systematic name:** 10-formyltetrahydrofolate amidohydrolase

**References:** [1206]

[EC 3.5.1.10 created 1961]

EC 3.5.1.11

Accepted name: penicillin amidase

**Reaction:** penicillin +  $H_2O$  = a carboxylate + 6-aminopenicillanate

Other name(s): penicillin acylase; benzylpenicillin acylase; novozym 217; semacylase; α-acylamino-β-lactam acylhy-

drolase; ampicillin acylase

Systematic name: penicillin amidohydrolase

References: [2500]

[EC 3.5.1.11 created 1961]

EC 3.5.1.12

Accepted name: biotinidase

**Reaction:** biotin amide  $+ H_2O = biotin + NH_3$ **Other name(s):** amidohydrolase biotinidase

Systematic name: biotin-amide amidohydrolase Comments: Also acts on biotin esters.

**References:** [1479, 2912]

[EC 3.5.1.12 created 1961]

EC 3.5.1.13

Accepted name: aryl-acylamidase

**Reaction:** an anilide  $+ H_2O = a$  carboxylate + aniline

Other name(s): AAA-1; AAA-2; brain acetylcholinesterase (is associated with AAA-2); pseudocholinesterase (asso-

ciated with arylacylamidase)

**Systematic name:** aryl-acylamide amidohydrolase **Comments:** Also acts on 4-substituted anilides.

**References:** [2090]

[EC 3.5.1.13 created 1965]

EC 3.5.1.14

**Accepted name:** *N*-acyl-aliphatic-L-amino acid amidohydrolase

**Reaction:** (1) an N-acyl-aliphatic-L-amino acid +  $H_2O$  = an aliphatic L-amino acid + a carboxylate

(2) an N-acetyl-L-cysteine-S-conjugate +  $H_2O$  = an L-cysteine-S-conjugate + acetate

Other name(s): aminoacylase 1; aminoacylase I; dehydropeptidase II; histozyme; hippuricase; benzamidase; acylase

I; hippurase; amido acid deacylase; L-aminoacylase; acylase; aminoacylase; L-amino-acid acylase; α-N-acylaminoacid hydrolase; long acyl amidoacylase; short acyl amidoacylase; ACY1 (gene name);

N-acyl-L-amino-acid amidohydrolase

**Systematic name:** *N*-acyl-aliphatic-L-amino acid amidohydrolase (carboxylate-forming)

Comments: Contains  $Zn^{2+}$ . The enzyme is found in animals and is involved in the hydrolysis of N-acylated or

N-acetylated amino acids (except L-aspartate). It acts on mercapturic acids (S-conjugates of N-acetyl-L-cysteine) and neutral aliphatic N-acyl- $\alpha$ -amino acids. Some bacterial aminoacylases demonstrate substrate specificity of both EC 3.5.1.14 and EC 3.5.1.114. *cf.* EC 3.5.1.15, aspartoacylase and EC

3.5.1.114, N-acyl-aromatic-L-amino acid amidohydrolase.

**References:** [219, 796, 1117, 1097, 2228, 3024, 1701]

[EC 3.5.1.14 created 1965, modified 2013]

EC 3.5.1.15

Accepted name: aspartoacylase

**Reaction:** N-acyl-L-aspartate +  $H_2O$  = a carboxylate + L-aspartate

Other name(s): aminoacylase II; N-acetylaspartate amidohydrolase; acetyl-aspartic deaminase; acylase II

**Systematic name:** *N*-acyl-L-aspartate amidohydrolase

**References:** [218, 219]

[EC 3.5.1.15 created 1965]

EC 3.5.1.16

**Accepted name:** acetylornithine deacetylase

**Reaction:**  $N^2$ -acetyl-L-ornithine +  $H_2O$  = acetate + L-ornithine

**Other name(s):** acetylornithinase; *N*-acetylornithinase; 2-*N*-acetyl-L-ornithine amidohydrolase

**Systematic name:**  $N^2$ -acetyl-L-ornithine amidohydrolase **Comments:** Also hydrolyses N-acetylmethionine.

**References:** [3072, 3073]

[EC 3.5.1.16 created 1965]

EC 3.5.1.17

Accepted name: acyl-lysine deacylase

**Reaction:**  $N^6$ -acyl-L-lysine + H<sub>2</sub>O = a carboxylate + L-lysine **Other name(s):**  $\varepsilon$ -lysine acylase;  $\varepsilon$ -N-acyl-L-lysine amidohydrolase

**Systematic name:**  $N^6$ -acyl-L-lysine amidohydrolase

**References:** [2223]

## [EC 3.5.1.17 created 1965]

EC 3.5.1.18

Accepted name: succinyl-diaminopimelate desuccinylase

**Reaction:** N-succinyl-LL-2,6-diaminoheptanedioate +  $H_2O$  = succinate + LL-2,6-diaminoheptanedioate

Other name(s): N-succinyl-L- $\alpha$ , $\varepsilon$ -diaminopimelic acid deacylase

**Systematic name:** N-succinyl-LL-2,6-diaminoheptanedioate amidohydrolase

**References:** [1459]

[EC 3.5.1.18 created 1965]

EC 3.5.1.19

Accepted name: nicotinamidase

**Reaction:** nicotinamide +  $H_2O$  = nicotinate +  $NH_3$ 

Other name(s): nicotinamide deaminase; nicotinamide amidase; YNDase

Systematic name: nicotinamide amidohydrolase

**References:** [2268, 2532]

[EC 3.5.1.19 created 1972]

EC 3.5.1.20

Accepted name: citrullinase

**Reaction:** L-citrulline +  $H_2O$  = L-ornithine +  $CO_2$  +  $NH_3$ 

**Other name(s):** citrulline ureidase; citrulline hydrolase; L-citrulline 5-N-carbamoyldihydrolase

**Systematic name:** L-citrulline  $N^5$ -carbamoyldihydrolase

**References:** [1140]

[EC 3.5.1.20 created 1972]

EC 3.5.1.21

**Accepted name:** N-acetyl- $\beta$ -alanine deacetylase

**Reaction:** N-acetyl- $\beta$ -alanine + H<sub>2</sub>O = acetate +  $\beta$ -alanine

**Systematic name:** N-acetyl- $\beta$ -alanine amidohydrolase

**References:** [842]

[EC 3.5.1.21 created 1972]

EC 3.5.1.22

Accepted name: pantothenase

**Reaction:** (R)-pantothenate + H<sub>2</sub>O = (R)-pantoate +  $\beta$ -alanine **Other name(s):** pantothenate hydrolase; pantothenate amidohydrolase

**Systematic name:** (*R*)-pantothenate amidohydrolase

**References:** [2122]

[EC 3.5.1.22 created 1972]

EC 3.5.1.23

Accepted name: ceramidase

**Reaction:** a ceramide  $+ H_2O = a$  carboxylate + sphingosine

Other name(s): acylsphingosine deacylase; glycosphingolipid ceramide deacylase

**Systematic name:** *N*-acylsphingosine amidohydrolase

**References:** [2088, 3279]

## [EC 3.5.1.23 created 1972, modified 1990]

EC 3.5.1.24

Accepted name: choloylglycine hydrolase

**Reaction:** glycocholate +  $H_2O$  = cholate + glycine

Other name(s): glycocholase; bile salt hydrolase; choloyltaurine hydrolase;  $3\alpha$ ,  $7\alpha$ ,  $12\alpha$ -trihydroxy-5 $\beta$ -cholan-24-

oylglycine amidohydrolase

Systematic name: glycocholate amidohydrolase

**Comments:** Also acts on the  $3\alpha$ ,  $12\alpha$ -dihydroxy-derivative, and on choloyl-taurine.

**References:** [2030, 2762]

[EC 3.5.1.24 created 1972]

EC 3.5.1.25

**Accepted name:** *N*-acetylglucosamine-6-phosphate deacetylase

**Reaction:** N-acetyl-D-glucosamine 6-phosphate + H<sub>2</sub>O = D-glucosamine 6-phosphate + acetate

Other name(s): acetylglucosamine phosphate deacetylase; acetylaminodeoxyglucosephosphate acetylhydrolase; 2-

acetamido-2-deoxy-D-glucose-6-phosphate amidohydrolase

**Systematic name:** *N*-acetyl-D-glucosamine-6-phosphate amidohydrolase

**References:** [3174, 3251]

[EC 3.5.1.25 created 1972 (EC 3.5.1.80 created 1999, incorporated 2002)]

EC 3.5.1.26

**Accepted name:**  $N^4$ -( $\beta$ -N-acetylglucosaminyl)-L-asparaginase

**Reaction:**  $N^4$ -( $\beta$ -N-acetyl-D-glucosaminyl)-L-asparagine + H<sub>2</sub>O = N-acetyl- $\beta$ -D-glucosaminylamine + L-

aspartate

Other name(s): aspartylglucosylamine deaspartylase; aspartylglucosylaminase; aspartylglucosylaminidase; as-

partylglycosylamine amidohydrolase; N-aspartyl-β-glucosaminidase; glucosylamidase; β-aspartylglucosylamine amidohydrolase; 4-N-(β-N-acetyl-D-glucosaminyl)-L-asparagine amidohydrolase;

drolase

**Systematic name:**  $N^4$ -( $\beta$ -N-acetyl-D-glucosaminyl)-L-asparagine amidohydrolase

**Comments:** Acts only on asparagine-oligosaccharides containing one amino acid, i.e., the asparagine has free  $\alpha$ -

amino and  $\alpha$ -carboxyl groups [cf. EC 3.5.1.52, peptide- $N^4$ -(N-acetyl- $\beta$ -glucosaminyl)asparagine ami-

dasel

**References:** [1497, 1774, 2887]

[EC 3.5.1.26 created 1972 (EC 3.5.1.37 created 1972, incorporated 1976)]

[3.5.1.27 Deleted entry. N-formylmethionylaminoacyl-tRNA deformylase. The activity is covered by EC 3.5.1.88, peptide deformylase]

[EC 3.5.1.27 created 1972, deleted 2014]

EC 3.5.1.28

**Accepted name:** *N*-acetylmuramoyl-L-alanine amidase

Reaction: Hydrolyses the link between N-acetylmuramoyl residues and L-amino acid residues in certain cell-

wall glycopeptides

**Other name(s):** acetylmuramyl-L-alanine amidase; *N*-acetylmuramyl-L-alanine amidase; *N*-acylmuramyl-L-alanine

amidase; acetylmuramoyl-alanine amidase; *N*-acetylmuramic acid L-alanine amidase; acetylmuramyl-alanine amidase; *N*-acetylmuramylalanine amidase; *N*-acetylmuramoyl-L-alanine

amidase type I; N-acetylmuramoyl-L-alanine amidase type II

**Systematic name:** peptidoglycan amidohydrolase

**References:** [908, 1127, 1126, 3126]

## [EC 3.5.1.28 created 1972 (EC 3.4.19.10 created 1992, incorporated 1997)]

EC 3.5.1.29

Accepted name: 2-(acetamidomethylene)succinate hydrolase

**Reaction:** 2-(acetamidomethylene)succinate +  $2 H_2O$  = acetate + succinate semialdehyde +  $NH_3 + CO_2$ 

Other name(s):  $\alpha$ -(N-acetylaminomethylene)succinic acid hydrolase

**Systematic name:** 2-(acetamidomethylene)succinate amidohydrolase (deaminating, decarboxylating)

**Comments:** Involved in the degradation of pyridoxin in *Pseudomonas*.

**References:** [1212, 2123]

[EC 3.5.1.29 created 1972]

EC 3.5.1.30

**Accepted name:** 5-aminopentanamidase

**Reaction:** 5-aminopentanamide +  $H_2O = 5$ -aminopentanoate +  $NH_3$ 

**Other name(s):** 5-aminovaleramidase; 5-aminonorvaleramidase

**Systematic name:** 5-aminopentanamide amidohydrolase

Comments: The enzyme from *Pseudomonas putida* also acts on 4-aminobutanamide and, more slowly, on 6-

aminohexanamide.

**References:** [2404, 2855]

[EC 3.5.1.30 created 1972, modified 1976]

EC 3.5.1.31

**Accepted name:** formylmethionine deformylase

**Reaction:** N-formyl-L-methionine +  $H_2O$  = formate + L-methionine

**Systematic name:** *N*-formyl-L-methionine amidohydrolase

**References:** [75]

[EC 3.5.1.31 created 1972]

EC 3.5.1.32

Accepted name: hippurate hydrolase

**Reaction:** hippurate + H<sub>2</sub>O = benzoate + glycine **Systematic name:** N-benzoylamino-acid amidohydrolase **Comments:** Acts on various N-benzoylamino acids.

**References:** [2452, 2453]

[EC 3.5.1.32 created 1972]

EC 3.5.1.33

**Accepted name:** *N*-acetylglucosamine deacetylase

**Reaction:** N-acetyl-D-glucosamine +  $H_2O = D$ -glucosamine + acetate

**Other name(s):** acetylaminodeoxyglucose acetylhydrolase; *N*-acetyl-D-glucosaminyl *N*-deacetylase

**Systematic name:** *N*-acetyl-D-glucosamine amidohydrolase

**References:** [2459]

[EC 3.5.1.33 created 1972]

[3.5.1.34 Deleted entry. acetylhistidine deacetylase. Identical with EC 3.4.13.5, Xaa-methyl-His dipeptidase]

[EC 3.5.1.34 created 1972, deleted 1981]

## EC 3.5.1.35

Accepted name: D-glutaminase

**Reaction:** D-glutamine +  $H_2O$  = D-glutamate +  $NH_3$ 

**Systematic name:** D-glutamine amidohydrolase

**References:** [631]

[EC 3.5.1.35 created 1972]

#### EC 3.5.1.36

**Accepted name:** *N*-methyl-2-oxoglutaramate hydrolase

**Reaction:** N-methyl-2-oxoglutaramate +  $H_2O = 2$ -oxoglutarate + methylamine

Other name(s): 5-hydroxy-*N*-methylpyroglutamate synthase Systematic name: *N*-methyl-2-oxoglutaramate methylamidohydrolase

**Comments:** In the reverse reaction, the product cyclizes non-enzymically to 2-hydroxy-*N*-methyl-5-oxo-L-proline.

**References:** [1132, 1133]

[EC 3.5.1.36 created 1972]

[3.5.1.37 Deleted entry. 4-L-aspartylglycosylamine amidohydrolase. Identical with EC 3.5.1.26  $N^4$ -( $\beta$ -N-acetylglucosaminyl)-L-asparaginase]

[EC 3.5.1.37 created 1972, deleted 1976]

## EC 3.5.1.38

Accepted name: glutamin-(asparagin-)ase

**Reaction:** (1) L-glutamine +  $H_2O = L$ -glutamate +  $NH_3$ 

(2) L-asparagine +  $H_2O$  = L-aspartate +  $NH_3$ 

**Other name(s):** glutaminase-asparaginase; ansB (gene name); L-asparagine/L-glutamine amidohydrolase; L-

ASNase/L-GLNase

**Systematic name:** L-glutamine(L-asparagine) amidohydrolase

**Comments:** The enzyme from the bacterium *Achromobacter* hydrolyses L-asparagine at 0.8 of the rate of L-

glutamine; the D-isomers are also hydrolysed, but more slowly. cf. EC 3.5.1.2, glutaminase and EC

3.5.1.1, asparaginase.

**References:** [2436, 2872, 1750, 2205]

[EC 3.5.1.38 created 1976]

## EC 3.5.1.39

Accepted name: alkylamidase

**Reaction:** N-methylhexanamide +  $H_2O$  = hexanoate + methylamine

**Systematic name:** *N*-methylhexanamide amidohydrolase

**Comments:** The enzyme hydrolyses *N*-monosubstituted and *N*,*N*-disubstituted amides, and there is some activity

towards primary amides. It has little or no activity towards short-chain substrates.

**References:** [416]

[EC 3.5.1.39 created 1976]

### EC 3.5.1.40

Accepted name: acylagmatine amidase

**Reaction:** benzoylagmatine  $+ H_2O =$  benzoate + agmatine **Other name(s):** acylagmatine amidohydrolase; acylagmatine deacylase

**Systematic name:** benzoylagmatine amidohydrolase

**Comments:** Also acts on acetylagmatine, propanoylagmatine and bleomycin B2

References: [3009]

## [EC 3.5.1.40 created 1976]

#### EC 3.5.1.41

**Accepted name:** chitin deacetylase

**Reaction:**  $chitin + H_2O = chitosan + acetate$ 

**Systematic name:** chitin amidohydrolase

**Comments:** Hydrolyses the *N*-acetamido groups of *N*-acetyl-D-glucosamine residues in chitin.

**References:** [69]

[EC 3.5.1.41 created 1976]

#### EC 3.5.1.42

Accepted name: nicotinamide-nucleotide amidase

**Reaction:**  $\beta$ -nicotinamide D-ribonucleotide +  $H_2O = \beta$ -nicotinate D-ribonucleotide +  $NH_3$ 

Other name(s): NMN deamidase; nicotinamide mononucleotide deamidase; nicotinamide mononucleotide amidohy-

drolase

**Systematic name:** nicotinamide-D-ribonucleotide amidohydrolase

**Comments:** Also acts more slowly on  $\beta$ -nicotinamide D-ribonucleoside.

**References:** [1235]

[EC 3.5.1.42 created 1976]

#### EC 3.5.1.43

Accepted name: peptidyl-glutaminase

**Reaction:**  $\alpha$ -*N*-peptidyl-L-glutamine + H<sub>2</sub>O =  $\alpha$ -*N*-peptidyl-L-glutamate + NH<sub>3</sub> **Other name(s):** peptidoglutaminase I; peptidoglutaminase; peptidoglutaminase

Systematic name: peptidyl-L-glutamine amidohydrolase

Comments: Specific for the hydrolysis of the  $\gamma$ -amide of glutamine substituted at the  $\alpha$ -amino group, e.g., glycyl-

L-glutamine, N-acetyl-L-glutamine and L-leucylglycyl-L-glutamine.

**References:** [1440]

[EC 3.5.1.43 created 1976]

## EC 3.5.1.44

Accepted name: protein-glutamine glutaminase

**Reaction:** protein L-glutamine +  $H_2O$  = protein L-glutamate +  $NH_3$ 

Other name(s): peptidoglutaminase II; glutaminyl-peptide glutaminase; destabilase; peptidylglutaminase II

**Systematic name:** protein-L-glutamine amidohydrolase

Comments: Specific for the hydrolysis of the  $\gamma$ -amide of glutamine substituted at the carboxyl position or both the

 $\alpha\text{-amino and carboxyl positions, e.g., L-glutaminylglycine and L-phenylalanyl-L-glutaminylglycine.}$ 

**References:** [1440]

[EC 3.5.1.44 created 1976, modified 1983]

[3.5.1.45 Deleted entry. urease (ATP-hydrolysing). Now listed only as EC 6.3.4.6 urea carboxylase]

[EC 3.5.1.45 created 1978, deleted 1986]

#### EC 3.5.1.46

**Accepted name:** 6-aminohexanoate-oligomer exohydrolase

**Reaction:** (1)  $[N-(6-\text{aminohexanoyl})]_n + H_2O = [N-(6-\text{aminohexanoyl})]_{n-1} + 6-\text{aminohexanoate}$ 

(2) N-(6-aminohexanoyl)-6-aminohexanoate +  $H_2O = 2$  6-aminohexanoate

Other name(s): 6-aminohexanoate-dimer hydrolase; nylB (gene name); 6-aminohexanoic acid oligomer hydrolase

(ambiguous); N-(6-aminohexanoyl)-6-aminohexanoate amidohydrolase; nylon-6 hydrolase (ambigu-

ous)

**Systematic name:** N-(6-aminohexanoyl)-6-aminohexanoate exoamidohydrolase

**Comments:** The enzyme is involved in degradation of nylon-6 oligomers. It degrades linear oligomers of 6-

aminohexanoate with a degree of polymerization of 2–20 by exo-type cleavage, removing residues sequentially from the N-terminus. Activity decreases with the increase of the polymerization number of the oligomer. *cf.* EC 3.5.1.117, 6-aminohexanoate-oligomer endohydrolase and EC 3.5.2.12,

6-aminohexanoate-cyclic-dimer hydrolase.

**References:** [1461]

[EC 3.5.1.46 created 1983, modified 2014]

EC 3.5.1.47

**Accepted name:** *N*-acetyldiaminopimelate deacetylase

**Reaction:** N-acetyl-LL-2,6-diaminoheptanedioate +  $H_2O$  = acetate + LL-2,6-diaminoheptanedioate

**Other name(s):** *N*-acetyl-L-diaminopimelic acid deacylase; *N*-acetyl-LL-diaminopimelate deacylase; 6-*N*-acetyl-LL-

2,6-diaminoheptanedioate amidohydrolase

**Systematic name:**  $N^6$ -acetyl-LL-2,6-diaminoheptanedioate amidohydrolase

**References:** [166, 2509, 2803]

[EC 3.5.1.47 created 1984 (EC 3.1.1.62 created 1989, incorporated 1992)]

EC 3.5.1.48

Accepted name: acetylspermidine deacetylase

**Reaction:**  $N^8$ -acetylspermidine + H<sub>2</sub>O = acetate + spermidine

Other name(s):  $N^8$ -monoacetylspermidine deacetylase;  $N^8$ -acetylspermidine deacetylase; N-acetylspermidine

deacetylase; N¹-acetylspermidine amidohydrolase (incorrect); 8-N-acetylspermidine amidohydrolase

**Systematic name:**  $N^8$ -acetylspermidine amidohydrolase

**Comments:** It was initially thought that  $N^1$ -acetylspermidine was the substrate for this deacetylase reaction [1680]

but this has since been disproved by Marchant et al. [1806].

**References:** [1680, 242, 1806]

[EC 3.5.1.48 created 1984, modified 2005]

EC 3.5.1.49

Accepted name: formamidase

**Reaction:** formamide  $+ H_2O = formate + NH_3$ 

**Systematic name:** formamide amidohydrolase

**Comments:** Also acts, more slowly, on acetamide, propanamide and butanamide.

**References:** [460, 821]

[EC 3.5.1.49 created 1984]

EC 3.5.1.50

Accepted name: pentanamidase

**Reaction:** pentanamide +  $H_2O$  = pentanoate +  $NH_3$ 

Other name(s): valeramidase

Systematic name: pentanamide amidohydrolase

Comments: Also acts, more slowly, on other short-chain aliphatic amides. Different from EC 3.5.1.49 formami-

dase.

**References:** [821]

## [EC 3.5.1.50 created 1984]

EC 3.5.1.51

**Accepted name:** 4-acetamidobutyryl-CoA deacetylase

**Reaction:** 4-acetamidobutanoyl-CoA +  $H_2O$  = acetate + 4-aminobutanoyl-CoA

**Other name(s):** aminobutyryl-CoA thiolesterase; deacetylase-thiolesterase

**Systematic name:** 4-acetamidobutanoyl-CoA amidohydrolase

**Comments:** The enzyme also hydrolyses 4-aminobutanoyl-CoA to aminobutanoate and coenzyme A.

**References:** [2168]

[EC 3.5.1.51 created 1984]

EC 3.5.1.52

**Accepted name:** peptide- $N^4$ -(N-acetyl- $\beta$ -glucosaminyl)asparagine amidase

**Reaction:** Hydrolysis of an  $N^4$ -(acetyl- $\beta$ -D-glucosaminyl)asparagine residue in which the glucosamine residue

may be further glycosylated, to yield a (substituted) N-acetyl-β-D-glucosaminylamine and a peptide

containing an aspartate residue

Other name(s): glycopeptida N-glycosidase; glycopeptidase; N-oligosaccharide glycopeptidase; N-glycanase; Jack-

bean glycopeptidase; PNGase A; PNGase F

Systematic name: N-linked-glycopeptide-(N-acetyl-β-D-glucosaminyl)-L-asparagine amidohydrolase

**Comments:** Does not act on (GlcNAc)Asn, because it requires the presence of more than two amino-acid residues

in the substrate [cf. EC 3.5.1.26,  $N^4$ -( $\beta$ -N-acetylglucosaminyl)-L-asparaginase]. The plant enzyme

was previously erroneously listed as EC 3.2.2.18.

**References:** [2295, 2841, 2843, 2886]

[EC 3.5.1.52 created 1984, modified 1989 (EC 3.2.2.18 created 1984, incorporated 1989)]

EC 3.5.1.53

**Accepted name:** *N*-carbamovlputrescine amidase

**Reaction:** N-carbamoylputrescine +  $H_2O$  = putrescine +  $CO_2$  +  $NH_3$ 

Other name(s): carbamoylputrescine hydrolase; NCP Systematic name: *N*-carbamoylputrescine amidohydrolase

**References:** [3253]

[EC 3.5.1.53 created 1986]

EC 3.5.1.54

Accepted name: allophanate hydrolase

**Reaction:** urea-1-carboxylate +  $H_2O = 2 CO_2 + 2 NH_3$ 

Other name(s): allophanate lyase; AtzF; TrzF

Systematic name: urea-1-carboxylate amidohydrolase

**Comments:** Along with EC 3.5.2.15 (cyanuric acid amidohydrolase) and EC 3.5.1.84 (biuret amidohydrolase),

this enzyme forms part of the cyanuric-acid metabolism pathway, which degrades *s*-triazide herbicides, such as atrazine [2-chloro-4-(ethylamino)-6-(isopropylamino)-1,3,5-triazine], in bacteria. The yeast enzyme (but not that from green algae) also catalyses the reaction of EC 6.3.4.6, urea carboxylase, thus bringing about the hydrolysis of urea to CO<sub>2</sub> and NH<sub>3</sub> in the presence of ATP and bicarbonate. The enzyme from *Pseudomonas* sp. strain ADP has a narrow substrate specificity, being unable to use the structurally analogous compounds urea, hydroxyurea or methylcarbamate as substrate [2621].

**References:** [1781, 2456, 2800, 1377, 423, 2621, 2619]

[EC 3.5.1.54 created 1986, modified 2008]

Accepted name: long-chain-fatty-acyl-glutamate deacylase

**Reaction:** N-long-chain-fatty-acyl-L-glutamate +  $H_2O$  = a long-chain carboxylate + L-glutamate

Other name(s): long-chain aminoacylase; long-chain-fatty-acyl-glutamate deacylase; long-chain acylglutamate ami-

dase; N-acyl-D-glutamate deacylase

**Systematic name:** *N*-long-chain-fatty-acyl-L-glutamate amidohydrolase

**Comments:** Does not act on acyl derivates of other amino acids. Optimum chain length of acyl residue is 12 to 16.

**References:** [857]

[EC 3.5.1.55 created 1986]

EC 3.5.1.56

**Accepted name:** *N,N*-dimethylformamidase

**Reaction:** N,N-dimethylformamide +  $H_2O$  = dimethylamine + formate

**Other name(s):** dimethylformamidase; DMFase

**Systematic name:** *N,N*-dimethylformamide amidohydrolase

**Comments:** An iron protein. Also acts on *N*-ethylformamide and *N*-methyl-formamide and, more slowly, on *N*,*N*-

diethylformamide, N,N-dimethylacetamide and unsubstituted acyl amides.

**References:** [2554]

[EC 3.5.1.56 created 1989]

EC 3.5.1.57

Accepted name: tryptophanamidase

**Reaction:** L-tryptophanamide +  $H_2O$  = L-tryptophan +  $NH_3$ 

**Other name(s):** tryptophan aminopeptidase; L-tryptophan aminopeptidase

**Systematic name:** L-tryptophanamide amidohydrolase

**Comments:** Requires Mn<sup>2+</sup>. Also acts on N-ethylformamide and L-tyrosinamide, and on some tryptophan dipep-

tides.

**References:** [1283]

[EC 3.5.1.57 created 1989]

EC 3.5.1.58

**Accepted name:** *N*-benzyloxycarbonylglycine hydrolase

**Reaction:** N-benzyloxycarbonylglycine +  $H_2O$  = benzyl alcohol +  $CO_2$  + glycine

**Other name(s):** benzyloxycarbonylglycine hydrolase;  $N^{\alpha}$ -carbobenzoxyamino acid amidohydrolase;  $N^{\alpha}$ -

benzyloxycarbonyl amino acid urethane hydrolase;  $N^{\alpha}$ -benzyloxycarbonyl amino acid urethane hy-

drolase I

**Systematic name:** *N*-benzyloxycarbonylglycine urethanehydrolase

**Comments:** Also acts, more slowly, on *N*-benzyloxycarbonylalanine, but not on the corresponding derivatives of

other amino acids or on N-benzyloxycarbonylpeptides. Requires  $Co^{2+}$  or  $Zn^{2+}$ . cf. EC 3.5.1.64,  $N^{\alpha}$ -

benzyloxycarbonylleucine hydrolase.

References: [2005]

[EC 3.5.1.58 created 1989]

EC 3.5.1.59

**Accepted name:** *N*-carbamoylsarcosine amidase

**Reaction:** N-carbamoylsarcosine +  $H_2O$  = sarcosine +  $CO_2$  +  $NH_3$ 

Other name(s): carbamovlsarcosine amidase

**Systematic name:** *N*-carbamovlsarcosine amidohydrolase

**References:** [568]

## [EC 3.5.1.59 created 1989]

EC 3.5.1.60

**Accepted name:** N-(long-chain-acyl)ethanolamine deacylase

**Reaction:** N-(long-chain-acyl)ethanolamine +  $H_2O$  = a long-chain carboxylate + ethanolamine

**Other name(s):** *N*-acylethanolamine amidohydrolase; acylethanolamine amidase

**Systematic name:** *N*-(long-chain-acyl)ethanolamine amidohydrolase

**Comments:** Does not act on *N*-acylsphingosine or *N*,*O*-diacylethanolamine.

**References:** [2569]

[EC 3.5.1.60 created 1989]

EC 3.5.1.61

Accepted name: mimosinase

**Reaction:** (S)-2-amino-3-(3-hydroxy-4-oxo-4H-pyridin-1-yl)propanoate +  $H_2O = 3$ -hydroxy-4H-pyrid-4-one +

L-serine

Systematic name: mimosine amidohydrolase

**Comments:** An enzyme from *Leucaena leucocephala* leaf, which also contains the toxic amino acid, mimosine.

References: [2882]

[EC 3.5.1.61 created 1989]

EC 3.5.1.62

Accepted name: acetylputrescine deacetylase

**Reaction:** N-acetylputrescine +  $H_2O$  = acetate + putrescine

**Systematic name:** *N*-acetylputrescine acetylhydrolase

Comments: The enzyme from *Micrococcus luteus* also acts on  $N^8$ -acetylspermidine and acetylcadaverine, but

more slowly.

**References:** [2813]

[EC 3.5.1.62 created 1989]

EC 3.5.1.63

**Accepted name:** 4-acetamidobutyrate deacetylase

**Reaction:** 4-acetamidobutanoate  $+ H_2O =$  acetate + 4-aminobutanoate

**Systematic name:** 4-acetamidobutanoate amidohydrolase

**Comments:** Also acts on N-acetyl- $\beta$ -alanine and 5-acetamidopentanoate.

**References:** [1090]

[EC 3.5.1.63 created 1989]

EC 3.5.1.64

**Accepted name:**  $N^{\alpha}$ -benzyloxycarbonylleucine hydrolase

**Reaction:**  $N^{\alpha}$ -benzyloxycarbonyl-L-leucine + H<sub>2</sub>O = benzyl alcohol + CO<sub>2</sub> + L-leucine

Other name(s): benzyloxycarbonylleucine hydrolase;  $N^{\alpha}$ -benzyloxycarbonyl amino acid urethane hydrolase IV;  $\alpha$ -N-

benzyloxycarbonyl-L-leucine urethanehydrolase

**Systematic name:**  $N^{\alpha}$ -benzyloxycarbonyl-L-leucine urethanehydrolase

**Comments:** Also acts on  $N^{\alpha}$ -t-butoxycarbonyl-L-leucine, and, more slowly, on the corresponding derivatives of

L-aspartate, L-methionine, L-glutamate and L-alanine. cf. EC 3.5.1.58 N-benzyloxycarbonylglycine

hydrolase.

**References:** [1838]

## [EC 3.5.1.64 created 1989]

EC 3.5.1.65

**Accepted name:** theanine hydrolase

**Reaction:**  $N^5$ -ethyl-L-glutamine + H<sub>2</sub>O = L-glutamate + ethylamine

**Other name(s):** L-theanine amidohydrolase; 5-N-ethyl-L-glutamine amidohydrolase

**Systematic name:**  $N^5$ -ethyl-L-glutamine amidohydrolase **Comments:** Also acts on other N-alkyl-L-glutamines.

**References:** [2991]

[EC 3.5.1.65 created 1989]

EC 3.5.1.66

**Accepted name:** 2-(hydroxymethyl)-3-(acetamidomethylene)succinate hydrolase

**Reaction:** 2-(hydroxymethyl)-3-(acetamidomethylene)succinate +  $2 H_2O$  = acetate + 2-(hydroxymethyl)-4-

oxobutanoate +  $NH_3 + CO_2$ 

Other name(s): compound B hydrolase; α-hydroxymethyl-α'-(*N*-acetylaminomethylene)succinic acid hydrolase Systematic name: 2-(hydroxymethyl)-3-(acetamidomethylene)succinate amidohydrolase (deaminating, decarboxylating)

**Comments:** Involved in the degradation of pyridoxin by *Pseudomonas* and *Arthrobacter*.

**References:** [1212]

[EC 3.5.1.66 created 1989]

EC 3.5.1.67

**Accepted name:** 4-methyleneglutaminase

**Reaction:** 4-methylene-L-glutamine +  $H_2O$  = 4-methylene-L-glutamate +  $NH_3$ **Other name(s):** 4-methyleneglutamine deamidase; 4-methyleneglutamine amidohydrolase

**Systematic name:** 4-methylene-L-glutamine amidohydrolase

**References:** [1217]

[EC 3.5.1.67 created 1989]

EC 3.5.1.68

**Accepted name:** *N*-formylglutamate deformylase

**Reaction:** N-formyl-L-glutamate +  $H_2O$  = formate + L-glutamate

**Other name(s):** β-citryl-L-glutamate hydrolase; formylglutamate deformylase; *N*-formylglutamate hydrolase; β-

 $citryl glutamate\ amidase;\ \beta\text{-}citryl\text{-}L\text{-}glutamate\ amidohydrolase};\ \beta\text{-}citryl\text{-}L\text{-}glutamate\ amidase};\ \beta\text{-}citryl\text{-}L\text{-}glutamate\ amid$ 

L-glutamate-hydrolyzing enzyme

**Systematic name:** *N*-formyl-L-glutamate amidohydrolase

 $\label{eq:comments:comments:comments:} The animal enzyme also acts on $\beta$-citryl-L-glutamate and $\beta$-citryl-L-glutamine.$ 

**References:** [1199, 1937]

[EC 3.5.1.68 created 1989]

EC 3.5.1.69

Accepted name: glycosphingolipid deacylase

Reaction: Hydrolysis of gangliosides and neutral glycosphingolipids, releasing fatty acids to form the lyso-

derivatives

Other name(s): glycosphingolipid ceramide deacylase Systematic name: glycosphingolipid amidohydrolase

**Comments:** Does not act on sphingolipids such as ceramide. Not identical with EC 3.5.1.23 ceramidase.

**References:** [1143]

## [EC 3.5.1.69 created 1990]

EC 3.5.1.70

Accepted name: aculeacin-A deacylase

Reaction: Hydrolysis of the amide bond in aculeacin A and related neutral lipopeptide antibiotics, releasing the

long-chain fatty acid side-chain

**Other name(s):** aculeacin A acylase

Systematic name: aculeacin-A amidohydrolase

**References:** [2859]

[EC 3.5.1.70 created 1992]

EC 3.5.1.71

**Accepted name:** *N*-feruloylglycine deacylase

**Reaction:** N-feruloylglycine +  $H_2O$  = ferulate + glycine

Other name(s): *N*-feruloylglycine hydrolase

Systematic name: *N*-feruloylglycine amidohydrolase

Comments: Hydrolyses a range of L-amino acids from the cinnamoyl and substituted cinnamoyl series. Not iden-

tical with EC 3.5.1.14 aminoacylase.

**References:** [1817, 1816]

[EC 3.5.1.71 created 1992]

EC 3.5.1.72

**Accepted name:** D-benzoylarginine-4-nitroanilide amidase

**Reaction:** N-benzoyl-D-arginine-4-nitroanilide +  $H_2O = N$ -benzoyl-D-arginine + 4-nitroaniline

Other name(s): benzoyl-D-arginine arylamidase; D-BAPA-ase Systematic name: *N*-benzoyl-D-arginine-4-nitroanilide amidohydrolase

**References:** [939]

[EC 3.5.1.72 created 1992]

EC 3.5.1.73

Accepted name: carnitinamidase

**Reaction:** L-carnitinamide +  $H_2O$  = L-carnitine +  $NH_3$ 

Other name(s): L-carnitinamidase; carnitine amidase; L-carnitine amidase

**Systematic name:** L-carnitinamide amidohydrolase **Comments:** Does not act on D-carnitinamide.

References: [2051]

[EC 3.5.1.73 created 1992]

EC 3.5.1.74

Accepted name: chenodeoxycholoyltaurine hydrolase

**Reaction:** chenodeoxycholoyltaurine +  $H_2O$  = chenodeoxycholate + taurine

**Systematic name:** chenodeoxycholoyltaurine amidohydrolase

**Comments:** Some other taurine conjugates are hydrolysed, but not glycine conjugates of bile acids.

**References:** [1406]

[EC 3.5.1.74 created 1992]

Accepted name: urethanase

**Reaction:** urethane +  $H_2O$  = ethanol +  $CO_2$  +  $NH_3$ 

**Other name(s):** urethane hydrolase

**Systematic name:** urethane amidohydrolase (decarboxylating)

**References:** [1484]

[EC 3.5.1.75 created 1992]

EC 3.5.1.76

Accepted name: arylalkyl acylamidase

**Reaction:** N-acetylarylalkylamine +  $H_2O$  = arylalkylamine + acetate

Other name(s): aralkyl acylamidase

**Systematic name:** *N*-acetylarylalkylamine amidohydrolase

Comments: Identified in *Pseudomonas putida*. Strict specificity for *N*-acetyl arylalkylamines, including *N*-acetyl-

2-phenylethylamine, *N*-acetyl-3-phenylpropylamine, *N*-acetyldopamine, *N*-acetyl-serotonin and melatonin. It also accepts arylalkyl acetates but not acetanilide derivatives, which are common substrates

of EC 3.5.1.13, aryl acylamidase.

**References:** [2634]

[EC 3.5.1.76 created 1999]

EC 3.5.1.77

**Accepted name:** *N*-carbamoyl-D-amino-acid hydrolase

**Reaction:** an *N*-carbamoyl-D-amino acid +  $H_2O$  = a D-amino acid +  $NH_3$  +  $CO_2$ 

**Other name(s):** D-N-carbamoylase; N-carbamoylase (ambiguous); N-carbamoyl-D-amino acid hydrolase

**Systematic name:** *N*-carbamoyl-D-amino-acid amidohydrolase

Comments: This enzyme, along with EC 3.5.1.87 (*N*-carbamoyl-L-amino-acid hydrolase), EC 5.1.99.5 (hydantoin

racemase) and hydantoinase, forms part of the reaction cascade known as the "hydantoinase process", which allows the total conversion of D,L-5-monosubstituted hydantoins into optically pure D- or L-amino acids [41]. It has strict stereospecificity for *N*-carbamoyl-D-amino acids and does not act upon the corresponding L-amino acids or on the *N*-formyl amino acids, *N*-carbamoyl-sarcosine, -citrulline,

-allantoin and -ureidopropanoate, which are substrates for other amidohydrolases.

**References:** [2146, 41]

[EC 3.5.1.77 created 1999, modified 2008]

EC 3.5.1.78

**Accepted name:** glutathionylspermidine amidase

**Reaction:** glutathionylspermidine +  $H_2O$  = glutathione + spermidine **Other name(s):** glutathionylspermidine amidohydrolase (spermidine-forming) **Systematic name:**  $\gamma$ -L-glutamyl-L-cysteinyl-glycine:spermidine amidase

**Comments:** Spermidine is numbered so that atom N-1 is in the amino group of the aminopropyl part of the

molecule. The enzyme from Escherichia coli is bifunctional and also catalyses the glutathionylsper-

midine synthase (EC 6.3.1.8) reaction, resulting in a net hydrolysis of ATP.

**References:** [257]

[EC 3.5.1.78 created 1999]

EC 3.5.1.79

**Accepted name:** phthalyl amidase

**Reaction:** a phthalylamide  $+ H_2O =$ phthalic acid +a substituted amine

**Systematic name:** phthalyl-amide amidohydrolase

**Comments:** In the entry, "phthalyl" is used to mean "2-carboxybenzoyl". The enzyme from *Xanthobacter agilis* 

hydrolyses phthalylated amino acids, peptides,  $\beta$ -lactams, aromatic and aliphatic amines. The substituent on nitrogen may be an alkyl group, but may also be complex, giving an amino acid or peptide derivative. Substitutions on the phthalyl ring include 6-F, 6-NH<sub>2</sub>, 3-OH, and a nitrogen in the aro-

matic ring ortho to the carboxy group attached to the amine. No cofactors are required

**References:** [291, 230, 495, 290]

[EC 3.5.1.79 created 1999]

[3.5.1.80 Deleted entry. N-acetylgalactosamine-6-phosphate deacetylase. Identical to EC 3.5.1.25, N-acetylglucosamine-6-phosphate deacetylase]

[EC 3.5.1.80 created 1999, deleted 2002]

EC 3.5.1.81

**Accepted name:** *N*-acyl-D-amino-acid deacylase

**Reaction:** N-acyl-D-amino acid +  $H_2O$  = a carboxylate + D-amino acid

**Systematic name:** *N*-acyl-D-amino acid amidohydrolase

**Comments:** The enzyme from *Alcaligenes denitrificans* subsp. *xylosoxydans* and *Alcaligenes xylosoxydans* subsp.

xylosoxydans has wide specificity; hydrolyses N-acyl derivative of neutral D-amino acids. Used in

separating D- and L-amino acids. Requires zinc.

**References:** [3088, 3087]

[EC 3.5.1.81 created 1999]

EC 3.5.1.82

**Accepted name:** *N*-acyl-D-glutamate deacylase

**Reaction:** N-acyl-D-glutamate +  $H_2O$  = a carboxylate + D-glutamate

**Systematic name:** *N*-acyl-D-glutamate amidohydrolase

**Comments:** The enzyme from *Alcaligenes xylosoxydans* subsp. *xylosoxydans* and *Pseudomonas* sp. is specific for

*N*-acyl-D-glutamate. Requires zinc.

**References:** [3086, 3089, 3090]

[EC 3.5.1.82 created 1999]

EC 3.5.1.83

**Accepted name:** *N*-acyl-D-aspartate deacylase

**Reaction:** N-acyl-D-aspartate +  $H_2O$  = a carboxylate + D-aspartate

**Systematic name:** *N*-acyl-D-aspartate amidohydrolase

**Comments:** The enzyme from *Alcaligenes xylosoxydans* subsp. *xylosoxydans* is specific for *N*-acyl-D-aspartate.

Requires zinc.

**References:** [1967, 3091]

[EC 3.5.1.83 created 1999]

EC 3.5.1.84

Accepted name: biuret amidohydrolase

**Reaction:** biuret +  $H_2O$  = urea-1-carboxylate +  $NH_3$ 

Systematic name: biuret amidohydrolase

Comments: Along with EC 3.5.2.15 (cyanuric acid amidohydrolase) and EC 3.5.1.54 (allophanate hydrolase), this

enzyme forms part of the cyanuric-acid metabolism pathway, which degrades s-triazide herbicides, such as atrazine [2-chloro-4-(ethylamino)-6-(isopropylamino)-1,3,5-triazine], in bacteria. Urea-1-carboxylate rather than urea (as was thought previously) is the 2-nitrogen intermediate in cyanuric-acid metabolism in bacteria [423]. The product, urea-1-carboxylate, can spontaneously decarboxylate under acidic conditions to form urea but, under physiological conditions, it can be converted into  ${\rm CO}_2$ 

and ammonia by the action of EC 3.5.1.54 [423].

**References:** [484, 423, 2621]

[EC 3.5.1.84 created 2000, modified 2008]

EC 3.5.1.85

**Accepted name:** (S)-N-acetyl-1-phenylethylamine hydrolase

**Reaction:** N-acetylphenylethylamine +  $H_2O$  = phenylethylamine + acetate

**Systematic name:** (S)-N-acetylphenylethylamine:H<sub>2</sub>O hydrolase

Comments: Inhibited by phenylmethanesulfonyl fluoride. Some related acetylated compounds are hydrolysed

with variable enantiomeric selectivities.

**References:** [317]

[EC 3.5.1.85 created 2000, modified 2002]

EC 3.5.1.86

Accepted name: mandelamide amidase

**Reaction:** (*R*)-mandelamide +  $H_2O = (R)$ -mandelate +  $NH_3$ 

**Other name(s):** Pseudomonas mandelamide hydrolase

Systematic name: mandelamide hydrolase

**References:** [3249]

[EC 3.5.1.86 created 2000]

EC 3.5.1.87

**Accepted name:** *N*-carbamoyl-L-amino-acid hydrolase

**Reaction:** an N-carbamoyl-L-2-amino acid (a 2-ureido carboxylate) +  $H_2O$  = an L-2-amino acid +  $NH_3$  +  $CO_2$ **Other name(s):** N-carbamyl L-amino acid amidohydrolase; N-carbamoyl-L-amino acid amidohydrolase; N-carbamoyl-L-amin

carbamoylase; N-carbamoylase (ambiguous)

**Systematic name:** *N*-carbamoyl-L-amino-acid amidohydrolase

Comments: This enzyme, along with EC 3.5.1.77 (*N*-carbamoyl-D-amino-acid hydrolase), EC 5.1.99.5 (hydantoin

racemase) and hydantoinase, forms part of the reaction cascade known as the "hydantoinase process", which allows the total conversion of D,L-5-monosubstituted hydantoins into optically pure D- or L-amino acids [41]. The enzyme from *Alcaligenes xylosoxidans* has broad specificity for carbamoyl-L-amino acids, although it is inactive on the carbamoyl derivatives of glutamate, aspartate, arginine, tyrosine or tryptophan. The enzyme from *Sinorhizobium meliloti* requires a divalent cation for activity and can hydrolyse *N*-carbamoyl-L-tryptophan as well as *N*-carbamoyl L-amino acids with aliphatic substituents [1822]. The enzyme is inactive on derivatives of D-amino acids. In addition to *N*-carbamoyl L-amino acids, the enzyme can also hydrolyse formyl and acetyl derivatives to varying

degrees [2145, 1822].

**References:** [2145, 1822, 41]

[EC 3.5.1.87 created 2001, modified 2008]

EC 3.5.1.88

**Accepted name:** peptide deformylase

**Reaction:** formyl-L-methionyl peptide  $+ H_2O =$  formate + methionyl peptide

Other name(s): *N*-formylmethionylaminoacyl-tRNA deformylase Systematic name: formyl-L-methionyl peptide amidohydrolase

Comments: Requires Fe(II). Also requires at least a dipeptide for an efficient rate of reaction. N-terminal L-

methionine is a prerequisite for activity but the enzyme has broad specificity at other positions. Dif-

fers in substrate specifity from EC 3.5.1.31 (formylmethionine deformylase).

**References:** [15, 1855, 391, 182, 181, 2361, 986, 2360, 1200, 2356, 916, 2252]

[EC 3.5.1.88 created 2001]

EC 3.5.1.89

**Accepted name:** *N*-acetylglucosaminylphosphatidylinositol deacetylase

**Reaction:** 6-(N-acetyl- $\alpha$ -D-glucosaminyl)-1-phosphatidyl-1D-myo-inositol + H<sub>2</sub>O = 6-( $\alpha$ -D-glucosaminyl)-1-

phosphatidyl-1D-myo-inositol + acetate

**Other name(s):** *N*-acetyl-D-glucosaminylphosphatidylinositol acetylhydrolase; *N*-

acetylglucosaminylphosphatidylinositol de-N-acetylase; GlcNAc-PI de-N-acetylase; GlcNAc-PI

deacetylase; acetylglucosaminylphosphatidylinositol deacetylase

**Systematic name:** 6-(*N*-acetyl-α-D-glucosaminyl)-1-phosphatidyl-1D-*myo*-inositol acetylhydrolase

**Comments:** Involved in the second step of glycosylphosphatidylinositol (GPI) anchor formation in all eukaryotes.

The enzyme appears to be composed of a single subunit (PIG-L in mammalian cells and GPI12 in yeast). In some species, the long-chain *sn*-1-acyl group of the phosphatidyl group is replaced by a

long-chain alkyl or alk-1-enyl group.

**References:** [626, 2045, 3133, 2702]

[EC 3.5.1.89 created 1992 as EC 3.1.1.69, transferred 2002 to EC 3.5.1.89, modified 2002]

EC 3.5.1.90

**Accepted name:** adenosylcobinamide hydrolase

**Reaction:** adenosylcobinamide +  $H_2O$  = adenosylcobyric acid + (R)-1-aminopropan-2-ol

Other name(s): CbiZ; AdoCbi amidohydrolase Systematic name: adenosylcobinamide amidohydrolase

**Comments:** Involved in the salvage pathway of cobinamide in archaea. *Archaea* convert adenosylcobinamide

(AdoCbi) into adenosylcobinamide phosphate (AdoCbi-P) in two steps. First, the amidohydrolase activity of CbiZ cleaves off the aminopropanol moiety of AdoCbi yielding adenosylcobyric acid (AdoCby); second, AdoCby is converted into AdoCbi-P by the action of EC 6.3.1.10,

adenosylcobinamide-phosphate synthase (CbiB).

**References:** [3213]

[EC 3.5.1.90 created 2004]

EC 3.5.1.91

**Accepted name:** *N*-substituted formamide deformylase

**Reaction:** N-benzylformamide +  $H_2O$  = formate + benzylamine

Other name(s): NfdA

**Systematic name:** *N*-benzylformamide amidohydrolase

**Comments:** Zinc is a cofactor. While N-benzylformamide is the best substrate, the enzyme from Arthrobacter

pascens can also act on the *N*-substituted formamides *N*-butylformamide, *N*-allylformamide, *N*-[2-(cyclohex-1-enyl)ethyl]formamide and *N*-(1-phenylethyl)formamide, but much more slowly. Amides

of other acids do not act as substrates.

**References:** [856]

[EC 3.5.1.91 created 2005]

EC 3.5.1.92

Accepted name: pantetheine hydrolase

**Reaction:** (R)-pantetheine +  $H_2O = (R)$ -pantothenate + 2-aminoethanethiol

Other name(s): pantetheinase; vanin; vanin-1 Systematic name: (*R*)-pantetheine amidohydrolase

Comments: The enzyme hydrolyses only one of the amide bonds of pantetheine. The substrate analogues phos-

phopantetheine and CoA are not substrates. The enzyme recycles pantothenate (vitamin B<sub>5</sub>) and pro-

duces 2-aminoethanethiol (cysteamine), a potent anti-oxidant [2287].

**References:** [669, 670, 1805, 103, 2287, 1818, 2220]

[EC 3.5.1.92 created 2006]

EC 3.5.1.93

Accepted name: glutaryl-7-aminocephalosporanic-acid acylase

**Reaction:** (7*R*)-7-(4-carboxybutanamido)cephalosporanate +  $H_2O = (7R)$ -7-aminocephalosporanate + glutarate **Other name(s):** 7β-(4-carboxybutanamido)cephalosporanic acid acylase; cephalosporin C acylase; glutaryl-7-ACA acylase; CA; GCA; GA; cephalosporin acylase; glutaryl-7-aminocephalosporanic acid acylase; GL-7-

ACA acylase

**Systematic name:** (7*R*)-7-(4-carboxybutanamido)cephalosporanate amidohydrolase

Comments: Forms 7-aminocephalosporanic acid, a key intermediate in the synthesis of cephem antibiotics. It re-

acts only weakly with cephalosporin C.

**References:** [1262, 1462, 1955, 1581, 1456, 1204, 1449]

[EC 3.5.1.93 created 2005]

EC 3.5.1.94

**Accepted name:**  $\gamma$ -glutamyl- $\gamma$ -aminobutyrate hydrolase

**Reaction:**  $4-(\gamma-L-glutamylamino)$ butanoate +  $H_2O = 4$ -aminobutanoate + L-glutamate

 $\textbf{Other name(s):} \quad \gamma\text{-glutamyl-GABA hydrolase; PuuD; YcjL; 4-($\gamma$-glutamylamino}) butanoate amidohydrolase; 4-($L-$\gamma$-respond to $L-$\gamma$-respond to $L-$$ 

glutamylamino)butanoate amidohydrolase

**Systematic name:** 4-(γ-L-glutamylamino)butanoate amidohydrolase

**Comments:** Forms part of a putrescine-utilizing pathway in *Escherichia coli*, in which it has been hypothe-

sized that putrescine is first glutamylated to form  $\gamma$ -glutamylatino)butanoate. The enzyme can also catalyse the

reactions of EC 3.5.1.35 (D-glutaminase) and EC 3.5.1.65 (theanine hydrolase).

**References:** [1566]

[EC 3.5.1.94 created 2006, modified 2011]

EC 3.5.1.95

**Accepted name:** *N*-malonylurea hydrolase

**Reaction:** 3-oxo-3-ureidopropanoate +  $H_2O$  = malonate + urea

Other name(s): ureidomalonase

**Systematic name:** 3-oxo-3-ureidopropanoate amidohydrolase (urea- and malonate-forming)

Comments: Forms part of the oxidative pyrimidine-degrading pathway in some microorganisms, along with EC

1.17.99.4 (uracil/thymine dehydrogenase) and EC 3.5.2.1 (barbiturase).

**References:** [2725, 2724]

[EC 3.5.1.95 created 2006]

EC 3.5.1.96

Accepted name: succinylglutamate desuccinylase

**Reaction:** N-succinyl-L-glutamate +  $H_2O$  = succinate + L-glutamate

Other name(s):  $N^2$ -succinylglutamate desuccinylase; SGDS; AstE

**Systematic name:** *N*-succinyl-L-glutamate amidohydrolase

**Comments:** Requires  $Co^{2+}$  for maximal activity [3139].  $N^2$ -Acetylglutamate is not a substrate. This is the final

enzyme in the arginine succinyltransferase (AST) pathway for the catabolism of arginine [3139]. This pathway converts the carbon skeleton of arginine into glutamate, with the concomitant production of ammonia and conversion of succinyl-CoA into succinate and CoA. The five enzymes involved in this pathway are EC 2.3.1.109 (arginine *N*-succinyltransferase), EC 3.5.3.23 (*N*-succinylarginine dihydrolase), EC 2.6.1.11 (acetylornithine transaminase), EC 1.2.1.71 (succinylglutamate-semialdehyde

dehydrogenase) and EC 3.5.1.96 (succinylglutamate desuccinylase).

**References:** [3139, 517, 518, 1276, 2575]

[EC 3.5.1.96 created 2006]

EC 3.5.1.97

Accepted name: acyl-homoserine-lactone acylase

**Reaction:** an *N*-acyl-L-homoserine lactone + H<sub>2</sub>O = L-homoserine lactone + a carboxylate

Other name(s): acyl-homoserine lactone acylase; AHL-acylase; AiiD; N-acyl-homoserine lactone acylase; PA2385

protein; quorum-quenching AHL acylase; quorum-quenching enzyme; QuiP

**Systematic name:** *N*-acyl-L-homoserine-lactone amidohydrolase

**Comments:** Acyl-homoserine lactones (AHLs) are produced by a number of bacterial species and are used by

them to regulate the expression of virulence genes in a process known as quorum-sensing. Each bacterial cell has a basal level of AHL and, once the population density reaches a critical level, it triggers AHL-signalling which, in turn, initiates the expression of particular virulence genes. Plants or animals capable of degrading AHLs would have a therapeutic advantage in avoiding bacterial infection as they could prevent AHL-signalling and the expression of virulence genes in quorum-sensing bacteria. This quorum-quenching enzyme removes the fatty-acid side chain from the homoserine lactone ring of AHL-dependent quorum-sensing signal molecules. It has broad specificity for AHLs with side changes ranging in length from 11 to 14 carbons. Substituents at the 3'-position, as found in *N*-(3-

oxododecanoyl)-L-homoserine lactone, do not affect this activity.

**References:** [1697, 2674]

[EC 3.5.1.97 created 2007]

EC 3.5.1.98

**Accepted name:** histone deacetylase

**Reaction:** Hydrolysis of an  $N^6$ -acetyl-lysine residue of a histone to yield a deacetylated histone

Other name(s): HDAC

Systematic name: histone amidohydrolase

Comments: A class of enzymes that remove acetyl groups from  $N^6$ -acetyl-lysine residues on a histone. The reac-

tion of this enzyme is opposite to that of EC 2.3.1.48, histone acetyltransferase. Histone deacetylases (HDACs) can be organized into three classes, HDAC1, HDAC2 and HDAC3, depending on sequence similarity and domain organization. Histone acetylation plays an important role in regulation of gene expression. In eukaryotes, HDACs play a key role in the regulation of transcription and cell prolifera-

tion [2722]. May be identical to EC 3.5.1.17, acyl-lysine deacylase.

**References:** [1535, 572, 2217, 2722, 770, 2274, 560]

[EC 3.5.1.98 created 2008]

EC 3.5.1.99

Accepted name: fatty acid amide hydrolase

**Reaction:** (1) an and a mide  $+ H_2O =$  arachidonic acid + ethanolamine

(2) oleamide +  $H_2O$  = oleic acid +  $NH_3$ 

**Other name(s):** FAAH; oleamide hydrolase; anandamide amidohydrolase

**Systematic name:** fatty acylamide amidohydrolase

Comments: Integral membrane protein, the enzyme is responsible for the catabolism of neuromodulatory fatty

acid amides, including anandamide and oleamide, occurs in mammalia.

**References:** [253, 2245, 2244]

[EC 3.5.1.99 created 2009]

EC 3.5.1.100

**Accepted name:** (R)-amidase

**Reaction:** (1) (*R*)-piperazine-2-carboxamide +  $H_2O = (R)$ -piperazine-2-carboxylate +  $NH_3$ 

(2)  $\beta$ -alaninamide + H<sub>2</sub>O =  $\beta$ -alanine + NH<sub>3</sub>

Other name(s): R-stereospecific amidase; R-amidase

**Systematic name:** (*R*)-piperazine-2-carboxamide amidohydrolase

**Comments:** In addition (*R*)-piperidine-3-carboxamide is hydrolysed to (*R*)-piperidine-3-carboxylic acid and NH<sub>3</sub>,

and (R)-N-tert-butylpiperazine-2-carboxamide is hydrolysed to (R)-piperazine-2-carboxylic acid and tert-butylamine with lower activity. The enzyme does not act on the other amide substrates which are

hydrolysed by EC 3.5.1.4 (amidase).

**References:** [1502]

[EC 3.5.1.100 created 2009, modified 2011]

EC 3.5.1.101

**Accepted name:** L-proline amide hydrolase

**Reaction:** (1) (S)-piperidine-2-carboxamide +  $H_2O = (S)$ -piperidine-2-carboxylate +  $NH_3$ 

(2) L-prolinamide +  $H_2O$  = L-proline +  $NH_3$ 

**Other name(s):** S-stereoselective piperazine-2-tert-butylcarboxamide hydrolase; LaaA; L-amino acid amidase

**Systematic name:** (S)-piperidine-2-carboxamide amidohydrolase

**References:** [1503]

[EC 3.5.1.101 created 2009]

EC 3.5.1.102

**Accepted name:** 2-amino-5-formylamino-6-ribosylaminopyrimidin-4(3*H*)-one 5'-monophosphate deformylase

**Reaction:** 2-amino-5-formylamino-6-(5-phospho-D-ribosylamino)pyrimidin-4(3H)-one +  $H_2O = 2,5$ -diamino-6-

(5-phospho-D-ribosylamino)pyrimidin-4(3*H*)-one + formate

Other name(s): ArfB

**Systematic name:** 2-amino-5-formylamino-6-(5-phospho-D-ribosylamino)pyrimidin-4(3*H*)-one amidohydrolase **Comments:** The enzyme catalyses the second step in archaeal riboflavin and 7,8-didemethyl-8-hydroxy-5-

deazariboflavin biosynthesis. The first step is catalysed by EC 3.5.4.29 (GTP cyclohydrolase IIa). The

bacterial enzyme, EC 3.5.4.25 (GTP cyclohydrolase II) catalyses both reactions.

**References:** [988]

[EC 3.5.1.102 created 2010, modified 2011]

EC 3.5.1.103

Accepted name: N-acetyl-1-D-myo-inositol-2-amino-2-deoxy-α-D-glucopyranoside deacetylase

**Reaction:**  $1-O-(2-\arctan ido-2-deoxy-\alpha-D-glucopyranosyl)-1D-myo-inositol + H<sub>2</sub>O = <math>1-O-(2-\arcsin o-2-deoxy-\alpha-D-glucopyranosyl)$ 

D-glucopyranosyl)-1D-*myo*-inositol + acetate

Other name(s): MshB

Systematic name: 1-(2-acetamido-2-deoxy-α-D-glucopyranosyl)-1D-*myo*-inositol acetylhydrolase

**Comments:** This enzyme is considered the key enzyme and rate limiting step in the mycothiol biosynthesis path-

way [2380]. In addition to acetylase activity, the enzyme possesses weak activity of EC 3.5.1.115, mycothiol *S*-conjugate amidase, and shares sequence similarity with that enzyme [2076]. The enzyme

requires a divalent transition metal ion for activity, believed to be  $Zn^{2+}$  [1854].

**References:** [2380, 2076, 1854]

## [EC 3.5.1.103 created 2010]

EC 3.5.1.104

Accepted name: peptidoglycan-N-acetylglucosamine deacetylase

peptidoglycan-N-acetyl-D-glucosamine + H<sub>2</sub>O = peptidoglycan-D-glucosamine + acetate **Reaction:** 

HP310; PgdA; SpPgdA; BC1960; peptidoglycan deacetylase; N-acetylglucosamine deacetylase; pep-Other name(s):

tidoglycan GlcNAc deacetylase; peptidoglycan N-acetylglucosamine deacetylase; PG N-deacetylase

**Systematic name:** peptidoglycan-N-acetylglucosamine amidohydrolase

**Comments:** Modification of peptidoglycan by N-deacetylation is an important factor in virulence of Helicobacter

pylori, Listeria monocytogenes and Streptococcus suis [3108, 2312, 778]. The enzyme from Streptococcus pneumoniae is a metalloenzyme using a His-His-Asp zinc-binding triad with a nearby aspartic

acid and histidine acting as the catalytic base and acid, respectively [233].

**References:** [2328, 2970, 233, 3108, 2312, 778]

[EC 3.5.1.104 created 2010]

EC 3.5.1.105

chitin disaccharide deacetylase Accepted name:

N,N'-diacetylchitobiose +  $H_2O = \textit{N}$ -acetyl- $\beta$ -D-glucosaminyl- $(1 \rightarrow 4)$ -D-glucosamine + acetate **Reaction:** Other name(s): chitobiose amidohydolase; COD; chitin oligosaccharide deacetylase; chitin oligosaccharide amidohydolase; 2-(acetylamino)-4-O-[2-(acetylamino)-2-deoxy-β-D-glucopyranosyl]-2-deoxy-D-

glucopyranose acetylhydrolase

*N*,*N*′-diacetylchitobiose acetylhydrolase **Systematic name:** 

> **Comments:** Chitin oligosaccharide deacetylase is a key enzyme in the chitin catabolic cascade of chitinolytic

Vibrio strains. Besides being a nutrient, the heterodisaccharide product 4-O-(N-acetyl-β-Dglucosaminyl)-D-glucosamine is a unique inducer of chitinase production in Vibrio parahemolyticus [1148]. In contrast to EC 3.5.1.41 (chitin deacetylase) this enzyme is specific for the chitin disaccha-

ride [1352, 2160]. It also deacetylates the chitin trisaccharide with lower efficiency [2160]. No activity with higher polymers of GlcNAc [1352, 2160].

[1352, 1148, 2160, 2159] References:

[EC 3.5.1.105 created 2010]

EC 3.5.1.106

Accepted name: N-formylmaleamate deformylase

N-formylmaleamic acid +  $H_2O$  = maleamate + formate **Reaction:** 

Other name(s):

Systematic name: N-formylmaleamic acid amidohydrolase

**Comments:** The reaction is involved in the aerobic catabolism of nicotinic acid.

**References:** [1322]

[EC 3.5.1.106 created 2010]

EC 3.5.1.107

Accepted name: maleamate amidohydrolase

> $maleamate + H_2O = maleate + NH_3$ **Reaction:**

Other name(s): NicF

**Systematic name:** maleamate amidohydrolase

> **Comments:** The reaction is involved in the aerobic catabolism of nicotinic acid.

**References:** [1322]

[EC 3.5.1.107 created 2010]

## EC 3.5.1.108

**Accepted name:** UDP-3-*O*-acyl-*N*-acetylglucosamine deacetylase

**Reaction:** UDP-3-O-[(3R)-3-hydroxymyristoyl]-N-acetyl- $\alpha$ -D-glucosamine + H<sub>2</sub>O = UDP-3-O-[(3R)-3-

hydroxymyristoyl]- $\alpha$ -D-glucosamine + acetate

Other name(s): LpxC protein; LpxC enzyme; LpxC deacetylase; deacetylase LpxC; UDP-3-O-acyl-GlcNAc deacety-

lase; UDP-3-*O*-((*R*)-3-hydroxymyristoyl)-*N*-acetylglucosamine deacetylase; UDP-(3-*O*-acyl)-*N*-acetylglucosamine deacetylase; UDP-3-*O*-(R-3-hydroxymyristoyl)-*N*-acetylglucosamine deacetylase; UDP-(3-*O*-(R-3-hydroxymyristoyl))-*N*-acetylglucosamine deacetylase; UDP-3-*O*-[(3*R*)-3-

hydroxymyristoyl]-*N*-acetylglucosamine amidohydrolase

**Systematic name:** UDP-3-O-[(3R)-3-hydroxymyristoyl]-N-acetyl- $\alpha$ -D-glucosamine amidohydrolase

**Comments:** A zinc protein. The enzyme catalyses a committed step in the biosynthesis of lipid A.

**References:** [1131, 1288, 1214, 3122, 3176, 1943]

[EC 3.5.1.108 created 2010]

#### EC 3.5.1.109

**Accepted name:** sphingomyelin deacylase

**Reaction:** (1) an N-acyl-sphingosylphosphorylcholine +  $H_2O$  = a fatty acid + sphingosylphosphorylcholine

(2) a D-glucosyl-N-acylsphingosine +  $H_2O$  = a fatty acid + D-glucosyl-sphingosine

Other name(s): SM deacylase; GcSM deacylase; glucosylceramide sphingomyelin deacylase; sphingomyelin gluco-

sylceramide deacylase; SM glucosylceramide GCer deacylase; SM-GCer deacylase; SMGCer deacy-

lase

**Systematic name:** *N*-acyl-sphingosylphosphorylcholine amidohydrolase

**Comments:** The enzyme is involved in the sphingolipid metabolism in the epidermis.

**References:** [1048, 1139, 1258]

[EC 3.5.1.109 created 2011]

# EC 3.5.1.110

**Accepted name:** peroxyureidoacrylate/ureidoacrylate amidohydrolase

**Reaction:** (1) (Z)-3-ureidoacrylate peracid +  $H_2O = (Z)$ -3-peroxyaminoacrylate +  $CO_2$  +  $NH_3$  (overall reaction)

(1a) (Z)-3-ureidoacrylate peracid +  $H_2O = (Z)$ -3-peroxyaminoacrylate + carbamate

(1b) carbamate =  $CO_2 + NH_3$  (spontaneous)

(2) (Z)-2-methylureidoacrylate peracid +  $H_2O = (Z)$ -2-methylperoxyaminoacrylate +  $CO_2 + NH_3$  (over-

all reaction)

(2a) (Z)-2-methylureidoacrylate peracid +  $H_2O = (Z)$ -2-methylperoxyaminoacrylate + carbamate

(2b) carbamate =  $CO_2 + NH_3$  (spontaneous)

Other name(s): RutB

**Systematic name:** (*Z*)-3-ureidoacrylate peracid amidohydrolase

**Comments:** The enzyme also shows activity towards ureidoacrylate. Part of the Rut pyrimidine catabolic pathway.

**References:** [1452]

[EC 3.5.1.110 created 2012]

#### EC 3.5.1.111

**Accepted name:** 2-oxoglutaramate amidase

**Reaction:** 2-oxoglutaramate +  $H_2O = 2$ -oxoglutarate +  $NH_3$ 

**Other name(s):** ω-amidase (ambiguous)

**Systematic name:** 5-amino-2,5-dioxopentanoate amidohydrolase

**Comments:** The enzyme, which is highly specific for its substrate, participates in the nicotine degradation pathway

of several Gram-positive bacteria.

**References:** [462]

[EC 3.5.1.111 created 2012]

EC 3.5.1.112

**Accepted name:** 2'-N-acetylparomamine deacetylase

**Reaction:** 2'-N-acetylparomamine +  $H_2O$  = paromamine + acetate **Other name(s):** btrD (gene name); neoL (gene name); kanN (gene name) **Systematic name:** 2'-N-acetylparomamine hydrolase (acetate-forming)

Comments: Involved in the biosynthetic pathways of several clinically important aminocyclitol antibiotics, includ-

ing kanamycin, butirosin, neomycin and ribostamycin. The enzyme from the bacterium *Streptomyces fradiae* can also accept 2"'-acetyl-6"'-hydroxyneomycin C as substrate, *cf.* EC 3.5.1.113, 2"'-acetyl-

6"'-hydroxyneomycin C deacetylase [3288].

**References:** [2967, 3288]

[EC 3.5.1.112 created 2012]

EC 3.5.1.113

**Accepted name:** 2'''-acetyl-6'''-hydroxyneomycin C deacetylase

**Reaction:** 2'''-acetyl-6'''-deamino-6'''-hydroxyneomycin C + H<sub>2</sub>O = 6'''-deamino-6'''-hydroxyneomycin C +

acetate

**Other name(s):** *neoL* (gene name)

**Systematic name:** 2'''-acetyl-6'''-hydroxyneomycin C hydrolase (acetate-forming)

Comments: Involved in the biosynthetic pathway of aminoglycoside antibiotics of the neomycin family. The en-

zyme from the bacterium Streptomyces fradiae also catalyses EC 3.5.1.112, 2'-N-acetylparomamine

deacetylase.

**References:** [3288]

[EC 3.5.1.113 created 2012]

EC 3.5.1.114

**Accepted name:** *N*-acyl-aromatic-L-amino acid amidohydrolase

**Reaction:** (1) an N-acyl-aromatic-L-amino acid +  $H_2O$  = an aromatic-L-amino acid + a carboxylate

(2) an N-acetyl-L-cysteine-S-conjugate +  $H_2O$  = an L-cysteine-S-conjugate + acetate

Other name(s): aminoacylase 3; aminoacylase III; ACY3 (gene name)

**Systematic name:** *N*-acyl-aromatic-L-amino acid amidohydrolase (carboxylate-forming)

**Comments:** This enzyme is found in animals and is involved in the hydrolysis of N-acylated or N-acetylated

amino acids (except L-aspartate). It preferentially deacetylates  $N^{\alpha}$ -acetylated aromatic amino acids and mercapturic acids (S-conjugates of *N*-acetyl-L-cysteine) that are usually not deacetylated by EC 3.5.1.14, *N*-acyl-aliphatic-L-amino acid amidohydrolase. The enzyme is significantly activated by Co<sup>2+</sup> and Ni<sup>2+</sup> [2975]. Some bacterial aminoacylases demonstrate substrate specificity for both EC 3.5.1.14 and EC 3.5.1.114. *cf.* EC 3.5.1.14, *N*-acyl-aliphatic-L-amino acid amidohydrolase and EC

3.5.1.15, aspartoacylase.

**References:** [2334, 2072, 2975, 1196, 2974]

[EC 3.5.1.114 created 2013]

EC 3.5.1.115

**Accepted name:** mycothiol S-conjugate amidase

**Reaction:** a mycothiol S-conjugate +  $H_2O$  = an N-acetyl L-cysteine-S-conjugate + 1-O-(2-amino-2-deoxy- $\alpha$ -D-

 ${\tt glucopyranosyl)-1D-} \textit{myo}\text{-}{\tt inositol}$ 

Other name(s): MCA

**Systematic name:** mycothiol *S*-conjugate 1D-*myo*-inositol 2-amino-2-deoxy-α-D-glucopyranosyl-hydrolase

**Comments:** The enzyme that is found in actinomycetes is involved in the detoxification of oxidizing agents

and electrophilic antibiotics. The enzyme has low activity with 1-O-(2-acetamido-2-deoxy- $\alpha$ -D-glucopyranosyl)-1D-myo-inositol as substrate (cf. EC 3.5.1.103, N-acetyl-1-D-myo-inositol-2-amino-

2-deoxy-α-D-glucopyranoside deacetylase) [2759].

**References:** [2075, 2759]

## [EC 3.5.1.115 created 2013]

EC 3.5.1.116

Accepted name: ureidoglycolate amidohydrolase

**Reaction:** (S)-ureidoglycolate +  $H_2O$  = glyoxylate + 2  $NH_3 + CO_2$ 

Other name(s): ureidoglycolate hydrolase; UAH (gene name)

**Systematic name:** (S)-ureidoglycolate amidohydrolase (decarboxylating)

**Comments:** This plant enzyme is involved in the degradation of ureidoglycolate, an intermediate of purine degra-

dation. Not to be confused with EC 4.3.2.3, ureidoglycolate lyase, which releases urea rather than

ammonia.

**References:** [3197, 3161, 3162]

[EC 3.5.1.116 created 1992 as EC 3.5.3.19, transferred 2014 to EC 3.5.1.116]

EC 3.5.1.117

**Accepted name:** 6-aminohexanoate-oligomer endohydrolase

**Reaction:**  $[N-(6-\text{aminohexanoyl})]_n + \text{H}_2\text{O} = [N-(6-\text{aminohexanoyl})]_{n-x} + [N-(6-\text{aminohexanoyl})]_x$ **Other name(s):** endo-type 6-aminohexanoate oligomer hydrolase; Ahx endo-type-oligomer hydrolase; 6-

aminohexanoate oligomer hydrolase; Ahx-oligomer hydrolase; nylon hydrolase; nylon-oligomer hy-

drolase; NylC; nylon-6 hydrolase (ambiguous)

Systematic name: 6-aminohexanoate oligomer endoamidohydrolase

**Comments:** The enzyme is involved in degradation of nylon-6 oligomers. It degrades linear or cyclic oligomers

of poly(6-aminohexanoate) with a degree of polymerization greater than three (n > 3) by endo-type cleavage, to oligomers of a length of two or more  $(2 \le x < n)$ . It shows negligible activity with N-(6-aminohexanoyl)-6-aminohexanoate (cf. EC 3.5.1.46, 6-aminohexanoate-oligomer exo hydrolase) or with 1,8-diazacyclotetradecane-2,9-dione (cf. EC 3.5.2.12, 6-aminohexanoate-cyclic-dimer hydro-

lase).

**References:** [1358, 3271, 2067]

[EC 3.5.1.117 created 2014]

EC 3.5.1.118

**Accepted name:**  $\gamma$ -glutamyl hercynylcysteine S-oxide hydrolase

**Reaction:**  $\gamma$ -L-glutamyl-S-(hercyn-2-yl)-L-cysteine S-oxide + H<sub>2</sub>O = S-(hercyn-2-yl)-L-cysteine S-oxide + L-

glutamate

**Other name(s):** EgtC

**Systematic name:**  $\gamma$ -glutamyl-S-(hercyn-2-yl)cysteine S-oxide amidohydrolase

**Comments:** The enzyme is part of the biosynthesis pathway of ergothioneine in mycobacteria.

**References:** [2599]

[EC 3.5.1.118 created 2015]

EC 3.5.1.119

Accepted name: Pup amidohydrolase

**Reaction:** [prokaryotic ubiquitin-like protein]-L-glutamine  $+ H_2O = [prokaryotic ubiquitin-like protein]-L-glutamine <math>+ H_2O = [prokaryotic ubiquitin-like protein]$ 

glutamate + NH<sub>3</sub>

**Other name(s):** *dop* (gene name); Pup deamidase; depupylase/deamidase; DPUP; depupylase

Systematic name: [prokaryotic ubiquitin-like protein]-L-glutamine amidohydrolase

**Comments:** The enzyme has been characterized from the bacterium *Mycobacterium tuberculosis*. It catalyses

the hydrolysis of the amido group of the C-terminal glutamine of prokaryotic ubiquitin-like protein (Pup), thus activating it for ligation to target proteins, a process catalysed by EC 6.3.1.19, prokaryotic ubiquitin-like protein ligase. The reaction requires ATP as cofactor but not its hydrolysis. The enzyme also catalyses the hydrolytic cleavage of the bond formed by the ligase, between an  $\epsilon$ -amino group of a lysine residue of the target protein and the  $\gamma$ -carboxylate of the C-terminal glutamate of the prokary-

otic ubiquitin-like protein.

**References:** [2782, 331, 2781]

[EC 3.5.1.119 created 2015]

[3.5.1.120 Transferred entry. 2-aminomuconate deaminase (2-hydroxymuconate-forming). Now EC 3.5.99.11, 2-aminomuconate deaminase (2-hydroxymuconate-forming) ]

[EC 3.5.1.120 created 2016, deleted 2017]

EC 3.5.1.121

**Accepted name:** protein N-terminal asparagine amidohydrolase

**Reaction:** N-terminal L-asparaginyl-[protein] + H<sub>2</sub>O = N-terminal L-aspartyl-[protein] + NH<sub>3</sub>

**Other name(s):** NTAN1 (gene name)

**Systematic name:** protein N-terminal asparagine amidohydrolase

**Comments:** This enzyme participates in the eukaryotic ubiquitin-dependent Arg/N-end rule pathway of protein

degradation, promoting the turnover of intracellular proteins that initiate with Met-Asn. Following the acetylation and removal of the initiator methionine, the exposed N-terminal asparagine is deaminated, resulting in its conversion to L-aspartate. The latter serves as a substrate for EC 2.3.2.8, arginyltransferase, making the protein susceptible to arginylation, polyubiquitination and degradation as specified

by the N-end rule.

**References:** [2770, 979, 367]

[EC 3.5.1.121 created 2016]

EC 3.5.1.122

Accepted name: protein N-terminal glutamine amidohydrolase

**Reaction:** N-terminal L-glutaminyl-[protein] +  $H_2O$  = N-terminal L-glutamyl-[protein] +  $NH_3$ 

Other name(s): NTAQ1 (gene name)

**Systematic name:** protein N-terminal glutamine amidohydrolase

**Comments:** This enzyme participates in the eukaryotic ubiquitin-dependent Arg/N-end rule pathway of protein

degradation, promoting the turnover of intracellular proteins that initiate with Met-Gln. Following the acetylation and removal of the initiator methionine, the exposed N-terminal glutamine is deaminated, resulting in its conversion to L-glutamate. The latter serves as a substrate for EC 2.3.2.8, arginyltransferase, making the protein susceptible to arginylation, polyubiquitination and degradation as specified

by the N-end rule.

**References:** [3110]

[EC 3.5.1.122 created 2016]

EC 3.5.1.123

**Accepted name:** γ-glutamylanilide hydrolase

**Reaction:**  $N^5$ -phenyl-L-glutamine + H<sub>2</sub>O = L-glutamate + aniline

**Other name(s):** *atdA*2 (gene name)

**Systematic name:**  $N^5$ -phenyl-L-glutamine amidohydrolase

**Comments:** The enzyme, characterized from the bacterium *Acinetobacter* sp. YAA, catalyses the opposite re-

action from that cayalysed by EC 6.3.1.18,  $\gamma$ -glutamylanilide synthase, which is part of an aniline degradation pathway. Its purpose is likely to maintain a low concentration of  $N^5$ -phenyl-L-glutamine,

which is potentially toxic.

**References:** [2858]

[EC 3.5.1.123 created 2016]

EC 3.5.1.124

Accepted name: protein deglycase

**Reaction:** (1) an  $N^{\omega}$ -(1-hydroxy-2-oxopropyl)-[protein]-L-arginine + H<sub>2</sub>O = a [protein]-L-arginine + lactate

(2) an  $N^6$ -(1-hydroxy-2-oxopropyl)-[protein]-L-lysine + H<sub>2</sub>O = a [protein]-L-lysine + lactate (3) an S-(1-hydroxy-2-oxopropyl)-[protein]-L-cysteine + H<sub>2</sub>O = a [protein]-L-cysteine + lactate

Other name(s): PARK7 (gene name); DJ-1 protein; yhbO (gene name); yajL (gene name); glyoxylase III (incorrect)

**Systematic name:** a [protein]-L-amino acid-1-hydroxypropan-2-one hydrolase [(R)-lactate-forming]

**Comments:** The enzyme, previously thought to be a glyoxalase, acts on glycated L-arginine, L-lysine, and L-

cysteine residues within proteins that have been attacked and modified by glyoxal or 2-oxopropanal. The attack forms hemithioacetal in the case of cysteines and aminocarbinols in the case of arginines and lysines. The enzyme repairs the amino acids, releasing glycolate or lactate (70-80% (S)-lactate) and (S)-lactate) depending on whether the attacking agent was glyoxal or 2-oxopropanal,

respectively [2421, 1908].

**References:** [1928, 2785, 2421, 1908, 2]

[EC 3.5.1.124 created 2016]

EC 3.5.1.125

**Accepted name:**  $N^2$ -acetyl-L-2,4-diaminobutanoate deacetylase

**Reaction:** (2S)-2-acetamido-4-aminobutanoate +  $H_2O = L-2,4$ -diaminobutanoate + acetate

**Other name(s):** *doeB* (gene name)

**Systematic name:** (2*S*)-2-acetamido-4-aminobutanoate amidohydrolase

**Comments:** The enzyme, found in bacteria, has no activity with (2S)-4-acetamido-2-aminobutanoate (cf. EC

3.5.4.44, ectoine hydrolase).

**References:** [2597]

[EC 3.5.1.125 created 2017]

EC 3.5.1.126

**Accepted name:** oxamate amidohydrolase

**Reaction:** oxamate +  $H_2O$  = oxalate +  $NH_3$ 

Other name(s): HpxW

**Systematic name:** oxamate amidohydrolase

**Comments:** The enzyme has been characterized from the bacterium *Klebsiella pneumoniae*.

**References:** [1137]

[EC 3.5.1.126 created 2017]

EC 3.5.1.127

**Accepted name:** jasmonoyl-L-amino acid hydrolase

**Reaction:** a jasmonoyl-L-amino acid + H<sub>2</sub>O = jasmonate + an L-amino acid **Other name(s):** IAR3 (gene name); ILL4 (gene name); ILL6 (gene name)

**Systematic name:** jasmonoyl-L amino acid amidohydrolase

Comments: This entry includes a family of enzymes that recyle jasmonoyl-amino acid conjugates back

to jasmonates. The enzymes from Arabidopsis thaliana have been shown to also act on 12-

hydroxyjasmonoyl-L-isoleucine, generating tuberonic acid.

**References:** [3179]

[EC 3.5.1.127 created 2017]

## EC 3.5.1.128

Accepted name: deaminated glutathione amidase

**Reaction:** N-(4-oxoglutaryl)-L-cysteinylglycine +  $H_2O$  = 2-oxoglutarate + L-cysteinylglycine

Other name(s): dGSH deaminase; NIT1 (gene name)

**Systematic name:** N-(4-oxoglutaryl)-L-cysteinylglycine amidohydrolase

**Comments:** The enzyme, present in animals, fungi and bacteria, is involved in clearing cells of the toxic com-

pound deaminated glutathione, which can be produced as an unwanted side product by several

transaminases.

**References:** [2257]

[EC 3.5.1.128 created 2018]

EC 3.5.1.129

**Accepted name:**  $N^5$ -(cytidine 5'-diphosphoramidyl)-L-glutamine hydrolase

**Reaction:**  $N^5$ -(cytidine 5'-diphosphoramidyl)-L-glutamine + H<sub>2</sub>O = cytidine 5'-diphosphoramidate + L-

glutamate

Other name(s):  $N^5$ -(cytidine 5'-diphosphoramidyl)-L-glutamine deacylase Systematic name:  $N^5$ -(cytidine 5'-diphosphoramidyl)-L-glutamine amidohydrolase

**Comments:** The enzyme, characterized from the bacterium *Campylobacter jejuni*, is involved in formation of a

unique O-methyl phosphoramidate modification on specific sugar residues within the bacterium's cap-

sular polysaccharides.

**References:** [2896]

[EC 3.5.1.129 created 2018]

# EC 3.5.2 In cyclic amides

## EC 3.5.2.1

Accepted name: barbiturase

**Reaction:** barbiturate +  $H_2O = 3$ -oxo-3-ureidopropanoate

**Systematic name:** barbiturate amidohydrolase (3-oxo-3-ureidopropanoate-forming)

**Comments:** Contains zinc and is specific for barbiturate as substrate [2724]. Forms part of the oxidative

pyrimidine-degrading pathway in some microorganisms, along with EC 1.17.99.4 (uracil/thymine dehydrogenase) and EC 3.5.1.95 (*N*-malonylurea hydrolase). It was previously thought that the end-products of the reaction were malonate and urea but this has since been disproved [2725]. May be involved in the regulation of pyrimidine metabolism, along with EC 2.4.2.9, uracil phosphoribosyl-

transferase.

**References:** [1083, 2725, 2724]

[EC 3.5.2.1 created 1961, modified 2006]

EC 3.5.2.2

Accepted name: dihydropyrimidinase

**Reaction:** 5,6-dihydrouracil +  $H_2O = 3$ -ureidopropanoate

Other name(s): hydantoinase; hydropyrimidine hydrase; hydantoin peptidase; pyrimidine hydrase; D-hydantoinase

**Systematic name:** 5,6-dihydropyrimidine amidohydrolase **Comments:** Also acts on dihydrothymine and hydantoin.

**References:** [305, 673]

[EC 3.5.2.2 created 1961]

EC 3.5.2.3

Accepted name: dihydroorotase

**Reaction:** (*S*)-dihydroorotate  $+ H_2O = N$ -carbamoyl-L-aspartate **Other name(s):** carbamoylaspartic dehydrase; dihydroorotate hydrolase

**Systematic name:** (*S*)-dihydroorotate amidohydrolase

**References:** [485, 1682]

[EC 3.5.2.3 created 1961]

EC 3.5.2.4

**Accepted name:** carboxymethylhydantoinase

**Reaction:** L-5-carboxymethylhydantoin +  $H_2O = N$ -carbamoyl-L-aspartate

**Other name(s):** hydantoin hydrolase

Systematic name: L-5-carboxymethylhydantoin amidohydrolase

References: [1682]

[EC 3.5.2.4 created 1961]

EC 3.5.2.5

Accepted name: allantoinase

**Reaction:** (S)-allantoin +  $H_2O$  = allantoate **Systematic name:** (S)-allantoin amidohydrolase

**References:** [786]

[EC 3.5.2.5 created 1961]

EC 3.5.2.6

**Accepted name: \beta**-lactamase

**Reaction:** a  $\beta$ -lactam + H<sub>2</sub>O = a substituted  $\beta$ -amino acid

Other name(s): penicillinase; cephalosporinase; neutrapen; penicillin $\beta$ -lactamase; exopenicillinase; ampicillinase;

penicillin amido-β-lactamhydrolase; penicillinase I, II; β-lactamase I-III; β-lactamase A, B, C; β-

lactamase AME I; cephalosporin-β-lactamase

**Systematic name:**  $\beta$ -lactam hydrolase

**Comments:** A group of enzymes of varying specificity hydrolysing  $\beta$ -lactams; some act more rapidly on

penicillins, some more rapidly on cephalosporins. The latter were formerly listed as EC 3.5.2.8,

cephalosporinase.

**References:** [454, 1112, 1578, 2306, 2307, 2462]

[EC 3.5.2.6 created 1961, modified 1981 (EC 3.5.2.8 created 1972, incorporated 1978)]

EC 3.5.2.7

Accepted name: imidazolonepropionase

**Reaction:** (S)-3-(5-oxo-4,5-dihydro-3*H*-imidazol-4-yl)propanoate +  $H_2O = N$ -formimidoyl-L-glutamate +  $H^+$ 

Other name(s): 4(5)-imidazolone-5(4)-propionic acid hydrolase; imidazolone propionic acid hydrolase

**Systematic name:** 3-(5-oxo-4,5-dihydro-3*H*-imidazol-4-yl)propanoate amidohydrolase

**References:** [2373, 2705]

[EC 3.5.2.7 created 1965, modified 2001]

[3.5.2.8 Deleted entry. cephalosporinase. Now included with EC 3.5.2.6  $\beta$ -lactamase]

[EC 3.5.2.8 created 1972, deleted 1978]

EC 3.5.2.9

**Accepted name:** 5-oxoprolinase (ATP-hydrolysing)

**Reaction:** ATP + 5-oxo-L-proline +  $2 \text{ H}_2\text{O}$  = ADP + phosphate + L-glutamate

Other name(s): pyroglutamase (ATP-hydrolysing); oxoprolinase; pyroglutamase; 5-oxoprolinase; pyroglutamate hy-

drolase; pyroglutamic hydrolase; L-pyroglutamate hydrolase; 5-oxo-L-prolinase; pyroglutamase

**Systematic name:** 5-oxo-L-proline amidohydrolase (ATP-hydrolysing)

**References:** [3043]

[EC 3.5.2.9 created 1976]

EC 3.5.2.10

**Accepted name:** creatininase

**Reaction:** creatinine  $+ H_2O =$  creatine **Other name(s):** creatinine hydrolase **Systematic name:** creatinine amidohydrolase

**References:** [2989]

[EC 3.5.2.10 created 1978]

EC 3.5.2.11

Accepted name: L-lysine-lactamase

**Reaction:** (S)-2-aminohexano-6-lactam +  $H_2O = L$ -lysine

**Other name(s):** L-α-aminocaprolactam hydrolase; L-lysinamidase; L-lysine-1,6-lactam lactamhydrolase

**Systematic name:** (S)-2-aminohexano-6-lactam lactamhydrolase

**Comments:** Also hydrolyses L-lysinamide.

**References:** [860, 2651]

[EC 3.5.2.11 created 1981, modified 1989]

EC 3.5.2.12

**Accepted name:** 6-aminohexanoate-cyclic-dimer hydrolase

**Reaction:** 1,8-diazacyclotetradecane-2,9-dione +  $H_2O = N$ -(6-aminohexanoyl)-6-aminohexanoate

**Systematic name:** 1,8-diazacyclotetradecane-2,9-dione lactamhydrolase

**Comments:** The cyclic dimer of 6-aminohexanoate is converted to the linear dimer.

**References:** [1460]

[EC 3.5.2.12 created 1983]

EC 3.5.2.13

**Accepted name:** 2,5-dioxopiperazine hydrolase

**Reaction:** 2,5-dioxopiperazine +  $H_2O$  = glycylglycine

Other name(s): cyclo(Gly-Gly) hydrolase; cyclo(glycylglycine) hydrolase

**Systematic name:** 2,5-dioxopiperazine amidohydrolase

**Comments:** Highly specific; does not hydrolyse other dioxopiperazines, glycylglycine, proteins or barbiturates.

**References:** [2818]

[EC 3.5.2.13 created 1989]

EC 3.5.2.14

**Accepted name:** N-methylhydantoinase (ATP-hydrolysing)

**Reaction:** ATP + N-methylhydantoin +  $2 \text{ H}_2\text{O} = \text{ADP} + \text{phosphate} + N$ -carbamoylsarcosine

**Other name(s):** *N*-methylhydantoin amidohydrolase; methylhydantoin amidase; *N*-methylhydantoin hydrolase; *N*-

methylhydantoinase; N-methylimidazolidine-2,4-dione amidohydrolase (ATP-hydrolysing)

**Systematic name:** *N*-methylhydantoin amidohydrolase (ATP-hydrolysing)

**References:** [1451]

[EC 3.5.2.14 created 1989]

EC 3.5.2.15

**Accepted name:** cyanuric acid amidohydrolase **Reaction:** cyanuric acid +  $H_2O$  = biuret +  $CO_2$ 

Other name(s): AtzD

Systematic name: cyanuric acid amidohydrolase

Comments: Along with EC 3.5.1.54 (allophanate hydrolase) and EC 3.5.1.84 (biuret amidohydrolase), this en-

zyme forms part of the cyanuric-acid metabolism pathway, which degrades *s*-triazide herbicides, such as atrazine [2-chloro-4-(ethylamino)-6-(isopropylamino)-1,3,5-triazine], in bacteria. This is a key enzyme in the pathway, catalysing the ring cleavage of cyanuric acid. The enzyme is specific for cyanuric acid as substrate as neither the structurally related compounds ammeline (2,4-diamino-6-hydroxy-*s*-triazine) and ammelide (2-amino-4,6-dihydroxy-*s*-triazine) nor a number of pyrimidine

compounds, such as uracil and cytosine, can act as substrates [1389].

**References:** [676, 675, 1389, 824]

[EC 3.5.2.15 created 2000, modified 2008]

EC 3.5.2.16

Accepted name: maleimide hydrolase

**Reaction:** maleimide  $+ H_2O =$  maleamic acid

Other name(s): imidase; cyclic imide hydrolase; cyclic-imide amidohydrolase (decyclicizing) [misprint]; cyclic-imide

amidohydrolase (decyclizing)

Systematic name: cyclic-imide amidohydrolase (ring-opening)

**Comments:** Succinimide and glutarimide, and sulfur-containing cyclic imides, such as rhodanine, can also act

as substrates for the enzyme from Blastobacter sp. A17p-4. The reverse, cyclization, reaction is also catalysed, but much more slowly. It has lower activity towards cyclic ureides, which are the substrates

of EC 3.5.2.2, dihydropyrimidinase.

**References:** [2147]

[EC 3.5.2.16 created 2001]

EC 3.5.2.17

**Accepted name:** hydroxyisourate hydrolase

**Reaction:** 5-hydroxyisourate +  $H_2O$  = 5-hydroxy-2-oxo-4-ureido-2,5-dihydro-1*H*-imidazole-5-carboxylate

Other name(s): HIUHase; 5-hydroxyisourate hydrolase Systematic name: 5-hydroxyisourate amidohydrolase

**Comments:** The reaction is the first stage in the conversion of 5-hydroxyisourate into S-allantoin. This reaction

will also occur spontaneously but more slowly.

**References:** [2387, 2386, 2531]

[EC 3.5.2.17 created 2004]

EC 3.5.2.18

Accepted name: enamidase

**Reaction:** 6-oxo-1,4,5,6-tetrahydronicotinate +  $2 \text{ H}_2\text{O} = 2$ -formylglutarate + NH<sub>3</sub>

**Systematic name:** 6-oxo-1,4,5,6-tetrahydronicotinate amidohydrolase

**Comments:** Contains iron and  $Zn^{2+}$ . Forms part of the nicotinate-fermentation catabolism pathway in *Eubac*-

terium barkeri. Other enzymes involved in this pathway are EC 1.17.1.5 (nicotinate dehydrogenase), EC 1.3.7.1 (6-hydroxynicotinate reductase), EC 1.1.1.291 (2-hydroxymethylglutarate dehydrogenase), EC 5.4.99.4 (2-methyleneglutarate mutase), EC 5.3.3.6 (methylitaconate  $\Delta$ -isomerase), EC 4.2.1.85

(dimethylmaleate hydratase) and EC 4.1.3.32 (2,3-dimethylmalate lyase).

**References:** [34]

[EC 3.5.2.18 created 2006]

EC 3.5.2.19

Accepted name: streptothricin hydrolase

**Reaction:** streptothricin-F +  $H_2O$  = streptothricin-F acid

Other name(s): *sttH* (gene name)

Systematic name: streptothricin-F hydrolase

Comments: The enzyme also catalyses the hydrolysis of streptothricin-D to streptothricin-D acid [1826]. The en-

zyme is responsible for streptothricin resistance in Streptomyces albulus and Streptomyces noursei

[1826, 1033].

**References:** [1826, 1033]

[EC 3.5.2.19 created 2011]

EC 3.5.2.20

**Accepted name:** isatin hydrolase

**Reaction:** isatin +  $H_2O$  = isatinate **Systematic name:** isatin amidohydrolase

**Comments:** Requires Mn<sup>2+</sup>. This enzyme, found in several bacterial species, is involved in the degradation of

indole-3-acetic acid.

**References:** [2713, 225]

[EC 3.5.2.20 created 2014]

### EC 3.5.3 In linear amidines

EC 3.5.3.1

**Accepted name:** arginase

**Reaction:** L-arginine +  $H_2O$  = L-ornithine + urea

Other name(s): arginine amidinase; canavanase; L-arginase; arginine transamidinase

**Systematic name:** L-arginine amidinohydrolase

**Comments:** Also hydrolyses  $\alpha$ -*N*-substituted L-arginines and canavanine.

**References:** [114, 345, 665, 971, 972]

[EC 3.5.3.1 created 1961]

EC 3.5.3.2

Accepted name: guanidinoacetase

**Reaction:** guanidinoacetate +  $H_2O$  = glycine + urea

Other name(s): glycocyaminase

Systematic name: guanidinoacetate amidinohydrolase

**Comments:** Requires Mn<sup>2+</sup>. **References:** [2444, 3297]

[EC 3.5.3.2 created 1961]

## EC 3.5.3.3

Accepted name: creatinase

**Reaction:** creatine +  $H_2O$  = sarcosine + urea

Systematic name: creatine amidinohydrolase

**References:** [2444, 3304]

[EC 3.5.3.3 created 1961]

### EC 3.5.3.4

Accepted name: allantoicase

**Reaction:** allantoate +  $H_2O = (S)$ -ureidoglycolate + urea

Systematic name: allantoate amidinohydrolase

**Comments:** Also hydrolyses (*R*)-ureidoglycolate to glyoxylate and urea.

**References:** [786, 2963, 3038, 2484]

[EC 3.5.3.4 created 1961]

## EC 3.5.3.5

Accepted name: formimidoylaspartate deiminase

**Reaction:** N-formimidoyl-L-aspartate +  $H_2O = N$ -formyl-L-aspartate +  $NH_3$ 

**Other name(s):** formiminoaspartate deiminase

**Systematic name:** *N*-formimidoyl-L-aspartate iminohydrolase

**References:** [1087]

[EC 3.5.3.5 created 1961, modified 2000]

## EC 3.5.3.6

Accepted name: arginine deiminase

**Reaction:** L-arginine +  $H_2O$  = L-citrulline +  $NH_3$ 

Other name(s): arginine dihydrolase; citrulline iminase; L-arginine deiminase

Systematic name: L-arginine iminohydrolase
Comments: Also acts on canavanine.
References: [2152, 2269, 2377]

[EC 3.5.3.6 created 1961]

## EC 3.5.3.7

Accepted name: guanidinobutyrase

**Reaction:** 4-guanidinobutanoate +  $H_2O = 4$ -aminobutanoate + urea

Other name(s): γ-guanidobutyrase; 4-guanidinobutyrate amidinobutyrase; γ-guanidinobutyrate amidinohydrolase; G-

Base; GBH; guanidinobutyrate ureahydrolase

Systematic name: 4-guanidinobutanoate amidinohydrolase

**Comments:** Requires Mn<sup>2+</sup>. Also acts, very slowly, on 5-guanidinopentanoate and 6-guanidinohexanoate.

**References:** [1959, 2911, 3294, 3295]

[EC 3.5.3.7 created 1972]

### EC 3.5.3.8

Accepted name: formimidoylglutamase

**Reaction:** N-formimidoyl-L-glutamate +  $H_2O = L$ -glutamate + formamide

Other name(s): formiminoglutamase; N-formiminoglutamate hydrolase; N-formimino-L-glutamate formiminohydro-

lase

**Systematic name:** *N*-formimidoyl-L-glutamate formimidoylhydrolase

**References:** [1372, 1752]

[EC 3.5.3.8 created 1972, modified 2000, modified 2001]

EC 3.5.3.9

Accepted name: allantoate deiminase

**Reaction:** allantoate +  $H_2O = (S)$ -ureidoglycine +  $NH_3 + CO_2$ 

**Other name(s):** allantoate amidohydrolase

**Systematic name:** allantoate amidinohydrolase (decarboxylating)

Comments: This enzyme is part of the ureide pathway, which permits certain organisms to recycle the nitrogen in

purine compounds. This enzyme, which liberates ammonia from allantoate, is present in plants and

bacteria. In plants it is localized in the endoplasmic reticulum. Requires manganese.

**References:** [3074, 2613]

[EC 3.5.3.9 created 1972, modified 2010]

EC 3.5.3.10

Accepted name: D-arginase

**Reaction:** D-arginine +  $H_2O$  = D-ornithine + urea

**Systematic name:** D-arginine amidinohydrolase

**References:** [2022]

[EC 3.5.3.10 created 1972]

EC 3.5.3.11

**Accepted name:** agmatinase

**Reaction:** agmatine + H<sub>2</sub>O = putrescine + urea **Other name(s):** agmatine ureohydrolase; SpeB **Systematic name:** agmatine amidinohydrolase

**References:** [1151, 3065]

[EC 3.5.3.11 created 1972]

EC 3.5.3.12

Accepted name: agmatine deiminase

**Reaction:** agmatine +  $H_2O = N$ -carbamoylputrescine +  $NH_3$ 

Other name(s): agmatine amidinohydrolase
Systematic name: agmatine iminohydrolase

**Comments:** The plant enzyme also catalyses the reactions of EC 2.1.3.3 (ornithine carbamoyltransferase), EC

2.1.3.6 (putrescine carbamoyltransferase) and EC 2.7.2.2 (carbamate kinase), thus functioning as a putrescine synthase, converting agmatine and ornithine into putrescine and citrulline, respectively.

**References:** [2701, 2751]

[EC 3.5.3.12 created 1972]

EC 3.5.3.13

Accepted name: formimidoylglutamate deiminase

**Reaction:** N-formimidoyl-L-glutamate +  $H_2O = N$ -formyl-L-glutamate +  $NH_3$ **Other name(s):** formiminoglutamate deiminase; formiminoglutamic iminohydrolase

**Systematic name:** *N*-formimidoyl-L-glutamate iminohydrolase

**References:** [3178]

## [EC 3.5.3.13 created 1975, modified 2000]

EC 3.5.3.14

Accepted name: amidinoaspartase

**Reaction:** N-amidino-L-aspartate +  $H_2O = L$ -aspartate + urea

Other name(s): amidinoaspartic amidinohydrolase

Systematic name: N-amidino-L-aspartate amidinohydrolase

Comments: Also acts slowly on N-amidino-L-glutamate.

**References:** [1919]

[EC 3.5.3.14 created 1976]

EC 3.5.3.15

Accepted name: protein-arginine deiminase

**Reaction:** protein L-arginine +  $H_2O$  = protein L-citrulline +  $NH_3$ 

Other name(s): peptidylarginine deiminase; PAD Systematic name: protein-L-arginine iminohydrolase

**Comments:** Also acts on *N*-acyl-L-arginine and, more slowly, on L-arginine esters.

**References:** [847]

[EC 3.5.3.15 created 1983]

EC 3.5.3.16

Accepted name: methylguanidinase

**Reaction:** methylguanidine +  $H_2O$  = methylamine + urea

**Other name(s):** methylguanidine hydrolase

Systematic name: methylguanidine amidinohydrolase

**Comments:** Acts on some other alkylguanidines, but very slowly.

**References:** [2039]

[EC 3.5.3.16 created 1984]

EC 3.5.3.17

Accepted name: guanidinopropionase

**Reaction:** 3-guanidinopropanoate +  $H_2O = \beta$ -alanine + urea

**Other name(s):** GPase; GPH

**Systematic name:** 3-guanidinopropanoate amidinopropionase

**Comments:** Requires Mn<sup>2+</sup>. Also acts, more slowly, on taurocyamine and 4-guanidinobutanoate.

References: [3296]

[EC 3.5.3.17 created 1989]

EC 3.5.3.18

Accepted name: dimethylargininase

**Reaction:**  $N^{\omega}$ ,  $N^{\omega'}$ -dimethyl-L-arginine + H<sub>2</sub>O = dimethylamine + L-citrulline

**Other name(s):** dimethylarginine dimethylaminohydrolase;  $N^G$ ,  $N^G$ -dimethylarginine dimethylaminohydrolase;

N<sup>G</sup>,N<sup>G</sup>-dimethyl-L-arginine dimethylamidohydrolase; ω,ω'-di-N-methyl-L-arginine dimethylami-

dohydrolase; N<sup>0</sup>,N<sup>0</sup>/-methyl-L-arginine dimethylamidohydrolase (incorrect)

**Systematic name:**  $N^{\omega}$ ,  $N^{\omega'}$ -dimethyl-L-arginine dimethylamidohydrolase

**Comments:** Also acts on  $N^{\omega}$ -methyl-L-arginine.

**References:** [2150]

## [EC 3.5.3.18 created 1992]

## [3.5.3.19 Transferred entry. ureidoglycolate hydrolase. Now EC 3.5.1.116, ureidoglycolate amidohydrolase]

[EC 3.5.3.19 created 1992, deleted 2014]

EC 3.5.3.20

Accepted name: diguanidinobutanase

**Reaction:** 1,4-diguanidinobutane +  $H_2O$  = agmatine + urea

**Systematic name:** 1,4-diguanidinobutane amidinohydrolase

**Comments:** Other diguanidinoalkanes with 3 to 10 methylene groups can also act, but more slowly.

**References:** [3293]

[EC 3.5.3.20 created 1992]

EC 3.5.3.21

Accepted name: methylenediurea deaminase

**Reaction:** methylenediurea +  $2 \text{ H}_2\text{O} = N$ -(hydroxymethyl)urea +  $2 \text{ NH}_3 + \text{CO}_2$  (overall reaction)

(1a) methylenediurea +  $H_2O = N$ -(carboxyaminomethyl)urea +  $NH_3$ 

(1b) N-(carboxyaminomethyl)urea = N-(aminomethyl)urea +  $CO_2$  (spontaneous) (1c) N-(aminomethyl)urea +  $H_2O = N$ -(hydroxymethyl)urea +  $NH_3$  (spontaneous)

**Other name(s):** methylenediurease

Systematic name: methylenediurea aminohydrolase

Comments: Methylenediurea is hydrolysed and decarboxylated to give an aminated methylurea, which then spon-

taneously hydrolyses to hydroxymethylurea. The enzyme from *Ochrobactrum anthropi* also hydrolyses dimethylenetriurea and trimethylenetetraurea as well as ureidoglycolate, which is hydrolysed to urea and glyoxylate, and allantoate, which is hydrolysed to ureidoglycolate, ammonia and carbon

dioxide.

**References:** [1297]

[EC 3.5.3.21 created 1999]

EC 3.5.3.22

Accepted name: proclavaminate amidinohydrolase

**Reaction:** amidinoproclavaminate +  $H_2O$  = proclavaminate + urea

Other name(s): PAH; proclavaminate amidino hydrolase Systematic name: amidinoproclavaminate amidinohydrolase

**Comments:** Forms part of the pathway for the biosythesis of the  $\beta$ -lactamase inhibitor clavulanate in *Strepto*-

myces clavuligerus. It carries out an intermediary reaction between the first reaction of EC 1.14.11.21,

clavaminate synthase, and the second and third reactions of that enzyme. Requires Mn<sup>2+</sup>.

**References:** [2510, 3335, 2955, 3222]

[EC 3.5.3.22 created 2003]

EC 3.5.3.23

**Accepted name:** *N*-succinylarginine dihydrolase

**Reaction:**  $N^2$ -succinyl-L-arginine + 2 H<sub>2</sub>O =  $N^2$ -succinyl-L-ornithine + 2 NH<sub>3</sub> + CO<sub>2</sub>

Other name(s):  $N^2$ -succinylarginine dihydrolase; arginine succinylhydrolase; SADH; AruB; AstB; 2-N-succinyl-L-

arginine iminohydrolase (decarboxylating)

**Systematic name:**  $N^2$ -succinyl-L-arginine iminohydrolase (decarboxylating)

Comments: Arginine,  $N^2$ -acetylarginine and  $N^2$ -glutamylarginine do not act as substrates [3139]. This is the sec-

ond enzyme in the arginine succinyltransferase (AST) pathway for the catabolism of arginine [2575]. This pathway converts the carbon skeleton of arginine into glutamate, with the concomitant production of ammonia and conversion of succinyl-CoA into succinate and CoA. The five enzymes involved in this pathway are EC 2.3.1.109 (arginine *N*-succinyltransferase), EC 3.5.3.23 (*N*-succinylarginine dihydrolase), EC 2.6.1.81 (succinylornithine transaminase), EC 1.2.1.71 (succinylglutamate semialde-

hyde dehydrogenase) and EC 3.5.1.96 (succinylglutamate desuccinylase).

**References:** [2575, 2927, 3139, 517, 1276]

[EC 3.5.3.23 created 2006]

EC 3.5.3.24

**Accepted name:**  $N^1$ -aminopropylagmatine ureohydrolase

**Reaction:**  $N^1$ -aminopropylagmatine +  $H_2O$  = spermidine + urea

**Systematic name:**  $N^1$ -aminopropylagmatine amidinohydrolase

Comments: The enzyme, which has been characterized from the hyperthermophilic archaeon *Pyrococcus ko-*

dakarensis and the thermophilic Gram-negative bacterium Thermus thermophilus, is involved in the

biosynthesis of spermidine.

**References:** [2165, 1975]

[EC 3.5.3.24 created 2013]

EC 3.5.3.25

**Accepted name:**  $N^{\omega}$ -hydroxy-L-arginine amidinohydrolase

**Reaction:**  $N^{\omega}$ -hydroxy-L-arginine + H<sub>2</sub>O = L-ornithine + hydroxyurea

**Other name(s):** *dcsB* (gene name)

**Systematic name:**  $N^{\omega}$ -hydroxy-L-arginine amidinohydrolase

Comments: The enzyme participates in the biosynthetic pathway of D-cycloserine, an antibiotic substance pro-

duced by several Streptomyces species.

**References:** [1547, 1548]

[EC 3.5.3.25 created 2013]

EC 3.5.3.26

**Accepted name:** (S)-ureidoglycine aminohydrolase

**Reaction:** (S)-2-ureidoglycine +  $H_2O = (S)$ -ureidoglycolate +  $NH_3$ 

Other name(s): UGlyAH; UGHY; *ylbA* (gene name)
Systematic name: (S)-ureidoglycine aminohydrolase

**Comments:** Binds Mn<sup>2+</sup>. This enzyme, found in plants and bacteria, is part of the ureide pathway, which enables

the recycling of the nitrogen in purine compounds. In plants it is localized in the endoplasmic reticu-

lum.

**References:** [2613, 3162, 2637]

[EC 3.5.3.26 created 2013]

# EC 3.5.4 In cyclic amidines

EC 3.5.4.1

Accepted name: cytosine deaminase

**Reaction:** cytosine +  $H_2O$  = uracil +  $NH_3$ 

Other name(s): isocytosine deaminase Systematic name: cytosine aminohydrolase **Comments:** Also acts on 5-methylcytosine.

**References:** [465, 1532]

[EC 3.5.4.1 created 1961]

EC 3.5.4.2

Accepted name: adenine deaminase

**Reaction:** adenine +  $H_2O$  = hypoxanthine +  $NH_3$ **Other name(s):** adenase; adenine aminase; ADase

Systematic name: adenine aminohydrolase

**References:** [243, 1124]

[EC 3.5.4.2 created 1961]

EC 3.5.4.3

Accepted name: guanine deaminase

**Reaction:** guanine +  $H_2O$  = xanthine +  $NH_3$  **Other name(s):** guanase; guanine aminase; GAH **Systematic name:** guanine aminohydrolase

**References:** guanine aminonydrolas [1152, 1361, 2348]

[EC 3.5.4.3 created 1961]

EC 3.5.4.4

**Accepted name:** adenosine deaminase

**Reaction:** adenosine  $+ H_2O = inosine + NH_3$  **Other name(s):** deoxyadenosine deaminase **Systematic name:** adenosine aminohydrolase

**References:** [1388, 2320]

[EC 3.5.4.4 created 1961]

EC 3.5.4.5

Accepted name: cytidine deaminase

**Reaction:** (1) cytidine +  $H_2O$  = uridine +  $NH_3$ 

(2) 2'-deoxycytidine +  $H_2O = 2'$ -deoxyuridine +  $NH_3$ 

**Other name(s):** cytosine nucleoside deaminase; (deoxy)cytidine deaminase; *cdd* (gene name); CDA (gene name)

**Systematic name:** cytidine/2'-deoxycytidine aminohydrolase

**Comments:** Contains zinc. Catalyses the deamination of cytidine and 2'-deoxycytidine with similar efficiencies.

The enzyme, which is widely distributed among organisms, is involved in salvage of both exogenous

and endogenous cytidine and 2'-deoxycytidine for UMP synthesis.

**References:** [2434, 3120, 2714, 1591, 3068]

[EC 3.5.4.5 created 1961, modified 2013]

EC 3.5.4.6

Accepted name: AMP deaminase

**Reaction:**  $AMP + H_2O = IMP + NH_3$ 

Other name(s): adenylic acid deaminase; AMP aminase; adenylic deaminase; adenylate deaminase; 5-AMP deami-

nase; adenosine 5-monophosphate deaminase; 5-adenylate deaminase; adenyl deaminase; 5-adenylic acid deaminase; adenosine monophosphate deaminase; adenylate aminohydrolase; adenylate desami-

nase; adenosine 5-phosphate aminohydrolase; 5-adenylate deaminase

**Systematic name:** AMP aminohydrolase

**Comments:** *cf.* EC 3.5.4.17 adenosine-phosphate deaminase. **References:** [1361, 1634, 1635, 1636, 1886, 3000, 3152]

[EC 3.5.4.6 created 1961]

EC 3.5.4.7

Accepted name: ADP deaminase

**Reaction:** ADP +  $H_2O = IDP + NH_3$ 

Other name(s): adenosine diphosphate deaminase; adenosinepyrophosphate deaminase

**Systematic name:** ADP aminohydrolase

**References:** [594]

[EC 3.5.4.7 created 1961]

EC 3.5.4.8

Accepted name: aminoimidazolase

**Reaction:** 4-aminoimidazole +  $H_2O$  = imidazol-4-one +  $NH_3$ 

Other name(s): 4-aminoimidazole hydrolase; 4-aminoimidazole deaminase

**Systematic name:** 4-aminoimidazole aminohydrolase

**Comments:** Requires  $Fe^{2+}$ . This enzyme forms part of the xanthine-degradation pathway in some bacteria. The

product of the reaction, imidazol-4-one, can be converted non-enzymically into formiminoglycine. An enzyme has been identified in *Clostridium cylindrosporum* that can perform this hydrolysis reaction

[811, 3075].

**References:** [2349, 811, 3075, 578]

[EC 3.5.4.8 created 1961]

EC 3.5.4.9

**Accepted name:** methenyltetrahydrofolate cyclohydrolase

**Reaction:** 5,10-methenyltetrahydrofolate +  $H_2O = 10$ -formyltetrahydrofolate

Other name(s): Citrovorum factor cyclodehydrase; cyclohydrolase; formyl-methenyl-methylenetetrahydrofolate syn-

thetase (combined); 5,10-methenyltetrahydrofolate 5-hydrolase (decyclizing)

**Systematic name:** 5,10-methenyltetrahydrofolate 5-hydrolase (ring-opening)

Comments: In eukaryotes, the enzyme occurs as a trifunctional enzyme that also has methylenetetrahydrofolate

dehydrogenase (NADP<sup>+</sup>) (EC 1.5.1.5) and formate—tetrahydrofolate ligase (EC 6.3.4.3) activity. In some prokaryotes, it occurs as a bifunctional enzyme that also has dehydrogenase (EC 1.5.1.5) activ-

ity or formimidoyltetrahydrofolate cyclodeaminase (EC 4.3.1.4) activity.

**References:** [2350, 2827]

[EC 3.5.4.9 created 1961]

EC 3.5.4.10

**Accepted name:** IMP cyclohydrolase

**Reaction:** IMP +  $H_2O$  = 5-formamido-1-(5-phospho-D-ribosyl)imidazole-4-carboxamide

Other name(s): inosinicase; inosinate cyclohydrolase; IMP 1,2-hydrolase (decyclizing)

**Systematic name:** IMP 1,2-hydrolase (ring-opening)

**References:** [780]

[EC 3.5.4.10 created 1961, modified 2000]

EC 3.5.4.11

**Accepted name:** pterin deaminase

**Reaction:** a 2-amino-4-hydroxypteridine +  $H_2O$  = a 2,4-dihydroxypteridine +  $NH_3$ 

Other name(s): acrasinase

**Systematic name:** 2-amino-4-hydroxypteridine aminohydrolase

**Comments:** The animal enzyme is specific for pterin, isoxanthopterin and tetrahydropterin.

**References:** [1654, 2405]

[EC 3.5.4.11 created 1965]

#### EC 3.5.4.12

**Accepted name:** dCMP deaminase

**Reaction:**  $dCMP + H_2O = dUMP + NH_3$ 

**Other name(s):** deoxycytidylate deaminase; deoxy-CMP-deaminase; deoxycytidylate aminohydrolase; deoxycytidine

monophosphate deaminase; deoxycytidine-5'-phosphate deaminase; deoxycytidine-5'-monophosphate

aminohydrolase

**Systematic name:** dCMP aminohydrolase

**Comments:** Also acts on some 5-substituted dCMPs.

**References:** [2550, 2551, 2612]

[EC 3.5.4.12 created 1965]

#### EC 3.5.4.13

**Accepted name:** dCTP deaminase

**Reaction:**  $dCTP + H_2O = dUTP + NH_3$ 

**Other name(s):** deoxycytidine triphosphate deaminase; 5-methyl-dCTP deaminase

Systematic name: dCTP aminohydrolase

**References:** [2934]

[EC 3.5.4.13 created 1972]

[3.5.4.14 Transferred entry. deoxycytidine deaminase. Now included in EC 3.5.4.5, (deoxy)cytidine deaminase]

[EC 3.5.4.14 created 1972, transferred 2013 to EC 3.5.4.5., deleted 2013]

## EC 3.5.4.15

Accepted name: guanosine deaminase

**Reaction:** guanosine +  $H_2O$  = xanthosine +  $NH_3$ 

Other name(s): guanosine aminase

Systematic name: guanosine aminohydrolase

**References:** [1264]

[EC 3.5.4.15 created 1972]

## EC 3.5.4.16

Accepted name: GTP cyclohydrolase I

**Reaction:** GTP +  $H_2O$  = formate + 7,8-dihydroneopterin 3'-triphosphate

**Other name(s):** GTP cyclohydrolase; guanosine triphosphate cyclohydrolase; guanosine triphosphate 8-deformylase;

dihydroneopterin triphosphate synthase; GTP 8-formylhydrolase

**Systematic name:** GTP 7,8-8,9-dihydrolase

Comments: The reaction involves hydrolysis of two C-N bonds and isomerization of the pentose unit; the re-

cyclization may be non-enzymic. This enzyme is involved in the de novo synthesis of tetrahydro-biopterin from GTP, with the other enzymes involved being EC 1.1.1.153 (sepiapterin reductase) and

EC 4.2.3.12 (6-pyruvoyltetrahydropterin synthase) [2805].

**References:** [326, 3208, 2805]

## [EC 3.5.4.16 created 1972]

EC 3.5.4.17

Accepted name: adenosine-phosphate deaminase

**Reaction:** (1) AMP +  $H_2O = IMP + NH_3$ 

(2) ADP +  $H_2O = IDP + NH_3$ (3) ATP +  $H_2O = ITP + NH_3$ 

Other name(s): adenylate deaminase; adenine nucleotide deaminase; adenosine (phosphate) deaminase

**Systematic name:** adenosine-phosphate aminohydrolase

**Comments:** Acts on AMP, ADP, ATP, NAD<sup>+</sup> and adenosine, in decreasing order of activity. The bacterial enzyme

can also accept the deoxy derivatives. cf. EC 3.5.4.6, AMP deaminase.

**References:** [2784, 3278]

[EC 3.5.4.17 created 1972, modified 1980, modified 2014]

EC 3.5.4.18

**Accepted name:** ATP deaminase

**Reaction:** ATP +  $H_2O = ITP + NH_3$ 

**Other name(s):** adenosine triphosphate deaminase

**Systematic name:** ATP aminohydrolase

**References:** [450]

[EC 3.5.4.18 created 1972]

EC 3.5.4.19

Accepted name: phosphoribosyl-AMP cyclohydrolase

**Reaction:**  $1-(5-\text{phospho-}\beta-\text{D-ribosyl})-\text{AMP} + \text{H}_2\text{O} = 1-(5-\text{phospho-}\beta-\text{D-ribosyl})-5-[(5-\text{phospho-}\beta-\text{D-ribosyl})$ 

ribosylamino)methylideneamino]imidazole-4-carboxamide

Other name(s): PRAMP-cyclohydrolase; phosphoribosyladenosine monophosphate cyclohydrolase; 1-(5-phospho-D-

ribosyl)-AMP 1,6-hydrolase

**Systematic name:** 1-(5-phospho-β-D-ribosyl)-AMP 1,6-hydrolase

**Comments:** The *Neurospora crassa* enzyme also catalyses the reactions of EC 1.1.1.23 (histidinol dehydrogenase)

and EC 3.6.1.31 (phosphoribosyl-ATP diphosphatase).

**References:** [1925]

[EC 3.5.4.19 created 1972, modified 1976, modified 1981, modified 2000]

EC 3.5.4.20

**Accepted name:** pyrithiamine deaminase

**Reaction:** 1-(4-amino-2-methylpyrimid-5-ylmethyl)-3-(2-hydroxyethyl)-2-methylpyridinium + H<sub>2</sub>O = 1-(4-methylpyridinium)

hydroxy-2-methylpyrimid-5-ylmethyl)-3-(2-hydroxyethyl)-2-methylpyridinium + NH<sub>3</sub>

Other name(s): 1-(4-amino-2-methylpyrimid-5-ylmethyl)-3-(β-hydroxyethyl)-2-methylpyridinium-bromide aminohy-

drolase

**Systematic name:** 1-(4-amino-2-methylpyrimid-5-ylmethyl)-3-(2-hydroxyethyl)-2-methylpyridinium aminohydrolase

**References:** [2673]

[EC 3.5.4.20 created 1972, modified 2014]

EC 3.5.4.21

Accepted name: creatinine deaminase

**Reaction:** creatinine +  $H_2O = N$ -methylhydantoin +  $NH_3$ **Other name(s):** creatinine hydrolase; creatinine desiminase Systematic name: creatinine iminohydrolase

References: [2823]

[EC 3.5.4.21 created 1972]

EC 3.5.4.22

**Accepted name:** 1-pyrroline-4-hydroxy-2-carboxylate deaminase

**Reaction:** 1-pyrroline-4-hydroxy-2-carboxylate +  $H_2O = 2,5$ -dioxopentanoate +  $NH_3$ 

Other name(s): HPC deaminase; 1-pyrroline-4-hydroxy-2-carboxylate aminohydrolase (decyclizing)

**Systematic name:** 1-pyrroline-4-hydroxy-2-carboxylate aminohydrolase (ring-opening)

**References:** [2671, 2672]

[EC 3.5.4.22 created 1976]

EC 3.5.4.23

Accepted name: blasticidin-S deaminase

**Reaction:** blasticidin  $S + H_2O = deaminohydroxyblasticidin <math>S + NH_3$ 

Systematic name: blasticidin-S aminohydrolase

Comments: Catalyses the deamination of the cytosine moiety of the antibiotics blasticidin S, cytomycin and

acetylblasticidin S.

**References:** [3245]

[EC 3.5.4.23 created 1976]

EC 3.5.4.24

Accepted name: sepiapterin deaminase

**Reaction:** sepiapterin +  $H_2O$  = xanthopterin-B2 +  $NH_3$ 

**Systematic name:** sepiapterin aminohydrolase

**Comments:** Also acts on isosepiapterin, but more slowly.

**References:** [2992]

[EC 3.5.4.24 created 1976]

EC 3.5.4.25

Accepted name: GTP cyclohydrolase II

**Reaction:** GTP + 3 H<sub>2</sub>O = formate + 2,5-diamino-6-hydroxy-4-(5-phospho-D-ribosylamino)pyrimidine +

diphosphate

Other name(s): guanosine triphosphate cyclohydrolase II; GTP-8-formylhydrolase

**Systematic name:** GTP 7,8-8,9-dihydrolase (diphosphate-forming)

Comments: Two C-N bonds are hydrolysed, releasing formate, with simultaneous removal of the terminal diphos-

phate.

**References:** [797]

[EC 3.5.4.25 created 1984, modified 2011]

EC 3.5.4.26

Accepted name: diaminohydroxyphosphoribosylaminopyrimidine deaminase

**Reaction:** 2,5-diamino-6-hydroxy-4-(5-phospho-D-ribosylamino)pyrimidine + H<sub>2</sub>O = 5-amino-6-(5-phospho-D-ribosylamino)

ribosylamino)uracil + NH<sub>3</sub>

Systematic name: 2,5-diamino-6-hydroxy-4-(5-phospho-D-ribosylamino)pyrimidine 2-aminohydrolase

**Comments:** The substrate is the product of EC 3.5.4.25 GTP cyclohydrolase II.

**References:** [332]

## [EC 3.5.4.26 created 1984, modified 2011]

EC 3.5.4.27

Accepted name: methenyltetrahydromethanopterin cyclohydrolase

5,10-methenyl-5,6,7,8-tetrahydromethanopterin +  $H_2O = 5$ -formyl-5,6,7,8-tetrahydromethanopterin **Reaction:** 5,10-methenyltetrahydromethanopterin cyclohydrolase; N<sup>5</sup>,N<sup>10</sup>-methenyltetrahydromethanopterin cy-Other name(s):

clohydrolase; methenyl-H<sub>4</sub>MPT cyclohydrolase; 5,10-methenyltetrahydromethanopterin 10-hydrolase

(decyclizing)

**Systematic name:** 5,10-methenyltetrahydromethanopterin 10-hydrolase (ring-opening)

**Comments:** Methanopterin is a pterin analogue. The enzyme is involved in the formation of methane from CO<sub>2</sub> in

Methanobacterium thermoautotrophicum.

References: [640]

[EC 3.5.4.27 created 1989]

EC 3.5.4.28

Accepted name: S-adenosylhomocysteine deaminase

S-adenosyl-L-homocysteine +  $H_2O = S$ -inosyl-L-homocysteine +  $NH_3$ Reaction:

Other name(s): adenosylhomocysteine deaminase

**Systematic name:** S-adenosyl-L-homocysteine aminohydrolase

**References:** [3351]

[EC 3.5.4.28 created 1992]

EC 3.5.4.29

Accepted name: GTP cyclohydrolase IIa

> **Reaction:** GTP + 3 H<sub>2</sub>O = 2-amino-5-formylamino-6-(5-phospho-D-ribosylamino)pyrimidin-4(3H)-one + 2

> > phosphate

**Systematic name:** GTP 8,9-hydrolase (phosphate-forming)

Requires Mg<sup>2+</sup>. This enzyme catalyses the hydrolysis of the imidazole ring of guanosine 5'-**Comments:** 

triphosphate, N<sup>7</sup>-methylguanosine 5'-triphosphate or inosine 5'-triphosphate. Xanthosine 5'triphosphate and ATP are not substrates. It also catalyses the hydrolysis of diphosphate to form two equivalents of phosphate. Unlike GTP cyclohydrolase II (EC 3.5.4.25), this enzyme does not release

formate, but does hydrolyse the diphosphate from GTP to phosphate.

**References:** [963]

[EC 3.5.4.29 created 2003, modified 2011]

EC 3.5.4.30

Accepted name: dCTP deaminase (dUMP-forming)

**Reaction:**  $dCTP + 2 H_2O = dUMP + diphosphate + NH_3$ **Systematic name:** dCTP aminohydrolase (dUMP-forming)

Requires Mg<sup>2+</sup>. Is highly specific for dCTP as substrate as dCMP, CTP, CDP, CMP, cytosine or de-**Comments:** 

oxycytosine are not deaminated. While most bacteria require two enzymes to form dUMP from dCTP (EC 3.5.4.13, dCTP deaminase and EC 3.6.1.23, dUTP diphosphatase), the archaeon Methanocaldococcus jannaschii uses a single enzyme to carry out both functions. This enzyme can also act as a

dUTP diphosphatase, but more slowly.

**References:** [1664]

[EC 3.5.4.30 created 2003]

EC 3.5.4.31

**Accepted name:** S-methyl-5'-thioadenosine deaminase

**Reaction:** S-methyl-5'-thioadenosine +  $H_2O = S$ -methyl-5'-thioinosine +  $NH_3$ 

**Other name(s):** MTA deaminase; 5-methylthioadenosine deaminase

**Systematic name:** S-methyl-5'-thioadenosine amidohydrolase

**Comments:** The enzyme from *Thermotoga maritima* also functions as *S*-adenosylhomocysteine deaminase

(EC 3.5.4.28) and has some activity against adenosine. Adenosine 5'-phosphate and S-adenosyl-L-

methionine (SAM) are not substrates.

**References:** [1130]

[EC 3.5.4.31 created 2011]

EC 3.5.4.32

Accepted name: 8-oxoguanine deaminase

**Reaction:** 8-oxoguanine +  $H_2O$  = urate +  $NH_3$ 

Other name(s): 8-OGD

**Systematic name:** 8-oxoguanine aminohydrolase

Comments:  $Zn^{2+}$  is bound in the active site. 8-Oxoguanine is formed via the oxidation of guanine within DNA

by reactive oxygen species. If uncorrected, this modification leads to the incorporation of 8-oxoG:A

mismatches and eventually to G:C to T:A transversions.

**References:** [1027]

[EC 3.5.4.32 created 2012]

EC 3.5.4.33

**Accepted name:** tRNA(adenine<sup>34</sup>) deaminase

**Reaction:** adenine<sup>34</sup> in tRNA +  $H_2O$  = hypoxanthine<sup>34</sup> in tRNA +  $NH_3$ 

Other name(s): tRNA:A34 deaminase; tadA protein; ADAT2-ADAT3 complex; TADA; tRNA adenosine deaminase

arginine; AtTadA; tadA/ecADAT2; tRNA A:34 deaminase

**Systematic name:** tRNA(adenine<sup>34</sup>) aminohydrolase

**Comments:** The enzyme is involved in editing of tRNA. The active site contains  $Zn^{2+}$  [2737].

**References:** [2737, 575, 1564, 3207, 1633, 2357]

[EC 3.5.4.33 created 2013]

EC 3.5.4.34

**Accepted name:** tRNA<sup>Ala</sup>(adenine<sup>37</sup>) deaminase

**Reaction:** adenine<sup>37</sup> in tRNA<sup>Ala</sup> + H<sub>2</sub>O = hypoxanthine<sup>37</sup> in tRNA<sup>Ala</sup> + NH<sub>3</sub>

**Other name(s):** ADAT1; Tad1p

**Systematic name:** tRNA<sup>Ala</sup>(adenine<sup>37</sup>) aminohydrolase

**Comments:** The enzyme deaminates adenosine<sup>37</sup> to inosine in eukaryotic tRNA<sup>Ala</sup> [1762]. tRNA editing is strictly

dependent on  $Mg^{2+}$  [892].

**References:** [1762, 892, 1410]

[EC 3.5.4.34 created 2013]

EC 3.5.4.35

**Accepted name:** tRNA(cytosine<sup>8</sup>) deaminase

**Reaction:** cytosine<sup>8</sup> in tRNA +  $H_2O$  = uracil<sup>8</sup> in tRNA +  $NH_3$ 

Other name(s): CDAT8

**Systematic name:** tRNA(cytosine<sup>8</sup>) aminohydrolase

**Comments:** The enzyme from *Methanopyrus kandleri* specifically catalyses the deamimation of cytosine at

poition 8 of tRNA in 30 different tRNAs. This cytosine-to-uracil editing guarantees the proper fold-

ing and functionality of the tRNAs.

**References:** [2370]

[EC 3.5.4.35 created 2013]

EC 3.5.4.36

**Accepted name:** mRNA(cytosine<sup>6666</sup>) deaminase

**Reaction:** cytosine<sup>6666</sup> in apolipoprotein B mRNA + H<sub>2</sub>O = uracil<sup>6666</sup> in apolipoprotein B mRNA + NH<sub>3</sub> **Other name(s):** APOBEC-1 (catalytic component of an RNA-editing complex); APOBEC1 (catalytic subunit);

apolipoprotein B mRNA-editing enzyme 1 (catalytic component of an RNA-editing complex); *apoB* mRNA-editing enzyme catalytic polypeptide 1 (catalytic component of an RNA-editing complex);

apoB mRNA editing complex; apolipoprotein B mRNA editing enzyme; REPR

**Systematic name:** mRNA(cytosine<sup>6666</sup>) aminohydrolase

Comments: The apolipoprotein B mRNA editing enzyme complex catalyses the editing of apolipoprotein B

mRNA at cytidine<sup>6666</sup> to uridine, thereby transforming the codon for glutamine-2153 to a termination codon. Editing results in translation of a truncated apolipoprotein B isoform (*apoB*-48) with distinct functions in lipid transport. The catalytic component (APOBEC-1) contains zinc at the active

site [145].

**References:** [427, 846, 145, 426]

[EC 3.5.4.36 created 2013]

EC 3.5.4.37

**Accepted name:** double-stranded RNA adenine deaminase

**Reaction:** adenine in double-stranded RNA +  $H_2O$  = hypoxanthine in double-stranded RNA +  $NH_3$  Other name(s): ADAR; double-stranded RNA adenosine deaminase; dsRAD; dsRNA adenosine deaminase;

DRADA1; double-stranded RNA-specific adenosine deaminase

**Systematic name:** double-stranded RNA adenine aminohydrolase

Comments: This eukaryotic enzyme is involved in RNA editing. It destabilizes double-stranded RNA through

conversion of adenosine to inosine. Inositol hexakisphosphate is required for activity [1763].

**References:** [1187, 2127, 3211, 1763]

[EC 3.5.4.37 created 2013]

EC 3.5.4.38

**Accepted name:** single-stranded DNA cytosine deaminase

**Reaction:** cytosine in single-stranded DNA +  $H_2O$  = uracil in single-stranded DNA +  $NH_3$ 

Other name(s): AID; activation-induced deaminase; AICDA (gene name); activation-induced cytidine deaminase

**Systematic name:** single-stranded DNA cytosine aminohydrolase

**Comments:** The enzyme exclusively catalyses deamination of cytosine in single-stranded DNA. It preferentially

deaminates five-nucleotide bubbles. The optimal target consists of a single-stranded NWRCN motif (W = A or T, R = A or G) [1601]. The enzyme initiates antibody diversification processes by deami-

nating immunoglobulin sequences.

**References:** [2709, 1601, 278, 1600, 3062]

[EC 3.5.4.38 created 2013]

EC 3.5.4.39

Accepted name: GTP cyclohydrolase IV

**Reaction:** GTP +  $H_2O = 7.8$ -dihydroneopterin 2',3'-cyclic phosphate + formate + diphosphate

Other name(s): MptA; GTP cyclohydrolase MptA

**Systematic name:** GTP 7,8-8,9-dihydrolase (cyclizing, formate-releasing, diphosphate-releasing)

**Comments:** Requires  $Fe^{2+}$ . A zinc protein. The enzyme is involved in methanopterin biosynthesis in

methanogenic archaea. cf. GTP cyclohydrolase I (EC 3.5.4.16), GTP cyclohydrolase II (EC 3.5.4.25)

and GTP cyclohydrolase IIa (EC 3.5.4.29).

**References:** [987]

[EC 3.5.4.39 created 2013]

EC 3.5.4.40

Accepted name: aminodeoxyfutalosine deaminase

**Reaction:** 6-amino-6-deoxyfutalosine +  $H_2O$  = futalosine +  $NH_3$ 

**Other name(s):** AFL deaminase; aminofutalosine deaminase; mqnX (gene name)

**Systematic name:** 6-amino-6-deoxyfutalosine deaminase

**Comments:** The enzyme, found in several bacterial species, is part of the futalosine pathway for menaquinone

biosynthesis.

**References:** [65, 935]

[EC 3.5.4.40 created 2014]

EC 3.5.4.41

**Accepted name:** 5'-deoxyadenosine deaminase

**Reaction:** 5'-deoxyadenosine +  $H_2O = 5'$ -deoxyinosine +  $NH_3$ 

Other name(s): MJ1541 (gene name); DadD
Systematic name: 5'-deoxyadenosine aminohydrolase

Comments: The enzyme from the archaeon *Methanocaldococcus jannaschii* is involved in the recycling of 5'-

deoxyadenosine.

**References:** [1912]

[EC 3.5.4.41 created 2014]

EC 3.5.4.42

**Accepted name:** *N*-isopropylammelide isopropylaminohydrolase

**Reaction:** N-isopropylammelide +  $H_2O$  = cyanuric acid + isopropylamine

**Other name(s):** *atzC* (gene name)

**Systematic name:** *N*-isopropylammelide isopropylaminohydrolase

**Comments:** Requires Zn<sup>2+</sup>. This bacterial enzyme is involved in degradation of the herbicide atrazine. It can hy-

drolyse other N-substituted amino dihydroxy-s-triazine molecules, and prefers substrates with linear

*N*-alkyl groups to those with branched alkyl groups.

**References:** [2490, 2620, 130]

[EC 3.5.4.42 created 2000 as EC 3.5.99.4, transferred 2016 to EC 3.5.4.42]

EC 3.5.4.43

Accepted name: hydroxydechloroatrazine ethylaminohydrolase

**Reaction:** hydroxyatrazine +  $H_2O = N$ -isopropylammelide + ethylamine

Other name(s): atzB (gene name); 2,4-dihydroxy-6-(isopropylamino)-1,3,5-triazine ethylaminohydrolase

Systematic name: hydroxyatrazine ethylaminohydrolase

**Comments:** Contains Zn<sup>2+</sup>. This bacterial enzyme is involved in degradation of the herbicide atrazine. The en-

zyme has a broad substrate range, and requires a monohydroxylated s-triazine ring with a minimum of one primary or secondary amine substituent and either a chloride or amine leaving group. It catalyses

both deamination and dechlorination reactions.

**References:** [269, 2601]

[EC 3.5.4.43 created 2000 as EC 3.5.99.3, transferred 2016 to EC 3.5.4.43]

EC 3.5.4.44

Accepted name: ectoine hydrolase

**Reaction:** ectoine +  $H_2O = (2S)$ -2-acetamido-4-aminobutanoate

Other name(s): doeA (gene name)
Systematic name: ectoine aminohydrolase

**Comments:** The enzyme, found in some halophilic bacteria, is involved in the degradation of the compatible so-

lute ectoine. The enzyme, which belongs to peptidase family M24, only acts in the direction of ectoine hydrolysis. It also produces smaller amounts of (2S)-4-acetamido-2-aminobutanoate, which is

recycled back to ectoine by EC 4.2.1.108, ectoine synthase.

**References:** [2597]

[EC 3.5.4.44 created 2017]

EC 3.5.4.45

Accepted name: melamine deaminase

**Reaction:** (1) melamine +  $H_2O$  = ammeline +  $NH_3$ 

(2) ammeline +  $H_2O$  = ammelide +  $NH_3$ 

**Other name(s):** *triA* (gene name)

Systematic name: melamine aminohydrolase

**Comments:** The enzyme, isolated from the bacterium *Acidovorax citrulli*, performs the deamination of melamine

15-fold faster than the deamination of ammeline. It also has activity with 2-chloro-4,6-diamino-s-triazine, but has no activity toward halo-substituted triazine ring compounds such as atrazine (cf. EC

3.8.1.8, atrazine chlorohydrolase).

**References:** [2602]

[EC 3.5.4.45 created 2017]

EC 3.5.4.46

**Accepted name:** cAMP deaminase

**Reaction:** 3',5'-cyclic AMP +  $H_2O = 3',5'$ -cyclic IMP +  $NH_3$ 

Other name(s): cyclic adenylate deaminase; CadD Systematic name: 3′,5′-cyclic AMP aminohydrolase

**Comments:** Requires Zn<sup>2+</sup>. The enzyme, isolated from the bacterium *Leptospira interrogans*, is specific for

cAMP.

**References:** [934]

[EC 3.5.4.46 created 2017]

## EC 3.5.5 In nitriles

EC 3.5.5.1

Accepted name: nitrilase

**Reaction:** a nitrile +  $2 H_2O = a carboxylate + NH_3$ 

Other name(s): acetonitrilase; benzonitrilase Systematic name: nitrile aminohydrolase

**Comments:** Acts on a wide range of aromatic nitriles including (indol-3-yl)acetonitrile, and also on some aliphatic

nitriles, and on the corresponding acid amides. cf. EC 4.2.1.84 nitrile hydratase.

**References:** [1058, 2910, 2220]

[EC 3.5.5.1 created 1965, modified 1989]

EC 3.5.5.2

Accepted name: ricinine nitrilase

**Reaction:** ricinine + 2  $H_2O$  = 3-carboxy-4-methoxy-*N*-methyl-2-pyridone +  $NH_3$ 

**Systematic name:** ricinine aminohydrolase **References:** [2442, 1171, 2220]

[EC 3.5.5.2 created 1972]

[3.5.5.3 Transferred entry. cyanate hydrolase. Now EC 4.2.1.104, cyanate hydratase]

[EC 3.5.5.3 created 1972, deleted 1990]

EC 3.5.5.4

Accepted name: cyanoalanine nitrilase

**Reaction:** 3-cyano-L-alanine +  $2 H_2O = L$ -aspartate +  $NH_3$  (overall reaction)

(1a) 3-cyano-L-alanine +  $H_2O$  = L-asparagine (1b) L-asparagine +  $H_2O$  = L-aspartate +  $NH_3$ 

Other name(s):  $\beta$ -cyanoalanine nitrilase

**Systematic name:** 3-cyano-L-alanine aminohydrolase

**Comments:** L-Asparagine is formed as an intermediate. cf. EC 4.2.1.65, 3-cyanoalanine hydratase and EC 3.5.1.1,

asparaginase.

**References:** [3256]

[EC 3.5.5.4 created 1986]

EC 3.5.5.5

Accepted name: arylacetonitrilase

**Reaction:** 4-chlorophenylacetonitrile + 2 H<sub>2</sub>O = 4-chlorophenylacetate + NH<sub>3</sub>

Systematic name: arylacetonitrile aminohydrolase

**Comments:** Requires thiol compounds. Also hydrolyses other 4-substituted phenylacetonitriles, thien-2-

ylacetonitrile, tolylacetonitriles, and, more slowly, benzyl cyanide.

**References:** [1848, 2024]

[EC 3.5.5.5 created 1992]

EC 3.5.5.6

Accepted name: bromoxynil nitrilase

**Reaction:** 3,5-dibromo-4-hydroxybenzonitrile + **2** H<sub>2</sub>O = 3,5-dibromo-4-hydroxybenzoate + NH<sub>3</sub>

**Systematic name:** 3,5-dibromo-4-hydroxybenzonitrile aminohydrolase

**Comments:** Involved in the bacterial degradation of the herbicide bromoxynil. Highly specific.

**References:** [2754]

[EC 3.5.5.6 created 1992]

EC 3.5.5.7

Accepted name: aliphatic nitrilase

**Reaction:** R-CN +  $\mathbf{2}$  H<sub>2</sub>O = R-COOH + NH<sub>3</sub> **Systematic name:** aliphatic nitrile aminohydrolase

**Comments:** Preferentially hydrolyses aliphatic nitriles, some of which are apparently not substrates for other

known nitrilases (EC 3.5.5.1). Substrates include crotononitrile, acrylonitrile and glutaronitrile.

**References:** [1491, 2220]

[EC 3.5.5.7 created 1999]

EC 3.5.5.8

Accepted name: thiocyanate hydrolase

**Reaction:** thiocyanate +  $2 H_2O$  = carbonyl sulfide +  $NH_3 + HO^-$ 

**Systematic name:** thiocyanate aminohydrolase

**Comments:** The enzyme from *Thiobacillus thioparus* catalyses the first step in the degradation of thiocyanate.

**References:** [1394, 1395]

[EC 3.5.5.8 created 2000]

## EC 3.5.99 In other compounds

EC 3.5.99.1

Accepted name: riboflavinase

**Reaction:** riboflavin +  $H_2O$  = ribitol + lumichrome

**Systematic name:** riboflavin hydrolase

**References:** [3254]

[EC 3.5.99.1 created 1961]

EC 3.5.99.2

Accepted name: aminopyrimidine aminohydrolase

**Reaction:** (1) 4-amino-5-aminomethyl-2-methylpyrimidine +  $H_2O = 4$ -amino-5-hydroxymethyl-2-

methylpyrimidine + NH<sub>3</sub>

(2) thiamine +  $H_2O$  = 4-amino-5-hydroxymethyl-2-methylpyrimidine + 5-(2-hydroxyethyl)-4-

methylthiazole

**Other name(s):** thiaminase; thiaminase II; *tenA* (gene name)

**Systematic name:** 4-amino-5-aminomethyl-2-methylpyrimidine aminohydrolase

**Comments:** Previously known as thiaminase II, this enzyme is involved in the regeneration of the thiamine pyrimi-

dine from degraded products, rather than in thiamine degradation, and participates in thiamine salvage

pathways.

**References:** [849, 1231, 2940, 191, 1317, 1318, 809]

[EC 3.5.99.2 created 1961, modified 2011]

[3.5.99.3 Transferred entry. hydroxydechloroatrazine ethylaminohydrolase. Now EC 3.5.4.43, hydroxydechloroatrazine ethylaminohydrolase]

[EC 3.5.99.3 created 2000, deleted 2016]

[3.5.99.4 Transferred entry. N-isopropylammelide isopropylaminohydrolase. Now EC 3.5.4.42, N-isopropylammelide isopropylaminohydrolase]

[EC 3.5.99.4 created 2000, deleted 2016]

EC 3.5.99.5

**Accepted name:** 2-aminomuconate deaminase

**Reaction:** 2-aminomuconate +  $H_2O = (3E)$ -2-oxohex-3-enedioate +  $NH_3$ 

Other name(s): *amnD* (gene name); *nbaF* (gene name)

Systematic name: 2-aminomuconate aminohydrolase

**Comments:** 2-Aminomuconate is an intermediate in the bacterial biodegradation of nitrobenzene. The enzyme

has been isolated from several species, including *Pseudomonas pseudocaligenes* JS45, *Pseudomonas fluorescens* KU-7, *Pseudomonas* sp. AP3 and *Burkholderia cenocepacia* J2315. The reaction is spon-

taneous in acid conditions.

**References:** [1092, 1093, 2857, 2004]

## [EC 3.5.99.5 created 2000, modified 2012]

EC 3.5.99.6

Accepted name: glucosamine-6-phosphate deaminase

**Reaction:**  $\alpha$ -D-glucosamine 6-phosphate + H<sub>2</sub>O = D-fructose 6-phosphate + NH<sub>3</sub>

Other name(s): glucosaminephosphate isomerase; glucosamine-6-phosphate isomerase; phosphoglucosaminiso-

merase; glucosamine phosphate deaminase; aminodeoxyglucosephosphate isomerase; phosphoglucosamine isomerase; 2-amino-2-deoxy-D-glucose-6-phosphate aminohydrolase (ketol isomerizing)

Systematic name: 2-amino-2-deoxy-α-D-glucose-6-phosphate aminohydrolase (ketol isomerizing)

**Comments:** The enzyme uses ring opening and isomerization of the aldose-ketose type to convert the -CH(-NH<sub>2</sub>)-

CH=O group of glucosamine 6-phosphate into -C(=NH)-CH<sub>2</sub>-OH, forming 2-deoxy-2-imino-D-arabino-hexitol, which then hydrolyses to yield fructose 6-phosphate and ammonia. *N*-Acetyl-D-

glucosamine 6-phosphate, which is not broken down, activates the enzyme.

**References:** [3209, 469, 2246, 1717]

[EC 3.5.99.6 created 1961 as EC 5.3.1.10, transferred 2000 to EC 3.5.99.6]

EC 3.5.99.7

**Accepted name:** 1-aminocyclopropane-1-carboxylate deaminase

**Reaction:** 1-aminocyclopropane-1-carboxylate  $+ H_2O = 2$ -oxobutanoate  $+ NH_3$  (overall reaction)

(1a) 1-aminocyclopropane-1-carboxylate = 2-aminobut-2-enoate(1b) 2-aminobut-2-enoate = 2-iminobutanoate (spontaneous)

(1c) 2-iminobutanoate +  $H_2O$  = 2-oxobutanoate +  $NH_3$  (spontaneous)

Other name(s): 1-aminocyclopropane-1-carboxylate endolyase (deaminating); ACC deaminase; 1-aminocyclopropane

carboxylic acid deaminase

**Systematic name:** 1-aminocyclopropane-1-carboxylate aminohydrolase (isomerizing)

**Comments:** A pyridoxal 5'-phosphate enzyme. The enzyme, found in certain soil bacteria and fungi, catalyses

the ring opening of 1-aminocyclopropane-1-carboxylate, the immediate precursor to ethylene, an important plant hormone that regulates fruit ripening and other processes. The enzyme releases an unstable enamine product that tautomerizes to an imine form, which undergoes a hydrolytic deamination. The latter reaction, which can occur spontaneously, can also be catalysed by EC 3.5.99.10, 2-iminobutanoate/2-iminopropanoate deaminase. The enzyme has been used to make fruit ripening dependent on externally added ethylene, as it removes the substrate for endogenous ethylene forma-

tion.

**References:** [1169, 3262, 2908]

[EC 3.5.99.7 created 1981 as EC 4.1.99.4, transferred 2002 to EC 3.5.99.7, modified 2014]

EC 3.5.99.8

**Accepted name:** 5-nitroanthranilic acid aminohydrolase

**Reaction:** 5-nitroanthranilate +  $H_2O = 5$ -nitrosalicylate +  $NH_3$ 

Other name(s): naaA (gene name); 5NAA deaminase Systematic name: 5-nitroanthranilate amidohydrolase

**Comments:** The enzyme catalyses the initial step in biodegradation of 5-nitroanthranilic acid by *Bradyrhizobium* 

sp. strain JS329.

**References:** [2338]

[EC 3.5.99.8 created 2011]

EC 3.5.99.9

Accepted name: 2-nitroimidazole nitrohydrolase

**Reaction:** 2-nitroimidazole +  $H_2O$  = imidazol-2-one + nitrite

Other name(s): NnhA; 2NI nitrohydrolase; 2NI denitrase

**Systematic name:** 2-nitroimidazole nitrohydrolase

**Comments:** The enzyme catalyses the initial step in the biodegradation of 2-nitroimidazole by the soil bacterium

Mycobacterium sp. JS330

**References:** [2339]

[EC 3.5.99.9 created 2012]

EC 3.5.99.10

Accepted name: 2-iminobutanoate/2-iminopropanoate deaminase Reaction: (1) 2-iminobutanoate +  $H_2O = 2$ -oxobutanoate +  $NH_3$ 

(2) 2-iminopropanoate +  $H_2O$  = pyruvate +  $NH_3$ 

Other name(s): yjgF (gene name); ridA (gene name); enamine/imine deaminase (ambiguous)

Systematic name: 2-iminobutanoate aminohydrolase

**Comments:** This enzyme, which has been found in all species and tissues examined, catalyses the hydrolytic

deamination of imine intermediates formed by several types of pyridoxal-5'-phosphate-dependent dehydratases, such as EC 4.3.1.19, threonine ammonia-lyase and EC 4.3.1.17, L-serine ammonia-lyase. The reactions, which can occur spontaneously, are accelerated to minimize the cellular damage that

could be caused by these reactive intermediates.

**References:** [1595]

[EC 3.5.99.10 created 2014]

EC 3.5.99.11

**Accepted name:** 2-aminomuconate deaminase (2-hydroxymuconate-forming)

**Reaction:** 2-aminomuconate +  $H_2O = (2Z,4E)$ -2-hydroxyhexa-2,4-dienedioate +  $NH_3$ 

**Other name(s):** *cnbZ* (gene name)

**Systematic name:** 2-aminomuconate aminohydrolase [(2Z,4E)-2-hydroxyhexa-2,4-dienedioate-forming]

**Comments:** The enzyme, characterized from the bacterium *Comamonas testosteroni* CNB-1, converts 2-

aminomuconate to 2-hydroxyhexa-2,4-dienedioate, unlike the enzymes from Pseudomonas, which produce (3E)-2-oxohex-3-enedioate (see EC 3.5.99.5, 2-aminomuconate deaminase). The enzyme

also acts on 2-amino-5-chloromuconate.

**References:** [1723]

[EC 3.5.99.11 created 2016 as EC 3.5.1.120, transferred 2017 to EC 3.5.99.11]

# EC 3.6 Acting on acid anhydrides

To this subclass belong mainly the enzymes acting on diphosphate bonds in compounds such as nucleoside di- and tri-phosphates (EC 3.6.1), on sulfonyl-containing anhydrides such as adenylylsulfate (EC 3.6.2) and on acid anhydrides; catalysing transmembrane movement of substances (EC 3.6.3).

## EC 3.6.1 In phosphorus-containing anhydrides

EC 3.6.1.1

Accepted name: inorganic diphosphatase

**Reaction:** diphosphate  $+ H_2O = 2$  phosphate **Systematic name:** diphosphate phosphohydrolase

**Comments:** Specificity varies with the source and with the activating metal ion. The enzyme from some sources

may be identical with EC 3.1.3.1 (alkaline phosphatase) or EC 3.1.3.9 (glucose-6-phosphatase). cf.

EC 7.1.3.1, H<sup>+</sup>-exporting diphosphatase.

**References:** [120, 1556, 2355]

#### EC 3.6.1.2

Accepted name: trimetaphosphatase

**Reaction:** trimetaphosphate  $+ H_2O = triphosphate$ 

Other name(s): inorganic trimetaphosphatase

Systematic name: trimetaphosphate hydrolase

**References:** [1513, 1898]

[EC 3.6.1.2 created 1961]

#### EC 3.6.1.3

Accepted name: adenosinetriphosphatase

**Reaction:** ATP +  $H_2O = ADP + phosphate$ 

Other name(s): adenylpyrophosphatase; ATP monophosphatase; triphosphatase; ATPase (ambiguous); SV40 T-

antigen; adenosine 5'-triphosphatase; ATP hydrolase; complex V (mitochondrial electron transport);

(Ca<sup>2+</sup> + Mg<sup>2+</sup>)-ATPase; HCO<sub>3</sub><sup>-</sup>-ATPase; adenosine triphosphatase

**Systematic name:** ATP phosphohydrolase

**Comments:** Many enzymes previously listed under this number are now listed separately under EC 3.6.3 and EC

3.6.4.

**References:** [888, 1437, 1821, 2101, 2427, 2925]

[EC 3.6.1.3 created 1961 (EC 3.6.1.4 created 1961, incorporated 1965)]

[3.6.1.4 Deleted entry. adenosinetriphosphatase (Mg-activated). Now included with EC 3.6.1.3 adenosinetriphosphatase]

[EC 3.6.1.4 created 1961, deleted 1965]

## EC 3.6.1.5

Accepted name: apyrase

**Reaction:** a nucleoside 5'-triphosphate +  $2 \text{ H}_2\text{O}$  = a nucleoside 5'-phosphate + 2 phosphate (overall reaction)

(1a) a nucleoside 5'-triphosphate +  $H_2O$  = a nucleoside 5'-diphosphate + phosphate (1b) a nucleoside 5'-diphosphate +  $H_2O$  = a nucleoside 5'-phosphate + phosphate

Other name(s): ATP-diphosphatase; adenosine diphosphatase; ADPase; ATP diphosphohydrolase [ambiguous]

**Systematic name:** nucleoside triphosphate phosphohydrolase (nucleoside monophosphoate-forming)

**Comments:** Apyrases are active against both di- and triphosphate nucleotides (NDPs and NTPs) and hydrolyse

NTPs to nucleotide monophosphates (NMPs) in two distinct successive phosphate-releasing steps, with NDPs as intermediates. They differ from ATPases, which specifically hydrolyse ATP, by hydrolysing both ATP and ADP. The eukaryotic enzymes requires Ca<sup>2+</sup>, but Mg<sup>2+</sup> can substitute. Most of the ecto-ATPases that occur on the cell surface and hydrolyse extracellular nucleotides belong to

this enzyme family.

**References:** [1536, 1681, 418, 444, 3119, 872, 3237]

[EC 3.6.1.5 created 1961, modified 1976, modified 2000, modified 2013]

#### EC 3.6.1.6

Accepted name: nucleoside diphosphate phosphatase

**Reaction:** a nucleoside diphosphate  $+ H_2O = a$  nucleoside phosphate + phosphate

Other name(s): nucleoside-diphosphatase; thiaminpyrophosphatase; UDPase; inosine diphosphatase; adenosine

diphosphatase; IDPase; ADPase; adenosinepyrophosphatase; guanosine diphosphatase; guanosine 5'-diphosphatase; inosine 5'-diphosphatase; uridine diphosphatase; uridine 5'-diphosphatase; type B nucleoside diphosphatase; GDPase; CDPase; nucleoside 5'-diphosphatase; type L nucleoside diphosphatase; t

phatase; NDPase; nucleoside diphosphate phosphohydrolase

**Systematic name:** nucleoside-diphosphate phosphohydrolase

**Comments:** The enzyme, which appears to be limited to metazoa, acts on multiple nucleoside diphosphates as

well as on D-ribose 5-diphosphate. Specificity depends on species and isoform.

**References:** [913, 1176, 3283, 731, 3005]

[EC 3.6.1.6 created 1961]

EC 3.6.1.7

Accepted name: acylphosphatase

**Reaction:** an acylphosphate  $+ H_2O = a$  carboxylate + phosphate

Other name(s): acetylphosphatase; 1,3-diphosphoglycerate phosphatase; acetic phosphatase; Ho 1-3; GP 1-3

**Systematic name:** acylphosphate phosphohydrolase

**References:** [2359, 2367, 2368, 2641]

[EC 3.6.1.7 created 1961]

EC 3.6.1.8

**Accepted name:** ATP diphosphatase

**Reaction:** ATP +  $H_2O = AMP + diphosphate$ 

Other name(s): ATPase (ambiguous); ATP pyrophosphatase; adenosine triphosphate pyrophosphatase; ATP diphos-

phohydrolase (ambiguous)

**Systematic name:** ATP diphosphohydrolase (diphosphate-forming)

**Comments:** Also acts on ITP, GTP, CTP and UTP.

**References:** [1123, 1329]

[EC 3.6.1.8 created 1961]

EC 3.6.1.9

**Accepted name:** nucleotide diphosphatase

**Reaction:** a nucleoside triphosphate  $+ H_2O = a$  nucleotide + diphosphate

Other name(s): ENPP1 (gene name); nucleotide pyrophosphatase; nucleotide-sugar pyrophosphatase; nucleoside-

triphosphate diphosphatase

**Systematic name:** nucleoside-triphosphate diphosphohydrolase

**Comments:** The enzyme preferentially hydrolyses ATP, but can also hydrolyse other nucleoside 5' triphosphates

such as GTP, CTP, TTP and UTP to their corresponding monophosphates. *In vitro* the enzyme also

acts as a nucleotidohydrolase on ADP, NAD+, NADP+, FAD, and CoA.

**References:** [424, 1418, 1597, 3325]

[EC 3.6.1.9 created 1961 (EC 3.6.1.19 created 1972, incorporated 2016), modified 2016]

EC 3.6.1.10

Accepted name: endopolyphosphatase

**Reaction:** polyphosphate + n H<sub>2</sub>O = (n+1) oligophosphate

**Other name(s):** polyphosphate depolymerase; metaphosphatase; polyphosphatase; polymetaphosphatase

**Systematic name:** polyphosphate polyphosphohydrolase

**Comments:** The product contains 4 or 5 phosphate residues.

**References:** [1793, 1844]

[EC 3.6.1.10 created 1961]

EC 3.6.1.11

Accepted name: exopolyphosphatase

**Reaction:**  $(polyphosphate)_n + H_2O = (polyphosphate)_{n-1} + phosphate$ 

**Other name(s):** metaphosphatase; acid phosphoanhydride phosphohydrolase; Gra-Pase

**Systematic name:** polyphosphate phosphohydrolase

**References:** [993, 1536, 1793]

[EC 3.6.1.11 created 1965]

EC 3.6.1.12

**Accepted name:** dCTP diphosphatase

**Reaction:**  $dCTP + H_2O = dCMP + diphosphate$ 

Other name(s): DCTPP1 (gene name); deoxycytidine-triphosphatase; dCTPase; dCTP pyrophosphatase; deoxycyti-

dine triphosphatase; deoxy-CTPase

**Systematic name:** dCTP nucleotidohydrolase

**Comments:** The mammalian enzyme also displays weak activity against dTTP and dATP, but none against dGTP.

Activity is highest with analogs including 5-iodo-dCTP and 5-methyl-dCTP.

**References:** [3343, 1982, 3217, 2109, 2410]

[EC 3.6.1.12 created 1965]

EC 3.6.1.13

**Accepted name:** ADP-ribose diphosphatase

**Reaction:** ADP-D-ribose +  $H_2O$  = AMP + D-ribose 5-phosphate

Other name(s): ADPribose pyrophosphatase; adenosine diphosphoribose pyrophosphatase; ADPR-PPase; ADP-ribose

ribophosphohydrolase

**Systematic name:** ADP-D-ribose ribophosphohydrolase

**References:** [630]

[EC 3.6.1.13 created 1965]

EC 3.6.1.14

**Accepted name:** adenosine-tetraphosphatase

**Reaction:** adenosine 5'-tetraphosphate +  $H_2O$  = ATP + phosphate

Systematic name: adenosine-tetraphosphate phosphohydrolase

**Comments:** Also acts on inosine tetraphosphate and tripolyphosphate but shows little or no activity with other

nucleotides or polyphosphates.

**References:** [2692]

[EC 3.6.1.14 created 1972]

EC 3.6.1.15

**Accepted name:** nucleoside-triphosphate phosphatase

**Reaction:** a nucleoside triphosphate +  $H_2O$  = a nucleoside diphosphate + phosphate

Other name(s): nucleoside-triphosphatase; nucleoside triphosphate phosphohydrolase; nucleoside-5-triphosphate

phosphohydrolase; nucleoside 5-triphosphatase; unspecific diphosphate phosphohydrolase

Systematic name: nucleoside-triphosphate phosphohydrolase

**Comments:** The enzyme is found in eukaryotes and thermophilic bacteria, but appears to be absent from

mesophilic bacteria. Also hydrolyses nucleoside diphosphates, thiamine diphosphate and FAD. The

enzyme from the plant *Pisum sativum* (garden pea) is regulated by calmodulin [1195].

**References:** [292, 1660, 1840, 2942, 1195, 1477, 2290]

[EC 3.6.1.15 created 1972]

EC 3.6.1.16

**Accepted name:** CDP-glycerol diphosphatase

**Reaction:** CDP-glycerol +  $H_2O = CMP + sn$ -glycerol 3-phosphate

**Other name(s):** CDP-glycerol pyrophosphatase; cytidine diphosphoglycerol pyrophosphatase

Systematic name: CDP-glycerol phosphoglycerohydrolase

**References:** [926]

[EC 3.6.1.16 created 1972]

#### EC 3.6.1.17

**Accepted name:** bis(5'-nucleosyl)-tetraphosphatase (asymmetrical)

**Reaction:**  $P^1$ ,  $P^4$ -bis(5'-guanosyl) tetraphosphate + H<sub>2</sub>O = GTP + GMP

Other name(s): bis(5'-guanosyl)-tetraphosphatase; bis(5'-adenosyl)-tetraphosphatase; diguanosinetetraphosphatase

(asymmetrical); dinucleosidetetraphosphatase (asymmetrical); diadenosine  $P^1$ ,  $P^4$ -tetraphosphatase;

dinucleoside tetraphosphatase; 1-P,4-P-bis(5'-nucleosyl)-tetraphosphate nucleotidohydrolase

**Systematic name:**  $P^1$ ,  $P^4$ -bis(5'-nucleosyl)-tetraphosphate nucleotidohydrolase

**Comments:** Also acts on bis(5'-xanthosyl)-tetraphosphate and, more slowly, on bis(5'-adenosyl)-tetraphosphate

and bis(5'-uridyl)-tetraphosphate [cf. EC 3.6.1.41 bis(5'-nucleosyl)-tetraphosphatase (symmetrical)]

**References:** [1302, 3029, 3128]

[EC 3.6.1.17 created 1972, modified 1976, modified 1986]

#### EC 3.6.1.18

**Accepted name:** FAD diphosphatase

**Reaction:**  $FAD + H_2O = AMP + FMN$ 

Other name(s): FAD pyrophosphatase; riboflavin adenine dinucleotide pyrophosphatase; flavin adenine dinucleotide

pyrophosphatase; riboflavine adenine dinucleotide pyrophosphatase; flavine adenine dinucleotide py-

rophosphatase

Systematic name: FAD nucleotidohydrolase

**Comments:** The plant enzyme also hydrolyses NAD<sup>+</sup> and NADH; the animal enzyme hydrolyses NAD<sup>+</sup> and CoA

at about half of the rate of hydrolysis of FAD. May be identical with EC 3.6.1.9 nucleotide diphos-

phatase.

**References:** [2379, 2636]

[EC 3.6.1.18 created 1972]

[3.6.1.19 Transferred entry. nucleoside-triphosphate diphosphatase. Now EC 3.6.1.9, nucleotide diphosphatase]

[EC 3.6.1.19 created 1972, deleted 2016]

## EC 3.6.1.20

**Accepted name:** 5'-acylphosphoadenosine hydrolase

**Reaction:** 5'-acylphosphoadenosine +  $H_2O$  = AMP + a carboxylate

**Other name(s):** 5-phosphoadenosine hydrolase

Systematic name: 5'-acylphosphoadenosine acylhydrolase
Comments: Also acts on inosine and uridine compounds.

**References:** [1414]

[EC 3.6.1.20 created 1972]

## EC 3.6.1.21

**Accepted name:** ADP-sugar diphosphatase

**Reaction:** ADP-sugar +  $H_2O$  = AMP +  $\alpha$ -D-aldose 1-phosphate

**Other name(s):** ADP-sugar pyrophosphatase; adenosine diphosphosugar pyrophosphatase

**Systematic name:** ADP-sugar sugarphosphohydrolase

**Comments:** Has a specificity that is distinct from that of UDP-sugar diphosphatase (EC 3.6.1.45).

**References:** [2451]

[EC 3.6.1.21 created 1972, modified 1999]

EC 3.6.1.22

**Accepted name:** NAD<sup>+</sup> diphosphatase

**Reaction:**  $NAD(H) + H_2O = AMP + NMN(H)$ 

Other name(s): NPY1 (gene name); nudC (gene name); NUDT7 (gene name); nicotinamide adenine dinucleotide

pyrophosphatase; NADP pyrophosphatase; NADH pyrophosphatase; NAD+ phosphohydrolase

**Systematic name:** NAD(H) phosphohydrolase

**Comments:** This enzyme, described from plants, animals, and bacteria, can act on both reduced and oxidized

forms of its substrate, although enzymes from different organisms have different preferences. Also acts on other dinucleotides, including NADP(H),  $FAD(H_2)$ , and the thionicotinamide analogues of

NAD<sup>+</sup> and NADP<sup>+</sup>.

**References:** [1512, 1291, 2821, 1551, 51, 2042, 815, 3236, 1304]

[EC 3.6.1.22 created 1972]

EC 3.6.1.23

**Accepted name:** dUTP diphosphatase

**Reaction:**  $dUTP + H_2O = dUMP + diphosphate$ 

Other name(s): DUT (gene name); deoxyuridine-triphosphatase; dUTPase; dUTP pyrophosphatase; desoxyuridine

5'-triphosphate nucleotidohydrolase; desoxyuridine 5'-triphosphatase

**Systematic name:** dUTP nucleotidohydrolase

**Comments:** The enzyme catalyses the Mg<sup>2+</sup>-dependent hydrolysis of dUTP to dUMP, providing the substrate for

EC 2.1.1.45, thymidylate synthase, leading to production of thymidine nucleotides. By reducing the effective ratio of dUTP to TTP, the enzyme also reduces the possibility of dUTP incorporation into

DNA.

**References:** [974, 205, 980, 2645, 924, 384, 1588, 123, 3052]

[EC 3.6.1.23 created 1972]

EC 3.6.1.24

Accepted name: nucleoside phosphoacylhydrolase

**Reaction:** Hydrolyses mixed phospho-anhydride bonds **Systematic name:** nucleoside-5'-phosphoacylate acylhydrolase

Comments: Attacks ribonucleoside 5'-nitrophenylphosphates, but is inactive against phosphodiesters.

**References:** [2736]

[EC 3.6.1.24 created 1972]

EC 3.6.1.25

Accepted name: triphosphatase

**Reaction:** triphosphate +  $H_2O$  = diphosphate + phosphate

Other name(s): inorganic triphosphatase
Systematic name: triphosphate phosphohydrolase

**References:** [1545, 3010]

[EC 3.6.1.25 created 1976]

EC 3.6.1.26

Accepted name: CDP-diacylglycerol diphosphatase

**Reaction:** CDP-diacylglycerol +  $H_2O$  = CMP + phosphatidate

Other name(s): cytidine diphosphodiacylglycerol pyrophosphatase; CDP diacylglycerol hydrolase

Systematic name: CDP-diacylglycerol phosphatidylhydrolase

**References:** [2354]

[EC 3.6.1.26 created 1976]

EC 3.6.1.27

Accepted name: undecaprenyl-diphosphate phosphatase

**Reaction:** ditrans, octacis-undecaprenyl diphosphate +  $H_2O = ditrans, octacis$ -undecaprenyl phosphate + phos-

phate

**Other name(s):** C<sub>55</sub>-isoprenyl diphosphatase; C<sub>55</sub>-isoprenyl pyrophosphatase; isoprenyl pyrophosphatase (ambigu-

ous); undecaprenyl pyrophosphate phosphatase; undecaprenyl pyrophosphate pyrophosphatase; UPP phosphatase; Und-PP pyrophosphatase; UppP (ambiguous); BacA; undecaprenyl-diphosphate phosphate phosph

phohydrolase; undecaprenyl-diphosphatase

**Systematic name:** ditrans,octacis-undecaprenyl-diphosphate phosphohydrolase

Comments: Isolated from the bacteria Micrococcus lysodeikticus [950], Escherichia coli [2,3,5,6] and Bacillus

subtilis [198]. The product of the reaction, ditrans, octacis-undecaprenyl phosphate, is essential for

cell wall polysaccharide biosynthesis in these strains.

**References:** [950, 900, 901, 198, 2892, 2952]

[EC 3.6.1.27 created 1978, modified 2002, modified 2012]

EC 3.6.1.28

Accepted name: thiamine-triphosphatase

**Reaction:** thiamine triphosphate +  $H_2O$  = thiamine diphosphate + phosphate

Systematic name: thiamine-triphosphate phosphohydrolase

**References:** [1071]

[EC 3.6.1.28 created 1978]

EC 3.6.1.29

**Accepted name:** bis(5'-adenosyl)-triphosphatase

**Reaction:**  $P^1$ ,  $P^3$ -bis(5'-adenosyl) triphosphate + H<sub>2</sub>O = ADP + AMP

**Other name(s):** dinucleosidetriphosphatase; diadenosine  $5.5-P^1$ ,  $P^3$ -triphosphatase; 1-P.3-P-bis(5'-adenosyl)-

triphosphate adenylohydrolase

**Systematic name:**  $P^1$ ,  $P^3$ -bis(5'-adenosyl)-triphosphate adenylohydrolase

**References:** [1302, 2659]

[EC 3.6.1.29 created 1978]

[3.6.1.30 Deleted entry.  $m^7G(5')pppN$  diphosphatase. Now covered by EC 3.6.1.59 [ $m^7GpppX$  diphosphatase] and EC 3.6.1.62 [ $m^7GpppN$ -mRNA hydrolase].]

[EC 3.6.1.30 created 1978, deleted 2012]

EC 3.6.1.31

**Accepted name:** phosphoribosyl-ATP diphosphatase

**Reaction:** 1-(5-phospho- $\beta$ -D-ribosyl)-ATP + H<sub>2</sub>O = 1-(5-phospho- $\beta$ -D-ribosyl)-AMP + diphosphate

Other name(s): phosphoribosyl-ATP pyrophosphatase; phosphoribosyladenosine triphosphate pyrophosphatase; 1-(5-

phosphoribosyl)-ATP diphosphohydrolase

**Systematic name:** 1-(5-phospho-β-D-ribosyl)-ATP diphosphohydrolase

**Comments:** The *Neurospora crassa* enzyme also catalyses the reactions of EC 1.1.1.23 (histidinol dehydrogenase)

and EC 3.5.4.19 (phosphoribosyl-AMP cyclohydrolase).

**References:** [2697]

[EC 3.6.1.31 created 1981]

[3.6.1.32 Transferred entry. myosin ATPase. Now EC 3.6.4.1, myosin ATPase]

[EC 3.6.1.32 created 1984, deleted 2000]

[3.6.1.33 Transferred entry. dynein ATPase. Now EC 3.6.4.2, dynein ATPase]

[EC 3.6.1.33 created 1984, deleted 2000]

[3.6.1.34 Transferred entry.  $H^+$ -transporting ATP synthase. Now EC 3.6.3.14,  $H^+$ -transporting two-sector ATPase]

[EC 3.6.1.34 created 1984, deleted 2000]

[3.6.1.35 Transferred entry. H<sup>+</sup>-transporting ATPase. Now EC 3.6.3.6, H<sup>+</sup>-exporting ATPase]

[EC 3.6.1.35 created 1984, deleted 2000]

[3.6.1.36 Transferred entry.  $H^+/K^+$  exchanging ATPase. Now EC 3.6.3.10,  $H^+/K^+$ -exchanging ATPase]

[EC 3.6.1.36 created 1984, deleted 2000]

[3.6.1.37 Transferred entry.  $Na^+/K^+$  exchanging ATPase. Now EC 3.6.3.9,  $Na^+/K^+$ -exchanging ATPase]

[EC 3.6.1.37 created 1984, deleted 2000]

[3.6.1.38 Transferred entry. Ca<sup>2+</sup>-transporting ATPase. Now EC 3.6.3.8, Ca<sup>2+</sup>-transporting ATPase]

[EC 3.6.1.38 created 1984, deleted 2000]

## EC 3.6.1.39

Accepted name: thymidine-triphosphatase

**Reaction:**  $dTTP + H_2O = dTDP + phosphate$ 

Other name(s): thymidine triphosphate nucleotidohydrolase; dTTPase; deoxythymidine-5'-triphosphatase

**Systematic name:** dTTP nucleotidohydrolase

**Comments:** Also acts, more slowly, on dUTP and UTP.

**References:** [528]

[EC 3.6.1.39 created 1984]

## EC 3.6.1.40

**Accepted name:** guanosine-5'-triphosphate,3'-diphosphate phosphatase

guanosine pentaphosphatase; guanosine pentaphosphate phosphatase; guanosine 5'-triphosphate 3'-

diphosphate 5'-phosphatase; guanosine pentaphosphate phosphohydrolase

**Systematic name:** guanosine-5'-triphosphate-3'-diphosphate 5'-phosphohydrolase

**Comments:** Also hydrolyses other guanosine 5'-triphosphate derivatives with at least one unsubstituted phosphate

group on the 3'-position, but not GTP, ATP or adenosine 5'-triphosphate 3'-diphosphate.

**References:** [1047]

[EC 3.6.1.40 created 1986, modified 2010]

## EC 3.6.1.41

**Accepted name:** bis(5'-nucleosyl)-tetraphosphatase (symmetrical)

**Reaction:**  $P^1$ ,  $P^4$ -bis(5'-adenosyl) tetraphosphate + H<sub>2</sub>O = **2** ADP

Other name(s): diadenosinetetraphosphatase (symmetrical); dinucleosidetetraphosphatase (symmetrical); symmetri-

cal diadenosine tetraphosphate hydrolase; adenosine tetraphosphate phosphodiesterase; Ap4A hydrolase; bis(5'-adenosyl) tetraphosphatase; diadenosine tetraphosphate hydrolase; diadenosine polyphosphate hydrolase; diadenosine 5',5'''- $P^1$ , $P^4$ -tetraphosphatase; diadenosinetetraphosphatase (symmetri-

cal); 1-P,4-P-bis(5'-nucleosyl)-tetraphosphate nucleosidebisphosphohydrolase

**Systematic name:**  $P^1$ ,  $P^4$ -bis(5'-nucleosyl)-tetraphosphate nucleosidebisphosphohydrolase

**Comments:** Also acts on bis(5'-guanosyl) tetraphosphate and bis(5'-adenosyl) pentaphosphate and, more slowly,

on some other polyphosphates, forming a nucleoside bisphosphate as one product in all cases [cf. EC

3.6.1.17 bis(5'-nucleosyl)-tetraphosphatase (asymmetrical)].

**References:** [148, 1002]

[EC 3.6.1.41 created 1986]

EC 3.6.1.42

Accepted name: guanosine-diphosphatase

**Reaction:**  $GDP + H_2O = GMP + phosphate$ 

Other name(s): GDPase

**Systematic name:** GDP phosphohydrolase

**Comments:** Also acts on UDP but not on other nucleoside diphosphates and triphosphates.

**References:** [2388]

[EC 3.6.1.42 created 1989]

EC 3.6.1.43

Accepted name: dolichyldiphosphatase

**Reaction:** dolichyl diphosphate +  $H_2O$  = dolichyl phosphate + phosphate

Other name(s): dolichol diphosphatase; dolichyl pyrophosphatase; dolichyl pyrophosphatase; dolichyl

diphosphate phosphohydrolase; Dol-P-P phosphohydrolase

Systematic name: dolichyl-diphosphate phosphohydrolase

**References:** [2058]

[EC 3.6.1.43 created 1989]

EC 3.6.1.44

Accepted name: oligosaccharide-diphosphodolichol diphosphatase

**Reaction:** oligosaccharide-diphosphodolichol + H<sub>2</sub>O = oligosaccharide phosphate + dolichyl phosphate

**Other name(s):** oligosaccharide-diphosphodolichol pyrophosphatase

Systematic name: oligosaccharide-diphosphodolichol phosphodolichohydrolase

**References:** [186]

[EC 3.6.1.44 created 1992]

EC 3.6.1.45

Accepted name: UDP-sugar diphosphatase

**Reaction:** UDP-sugar +  $H_2O$  = UMP +  $\alpha$ -D-aldose 1-phosphate

Other name(s): nucleosidediphosphate-sugar pyrophosphatase; nucleosidediphosphate-sugar diphosphatase; UDP-

sugar hydrolase; UDP-sugar pyrophosphatase

**Systematic name:** UDP-sugar sugarphosphohydrolase

**Comments:** A divalent cation is required for activity. UDP-sugar is the best substrate, although other nucleoside-

sugar diphosphates are used as substrates with similar  $K_m$  values but much lower maximum velocities. Thus, this enzyme has a specificity distinct from that of ADP-sugar diphosphatase (EC 3.6.1.21). Some but not all enzymes of this class also appear to have 5'-nucleotidase (see EC 3.1.3.5) activity.

**References:** [881, 927]

## [EC 3.6.1.45 created 1999]

[3.6.1.46	Transferred entry. heterotrimeric G-protein GTPase. Now EC 3.6.5.1, heterotrimeric G-protein GTPase]
	[EC 3.6.1.46 created 2000, deleted 2003]
[3.6.1.47	Transferred entry. small monomeric GTPase. Now EC 3.6.5.2, small monomeric GTPase]
	[EC 3.6.1.47 created 2000, deleted 2003]
[3.6.1.48	Transferred entry. protein-synthesizing GTPase. Now EC 3.6.5.3, protein-synthesizing GTPase]
	[EC 3.6.1.48 created 2000, deleted 200]
[3.6.1.49	Transferred entry. signal-recognition-particle GTPase. Now EC 3.6.5.4, signal-recognition-particle GTPase]
	[EC 3.6.1.49 created 2000, deleted 2003]
[3.6.1.50	Transferred entry. dynamin GTPase. Now EC 3.6.5.5, dynamin GTPase]
	[EC 3.6.1.50 created 2000, deleted 2003]
[3.6.1.51	Transferred entry. tubulin GTPase. Now EC 3.6.5.6, tubulin GTPase]
	[EC 3.6.1.51 created 2000, deleted 2003]

#### EC 3.6.1.52

**Accepted name:** diphosphoinositol-polyphosphate diphosphatase

**Reaction:** diphospho-myo-inositol polyphosphate +  $H_2O = myo$ -inositol polyphosphate + phosphate

**Other name(s):** diphosphoinositol-polyphosphate phosphohydrolase; DIPP **Systematic name:** diphospho-*myo*-inositol-polyphosphate diphosphohydrolase

**Comments:** This enzyme hydrolyses the diphosphate bond, leaving a phospho group where a diphospho group had

been. It can also act on bis(adenosine) diphosphate.

**References:** [2491, 352]

[EC 3.6.1.52 created 2002]

## EC 3.6.1.53

**Accepted name:** Mn<sup>2+</sup>-dependent ADP-ribose/CDP-alcohol diphosphatase

**Reaction:** (1) CDP-choline +  $H_2O = CMP + phosphocholine$ 

(2) ADP-D-ribose +  $H_2O$  = AMP + D-ribose 5-phosphate

**Other name(s):** Mn<sup>2+</sup>-dependent ADP-ribose/CDP-alcohol pyrophosphatase; ADPRibase-Mn

**Systematic name:** CDP-choline phosphohydrolase

**Comments:** Requires Mn<sup>2+</sup>. Unlike EC 3.6.1.13, ADP-ribose diphosphatase, it cannot utilize Mg<sup>2+</sup>. ADP-D-

ribose, CDP-choline, CDP-ethanolamine and ADP are substrates for this enzyme but ADP-D-glucose, UDP-D-glucose, CDP-D-glucose, CDP, CMP and AMP are not hydrolysed [362]. The mammalian enzyme hydrolyses cyclic ADP-ribose to 1-(5-phospho-β-D-ribosyl)-AMP with 100-fold lower efficiency than ADP-D-ribose [363]. In rat, the enzyme is found predominantly in thymus and spleen.

**References:** [364, 362, 363, 2448]

[EC 3.6.1.53 created 2008]

## EC 3.6.1.54

**Accepted name:** UDP-2,3-diacylglucosamine diphosphatase

**Reaction:** UDP-2-N,3-O-bis[(3R)-3-hydroxytetradecanoyl]- $\alpha$ -D-glucosamine + H<sub>2</sub>O = 2-N,3-O-bis[(3R)-3-

hydroxytetradecanoyl ]-α-D-glucosaminyl 1-phosphate + UMP

**Other name(s):** UDP-2,3-diacylglucosamine hydrolase; UDP-2,3-diacylglucosamine pyrophosphatase; *ybbF* (gene

name); lpxH (gene name); UDP-2,3-bis[(3R)-3-hydroxymyristoyl $]-\alpha$ -D-glucosamine 2,3-bis[(3R)-3-

hydroxymyristoyl]-β-D-glucosaminyl 1-phosphate phosphohydrolase (incorrect)

Systematic name: UDP-2-N,3-O-bis[(3R)-3-hydroxytetradecanoyl]- $\alpha$ -D-glucosamine 2-N,3-O-bis[(3R)-3-

hydroxytetradecanoyl]-α-D-glucosaminyl 1-phosphate phosphohydrolase

**Comments:** The enzyme catalyses a step in the biosynthesis of lipid A.

**References:** [111, 110]

[EC 3.6.1.54 created 2010]

#### EC 3.6.1.55

**Accepted name:** 8-oxo-dGTP diphosphatase

**Reaction:** 8-oxo-dGTP +  $H_2O = 8$ -oxo-dGMP + diphosphate

Other name(s): MutT; 7,8-dihydro-8-oxoguanine triphosphatase; 8-oxo-dGTPase; 7,8-dihydro-8-oxo-dGTP py-

rophosphohydrolase

**Systematic name:** 8-oxo-dGTP diphosphohydrolase

Comments: This enzyme hydrolyses the phosphoanhydride bond between the  $\alpha$  and  $\beta$  phosphate of 8-oxoguanine-

containing nucleoside di- and triphosphates thereby preventing misincorporation of the oxidized purine nucleoside triphosphates into DNA. It does not hydrolyse 2-hydroxy-dATP (*cf.* EC 3.6.1.56,

2-hydroxy-dATP diphosphatase) [3289]. Requires Mg<sup>2+</sup>.

**References:** [1267, 3305, 2047, 3289]

[EC 3.6.1.55 created 2011]

#### EC 3.6.1.56

**Accepted name:** 2-hydroxy-dATP diphosphatase

**Reaction:** 2-hydroxy-dATP +  $H_2O = 2$ -hydroxy-dAMP + diphosphate

Other name(s): NUDT1; MTH1; MTH<sub>2</sub>; oxidized purine nucleoside triphosphatase; (2'-deoxy) ribonucleoside 5'-

triphosphate pyrophosphohydrolase

**Systematic name:** 2-hydroxy-dATP diphosphohydrolase

**Comments:** The enzyme hydrolyses oxidized purine nucleoside triphosphates such as 2-hydroxy-dATP, thereby

preventing their misincorporation into DNA. It can also recognize 8-oxo-dGTP and 8-oxo-dATP, but

with lower efficiency (cf. EC 3.6.1.55, 8-oxo-dGTP diphosphatase) [838].

**References:** [2506, 1360, 838, 2503, 839]

[EC 3.6.1.56 created 2011]

## EC 3.6.1.57

**Accepted name:** UDP-2,4-diacetamido-2,4,6-trideoxy-β-L-altropyranose hydrolase

**Reaction:** UDP-2,4-diacetamido-2,4,6-trideoxy- $\beta$ -L-altropyranose + H<sub>2</sub>O = 2,4-diacetamido-2,4,6-trideoxy- $\beta$ -L-

altropyranose + UDP

Other name(s): PseG; UDP-6-deoxy-AltdiNAc hydrolase; Cj1312; UDP-2,4-bis(acetamido)-2,4,6-trideoxy-β-L-

altropyranose hydrolase

**Systematic name:** UDP-2,4-diacetamido-2,4,6-trideoxy-β-L-altropyranose hydrolase

**Comments:** The enzyme is involved in biosynthesis of pseudaminic acid.

**References:** [1719, 2581]

[EC 3.6.1.57 created 2011]

#### EC 3.6.1.58

Accepted name: 8-oxo-dGDP phosphatase

**Reaction:** 8-oxo-dGDP +  $H_2O = 8$ -oxo-dGMP + phosphate

Other name(s): NUDT5; MTH3 (gene name); NUDT18

**Systematic name:** 8-oxo-dGDP phosphohydrolase

**Comments:** The enzyme catalyses the hydrolysis of both 8-oxo-dGDP and 8-oxo-GDP thereby preventing trans-

lational errors caused by oxidative damage. The preferred *in vivo* substrate is not known. The enzyme does not degrade 8-oxo-dGTP and 8-oxo-GTP to the monophosphates (*cf.* EC 3.6.1.55, 8-oxo-dGTP diphosphatase) [1259, 1260]. Ribonucleotide diphosphates and deoxyribonucleotide diphosphates are hydrolysed with broad specificity. The bifunctional enzyme NUDT5 also hydrolyses ADP-ribose to AMP and D-ribose 5-phosphate (*cf.* EC 3.6.1.13, ADP-ribose diphosphatase) [1268]. The human enzyme NUDT18 also hydrolyses 8-oxo-dADP and 2-hydroxy-dADP, the latter at a slower rate [2835].

**References:** [1259, 1260, 1373, 1268, 3323, 2835]

[EC 3.6.1.58 created 2012]

EC 3.6.1.59

**Accepted name:** 5'- $(N^7$ -methyl 5'-triphosphoguanosine)-[mRNA] diphosphatase

**Reaction:** a 5'-( $N^7$ -methyl 5'-triphosphoguanosine)-[mRNA] +  $H_2O = N^7$ -methylguanosine 5'-phosphate + a

5'-diphospho-[mRNA]

**Other name(s):** DcpS; m<sup>7</sup>GpppX pyrophosphatase; m<sup>7</sup>GpppN m<sup>7</sup>GMP phosphohydrolase; m<sup>7</sup>GpppX diphosphatase;

m<sup>7</sup>G5'ppp5'N m<sup>7</sup>GMP phosphohydrolase

**Systematic name:** 5'- $(N^7$ -methyl 5'-triphosphoguanosine)-[mRNA]  $N^7$ -methylguanosine 5'-phosphate phosphohydro-

lase

Comments: The enzyme removes (decaps) the  $N^7$ -methylguanosine 5-phosphate cap from an mRNA degraded

to a maximal length of 10 nucleotides [1720, 464]. Decapping is an important process in the control of eukaryotic mRNA degradation. The enzyme functions to clear the cell of cap structure following decay of the RNA body [1725]. The nematode enzyme can also decap triply methylated substrates,

 $5'-(N^2,N^2,N^7-\text{trimethyl }5'-\text{triphosphoguanosine})-[\text{mRNA}]$  [3045].

**References:** [1795, 1725, 1720, 3045, 414, 464, 3227]

[EC 3.6.1.59 created 2012, modified 2013]

EC 3.6.1.60

**Accepted name:** diadenosine hexaphosphate hydrolase (AMP-forming)

**Reaction:** (1)  $P^1$ ,  $P^6$ -bis(5'-adenosyl)hexaphosphate + H<sub>2</sub>O = adenosine 5'-pentaphosphate + AMP

(2)  $P^1$ ,  $P^5$ -bis(5'-adenosyl)pentaphosphate + H<sub>2</sub>O = adenosine 5'-tetraphosphate + AMP

Other name(s): hAps1; NUDT11 (gene name); hAps2; NUDT10 (gene name)

**Systematic name:**  $P^1$ , $P^6$ -bis(5'-adenosyl)hexaphosphate nucleotidohydrolase (AMP-forming)

Comments: A divalent cation is essential for activity. Mn<sup>2+</sup> (2–6 mM) is most effective. The enzyme controls in-

tracellular levels of  $P^1$ ,  $P^5$ -bis(5'-adenosyl)pentaphosphate and P1,P6-bis(5'-adenosyl)hexaphosphate. Weak activity with P1,P4-bis(5'-adenosyl)tetraphosphate. Marked preference for adenine over gua-

nine nucleotides.

**References:** [1651, 2492]

[EC 3.6.1.60 created 2012]

EC 3.6.1.61

**Accepted name:** diadenosine hexaphosphate hydrolase (ATP-forming)

**Reaction:** (1)  $P^1$ ,  $P^6$ -bis(5'-adenosyl)hexaphosphate + H<sub>2</sub>O = **2** ATP

(2)  $P^1$ ,  $P^5$ -bis(5'-adenosyl)pentaphosphate + H<sub>2</sub>O = ATP + ADP

(3)  $P^1$ ,  $P^4$ -bis(5'-adenosyl)tetraphosphate + H<sub>2</sub>O = ATP + AMP

Other name(s): Ndx1

**Systematic name:**  $P^1, P^6$ -bis(5'-adenosyl)hexaphosphate nucleotidohydrolase (ATP-forming)

Comments: The enzyme requires the presence of the divalent cations  $(Mn^{2+}, Mg^{2+}, Zn^{2+}, and Co^{2+})$ . It hy-

drolyses  $P^1$ ,  $P^4$ -bis(5-guanosyl) tetraphosphate very slowly [cf. EC 3.6.1.17, bis(5-nucleosyl)-

tetraphosphatase (asymmetrical)].

**References:** [1281]

## [EC 3.6.1.61 created 2012]

EC 3.6.1.62

**Accepted name:** 5'- $(N^7$ -methylguanosine 5'-triphospho)-[mRNA] hydrolase

**Reaction:** a 5'- $(N^7$ -methylguanosine 5'-triphospho)-[mRNA] + H<sub>2</sub>O =  $N^7$ -methylguanosine 5'-diphosphate + a

5'-phospho-[mRNA]

Other name(s): Dcp2; NUDT16; D10 protein; D9 protein; D10 decapping enzyme; decapping enzyme; m<sup>7</sup>GpppN-

mRNA hydrolase; m<sup>7</sup>GpppN-mRNA m<sup>7</sup>GDP phosphohydrolase

**Systematic name:** 5'- $(N^7$ -methylguanosine 5'-triphospho)-[mRNA]  $N^7$ -methylguanosine-5'-diphosphate phosphohydro-

lase

Comments: Decapping of mRNA is a critical step in eukaryotic mRNA turnover. The enzyme is unable to cleave

a free cap structure (m<sup>7</sup>GpppG) [3044]. The enzyme from Vaccinia virus is synergistically activated

in the presence of  $Mg^{2+}$  and  $Mn^{2+}$  [2732].

**References:** [3233, 1744, 3044, 2239, 2732, 2238, 2720]

[EC 3.6.1.62 created 2012, modified 2013]

EC 3.6.1.63

**Accepted name:** α-D-ribose 1-methylphosphonate 5-triphosphate diphosphatase

**Reaction:**  $\alpha$ -D-ribose 1-methylphosphonate 5-triphosphate + H<sub>2</sub>O =  $\alpha$ -D-ribose 1-methylphosphonate 5-

phosphate + diphosphate

**Other name(s):** *phnM* (gene name)

**Systematic name:** α-D-ribose-1-methylphosphonate-5-triphosphate diphosphohydrolase

**Comments:** Isolated from the bacterium *Escherichia coli*.

**References:** [1370]

[EC 3.6.1.63 created 2012]

EC 3.6.1.64

**Accepted name:** inosine diphosphate phosphatase

**Reaction:** (1) IDP +  $H_2O$  = IMP + phosphate

(2)  $dIDP + H_2O = dIMP + phosphate$ 

Other name(s): (deoxy)inosine diphosphatase; NUDT16

Systematic name: inosine diphosphate phosphatase

**Comments:** The human enzyme also hydrolyses GDP and dGDP, and to a lesser extent ITP, dITP and XTP.

**References:** [1284]

[EC 3.6.1.64 created 2013]

EC 3.6.1.65

Accepted name: (d)CTP diphosphatase

**Reaction:** (1) CTP +  $H_2O = CMP + diphosphate$ 

(2)  $dCTP + H_2O = dCMP + diphosphate$ 

**Other name(s):** (d)CTP pyrophosphohydrolase; (d)CTP diphosphohydrolase; *nudG* (gene name)

**Systematic name:** (deoxy)cytidine 5'-triphosphate diphosphohydrolase

**Comments:** The enzyme, characterized from the bacterium *Escherichia coli*, is specific for the pyrimidine nu-

cleotides CTP and dCTP. It also acts on 5-methyl-dCTP, 5-hydroxy-dCTP and 8-hydroxy-dGTP.

**References:** [2155, 840, 1374, 1224]

[EC 3.6.1.65 created 2013]

EC 3.6.1.66

Accepted name: XTP/dITP diphosphatase

**Reaction:** (1) XTP +  $H_2O = XMP + diphosphate$ 

(2)  $dITP + H_2O = dIMP + diphosphate$ (2)  $ITP + H_2O = IMP + diphosphate$ 

**Other name(s):** hypoxanthine/xanthine dNTP pyrophosphatase; rdgB (gene name)

**Systematic name:** XTP/dITP diphosphohydrolase (diphosphate-forming)

Comments: The enzymes from the bacterium Escherichia coli and the archaea Methanococcus jannaschii and Ar-

chaeoglobus fulgidus are highly specific for XTP, dITP and ITP. The activity is dependent on divalent

cations, Mg<sup>2+</sup> is preferred.

**References:** [1213, 448, 449, 2537]

[EC 3.6.1.66 created 2013]

EC 3.6.1.67

**Accepted name:** dihydroneopterin triphosphate diphosphatase

**Reaction:** 7,8-dihydroneopterin 3'-triphosphate +  $H_2O = 7$ ,8-dihydroneopterin 3'-phosphate + diphosphate **Other name(s):** folQ (gene name); nudB (gene name); NUDT1 (gene name); dihydroneopterin triphosphate pyrophos-

phohydrolase

**Systematic name:** 7,8-dihydroneopterin 3'-triphosphate diphosphohydrolase

Comments: The enzyme participates in a folate biosynthesis pathway, which is found in bacteria, fungi, and

plants. Requires Mg<sup>2+</sup>.

**References:** [2816, 2156, 1472, 867]

[EC 3.6.1.67 created 2014]

EC 3.6.1.68

Accepted name: geranyl diphosphate phosphohydrolase

**Reaction:** geranyl diphosphate +  $H_2O$  = geranyl phosphate + phosphate

Other name(s): NUDX1 (gene name)

**Systematic name:** geranyl-diphosphate phosphohydrolase

**Comments:** The enzyme, characterized from roses, is involved in a cytosolic pathway for the biosynthesis of free

monoterpene alcohols that contribute to fragrance. In vitro the enzyme also acts on (2E,6E)-farnesyl

diphosphate.

**References:** [1771]

[EC 3.6.1.68 created 2015 as EC 3.1.3.98, transferred 2016 to EC 3.6.1.68]

## EC 3.6.2 In sulfonyl-containing anhydrides

EC 3.6.2.1

Accepted name: adenylylsulfatase

**Reaction:** adenylyl sulfate +  $H_2O = AMP + sulfate$ 

**Other name(s):** adenosine 5-phosphosulfate sulfohydrolase; adenylylsulfate sulfohydrolase

**Systematic name:** adenylyl-sulfate sulfohydrolase

**References:** [122]

[EC 3.6.2.1 created 1972]

EC 3.6.2.2

**Accepted name:** phosphoadenylylsulfatase

**Reaction:** 3'-phosphoadenylyl sulfate + H<sub>2</sub>O = adenosine 3'.5'-bisphosphate + sulfate

Other name(s): 3-phosphoadenylyl sulfatase; 3-phosphoadenosine 5-phosphosulfate sulfatase; PAPS sulfatase; 3'-

phosphoadenylylsulfate sulfohydrolase

**Systematic name:** 3'-phosphoadenylyl-sulfate sulfohydrolase

**Comments:** Requires  $Mn^{2+}$ .

**References:** [128]

[EC 3.6.2.2 created 1972]

## EC 3.6.3 Acting on acid anhydrides to catalyse transmembrane movement of substances

Several types of ATP phosphohydrolase are listed here. Entries EC 3.6.3.1 to EC 3.6.3.12 and EC 3.6.3.53 are enzymes undergoing covalent phosphorylation of an aspartate residue during the transport cycle; entries EC 3.6.3.14 and EC 3.6.3.15 refer to enzymes of complicated membrane and non-membrane location that can also serve in ATP synthesis; entry EC 3.6.3.16 is a multisubunit enzyme that is involved in arsenite transport only; entries EC 3.6.3.17 to EC 3.6.3.50 are two-domain enzymes of the ABC family; entries EC 3.6.3.51 and EC 3.6.3.52 are parts of a complex protein-transporting machinery in mitochondria and chloroplasts.

[3.6.3.1 Transferred entry. phospholipid-translocating ATPase. Now EC 7.6.2.1, P-type phospholipid transporter]

[EC 3.6.3.1 created 2000 (EC 3.6.3.13 created 2000, incorporated 2001), deleted 2018]

[3.6.3.2 Transferred entry.  $Mg^{2+}$ -importing ATPase. Now EC 7.2.2.14, P-type  $Mg^{2+}$  transporter]

[EC 3.6.3.2 created 2000, modified 2001, deleted 2018]

## EC 3.6.3.3

**Accepted name:** Cd<sup>2+</sup>-exporting ATPase

**Reaction:** ATP +  $H_2O + Cd^{2+}_{in} = ADP + phosphate + <math>Cd^{2+}_{out}$ 

**Systematic name:** ATP phosphohydrolase ( $Cd^{2+}$ -exporting)

**Comments:** A P-type ATPase that undergoes covalent phosphorylation during the transport cycle. This enzyme

occurs in protozoa, fungi and plants.

**References:** [2660, 2968]

[EC 3.6.3.3 created 2000]

[3.6.3.4	Transferred entry.	$Cu^{2+}$ -exporting ATPase.	Now EC 7.2.2.9,	Cu <sup>2+</sup> -exporting ATPase]
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[EC 3.6.3.4 created 2000, modified 2013, deleted 2018]

[3.6.3.5 Transferred entry. 
$$Zn^{2+}$$
-exporting ATPase. Now EC 7.2.2.12,  $Zn^{2+}$ -exporting ATPase]

[EC 3.6.3.5 created 2000, modified 2001, modified 2006, deleted 2018]

[3.6.3.6 Transferred entry.  $H^+$ -exporting ATPase. Now EC 7.1.2.1, P-type  $H^+$ -exporting transporter]

 $[EC\ 3.6.3.6\ created\ 1984\ as\ EC\ 3.6.1.35,\ transferred\ 2000\ to\ EC\ 3.6.3.6,\ deleted\ 2018]$ 

[3.6.3.7 Transferred entry. Na<sup>+</sup>-exporting ATPase. Now EC 7.2.2.3, P-type Na<sup>+</sup> transporter]

 $[EC\ 3.6.3.7\ created\ 2000,\ modified\ 2001,\ transferred\ 2018\ to\ EC\ 7.2.2.3,\ deleted\ 2018]$ 

[3.6.3.8 Transferred entry. Ca<sup>2+</sup>-transporting ATPase. Now EC 7.2.2.10, Ca<sup>2+</sup>-transporting ATPase]

[EC 3.6.3.8 created 1984 as EC 3.6.1.38, transferred 2000 to EC 3.6.3.8, modified 2001, modified 2011, deleted 2018]

[3.6.3.9 Transferred entry, Na<sup>+</sup>/K<sup>+</sup>-exchanging ATPase. Now EC 7.2.2.13, Na<sup>+</sup>/K<sup>+</sup>-exchanging ATPase.]

[EC 3.6.3.9 created 1984 as EC 3.6.1.37, transferred 2000 to EC 3.6.3.9, modified 2001, deleted 2018]

[3.6.3.10 Transferred entry.  $H^+/K^+$ -exchanging ATPase. Now EC 7.2.2.19,  $H^+/K^+$ -exchanging ATPase]

[EC 3.6.3.10 created 1984 as EC 3.6.1.36, transferred 2000 to EC 3.6.3.10, deleted 2018]

## EC 3.6.3.11

**Accepted name:** Cl<sup>-</sup>-transporting ATPase

**Reaction:** ATP +  $H_2O$  +  $Cl_{out}$  = ADP + phosphate +  $Cl_{in}$ **Other name(s):** Cl<sup>-</sup>-translocating ATPase; Cl<sup>-</sup>-motive ATPase

**Systematic name:** ATP phosphohydrolase (Cl<sup>-</sup>-importing)

**Comments:** A P-type ATPase that undergoes covalent phosphorylation during the transport cycle. An animal and

plant enzyme involved in the import of chloride anions.

**References:** [2158, 893, 1245]

[EC 3.6.3.11 created 2000]

[3.6.3.12 Transferred entry. K<sup>+</sup>-transporting ATPase. Now EC 7.2.2.6, K<sup>+</sup>-transporting ATPase]

[EC 3.6.3.12 created 2000, deleted 2018]

[3.6.3.13 Deleted entry. aminophospholipid-transporting ATPase. Identical to EC 3.6.3.1, phospholipid-translocating AT-Pase]

[EC 3.6.3.13 created 2000, deleted 2001]

[3.6.3.14 Transferred entry.  $H^+$ -transporting two-sector ATPase. Now EC 7.1.2.2,  $H^+$ -transporting two-sector ATPase]

[EC 3.6.3.14 created 1984 as EC 3.6.1.34, transferred 2000 to EC 3.6.3.14, deleted 2018]

[3.6.3.15 Transferred entry. Na<sup>+</sup>-transporting two-sector ATPase. Now EC 7.2.2.1, Na<sup>+</sup>-transporting two-sector ATPase]

[EC 3.6.3.15 created 2000, transferred 2018 to EC 7.2.2.1, deleted 2018]

## EC 3.6.3.16

Accepted name: arsenite-transporting ATPase

**Reaction:** ATP +  $H_2O$  + arsenite<sub>in</sub> = ADP + phosphate + arsenite<sub>out</sub>

**Systematic name:** ATP phosphohydrolase (arsenite-exporting)

Comments: A multisubunit non-phosphorylated ATPase that is involved in the transport of ions. A bacterial en-

zyme that usually contains two subunits where one (with 12 membrane-spanning segments) forms the 'channel' part and the other (occurring in pairs peripherally to the membrane) contains the ATP-

binding site. Exports arsenite and antimonite anions.

**References:** [2661, 2460, 316, 3338]

[EC 3.6.3.16 created 2000]

## EC 3.6.3.17

Accepted name: monosaccharide-transporting ATPase

**Reaction:** ATP +  $H_2O$  + monosaccharide<sub>out</sub> = ADP + phosphate + monosaccharide<sub>in</sub>

**Systematic name:** ATP phosphohydrolase (monosaccharide-importing)

**Comments:** ABC-type (ATP-binding cassette-type) ATPase, characterized by the presence of two similar ATP-

binding domains. Does not undergo phosphorylation during the transport process. Family of bacterial

enzymes importing ribose, xylose, arabinose, galactose and methylgalactoside.

**References:** [1138, 1541, 1419, 2495, 2721, 978]

[EC 3.6.3.17 created 2000]

[3.6.3.18 Transferred entry. oligosaccharide-transporting ATPase. Now EC 7.5.2.2, ABC-type oligosaccharide transporter]

[EC 3.6.3.18 created 2000, deleted 2018]

[3.6.3.19 Transferred entry. maltose-transporting ATPase. Now EC 7.5.2.1, ABC-type maltose transporter]

[EC 3.6.3.19 created 2000, deleted 2018]

[3.6.3.20 ATPase]	$Transferred\ entry.\ glycerol-3-phosphate-transporting\ ATPase.\ Now\ EC\ 7.6.2.10,\ glycerol-3-phosphate-transporting$
	[EC 3.6.3.20 created 2000, deleted 2018]
[3.6.3.21 porter]	Transferred entry. polar-amino-acid-transporting ATPase. Now EC 7.4.2.1, ABC-type polar-amino-acid trans-
	[EC 3.6.3.21 created 2000, deleted 2018]
[3.6.3.22 transporter]	Transferred entry. nonpolar-amino-acid-transporting ATPase. Now EC 7.4.2.2, ABC-type nonpolar-amino-acid
	[EC 3.6.3.22 created 2000, deleted 2018]
[3.6.3.23	Transferred entry. oligopeptide-transporting ATPase. Now EC 7.4.2.6, oligopeptide-transporting ATPase]
	[EC 3.6.3.23 created 2000, deleted 2018]
[3.6.3.24	Transferred entry. nickel-transporting ATPase. Now EC 7.2.2.11, nickel-transporting ATPase]
	[EC 3.6.3.24 created 2000, deleted 2018]
[3.6.3.25	Transferred entry. sulfate-transporting ATPase. Now EC 7.3.2.3, sulfate-transporting ATPase]
	[EC 3.6.3.25 created 2000, deleted 2018]
[3.6.3.26	Transferred entry. nitrate-transporting ATPase. Now EC 7.3.2.4, nitrate-transporting ATPase]
	[EC 3.6.3.26 created 2000, deleted 2018]
[3.6.3.27	Transferred entry. phosphate-transporting ATPase. Now EC 7.3.2.1, ABC-type phosphate transporter ]
	[EC 3.6.3.27 created 2000, deleted 2018]
[3.6.3.28	Transferred entry. phosphonate-transporting ATPase. Now EC 7.3.2.2, ABC-type phosphonate transporter]
	[EC 3.6.3.28 created 2000, deleted 2018]
[3.6.3.29	Transferred entry. molybdate-transporting ATPase. Now EC 7.3.2.5, molybdate-transporting ATPase]
	[EC 3.6.3.29 created 2000, deleted 2018]
[3.6.3.30	Transferred entry. Fe <sup>3+</sup> -transporting ATPase. Now EC 7.2.2.7, Fe <sup>3+</sup> -transporting ATPase]
	[EC 3.6.3.30 created 2000, deleted 2018]
[3.6.3.31	Transferred entry. polyamine-transporting ATPase. Now EC 7.6.2.11, polyamine-transporting ATPase]
	[EC 3.6.3.31 created 2000, deleted 2018]
[3.6.3.32 Pase]	Transferred entry. quaternary-amine-transporting ATPase. Now EC 7.6.2.9, quaternary-amine-transporting AT-
	[EC 3.6.3.32 created 2000, deleted 2018]
[3.6.3.33	Transferred entry. vitamin $B_{12}$ -transporting ATPase. Now EC 7.6.2.8, vitamin $B_{12}$ -transporting ATPase]
	[EC 3.6.3.33 created 2000, deleted 2018]
[3.6.3.34 7.2.2.16, iron citrate ABC 1	Transferred entry. iron-chelate-transporting ATPase; now recognized to be at least 3 separate enzymes EC n(III) hydroxamate ABC transporter, EC 7.2.2.17, ferric enterobactin ABC transporter, and EC 7.2.2.18, ferric transporter]
	[EC 3.6.3.34 created 2000, deleted 2018]
[3.6.3.35	Transferred entry. manganese-transporting ATPase. Now EC 7.2.2.5, manganese-transporting ATPase]

	[EC 3.6.3.35 created 2000, deleted 2018]
[3.6.3.36	Transferred entry. taurine-transporting ATPase. Now EC 7.6.2.7, taurine-transporting ATPase]
[	[EC 3.6.3.36 created 2000, deleted 2018]
[3.6.3.37	Transferred entry. guanine-transporting ATPase. Now EC 7.6.2.6, guanine-transporting ATPase]
[5.0.5.57	[EC 3.6.3.37 created 2000, deleted 2018]
[3.6.3.38	
transporter]	Transferred entry. capsular-polysaccharide-transporting ATPase. Now EC 7.6.2.2, ABC-type capsular-polysaccharide
	[EC 3.6.3.38 created 2000, deleted 2018]
[3.6.3.39 ATPase]	Transferred entry. lipopolysaccharide-transporting ATPase. Now EC 7.5.2.5, lipopolysaccharide-transporting
	[EC 3.6.3.39 created 2000, deleted 2018]
[3.6.3.40	Transferred entry. teichoic-acid-transporting ATPase. Now EC 7.5.2.4, teichoic-acid-transporting ATPase]
	[EC 3.6.3.40 created 2000, deleted 2018]
[3.6.3.41	Transferred entry. heme-transporting ATPase. Now EC 7.6.2.5, heme-transporting ATPase]
	[EC 3.6.3.41 created 2000, deleted 2018]
[3.6.3.42	Transferred entry. $\beta$ -glucan-transporting ATPase. Now EC 7.5.2.3, $\beta$ -glucan-transporting ATPase]
	[EC 3.6.3.42 created 2000, deleted 2018]
[3.6.3.43	Transferred entry. peptide-transporting ATPase. Now EC 7.4.2.5, peptide-transporting ATPase]
	[EC 3.6.3.43 created 2000, deleted 2018]
[3.6.3.44	Transferred entry. xenobiotic-transporting ATPase. Now EC 7.6.2.2, ABC-type xenobiotic transporter]
	[EC 3.6.3.44 created 2000 (EC 3.6.3.45 incorporated 2006), modified 2006, deleted 2018]
[3.6.3.45	Deleted entry. steroid-transporting ATPase. Now included with EC 3.6.3.44, xenobiotic-transporting ATPase]
	[EC 3.6.3.45 created 2000, deleted 2006]
[3.6.3.46	Transferred entry. cadmium-transporting ATPase. Now EC 7.2.2.2, ABC-type Cd <sup>2+</sup> transporter]
	[EC 3.6.3.46 created 2000, transferred 2018 to EC 7.2.2.2, deleted 2018]
[3.6.3.47	Transferred entry. fatty-acyl-CoA-transporting ATPase. Now EC 7.6.2.4, fatty-acyl-CoA-transporting ATPase]
	[EC 3.6.3.47 created 2000, deleted 2018]
[3.6.3.48	Transferred entry. α-factor-transporting ATPase. Now EC 7.4.2.7 as α-factor-pheromone transporting ATPase]
	[EC 3.6.3.48 created 2000, deleted 2018]
[3.6.3.49 ATPase]	Transferred entry. channel-conductance-controlling ATPase. Now EC 5.6.1.6, channel-conductance-controlling
	[EC 3.6.3.49 created 2000, deleted 2018]
[3.6.3.50	Transferred entry. protein-secreting ATPase. Now EC 7.4.2.8, protein-secreting ATPase]
	[EC 3.6.3.50 created 2000, deleted 2018]
[3.6.3.51 ATPase]	$Transferred\ entry.\ mitochondrial\ protein-transporting\ ATP ase.\ Now\ EC\ 7.4.2.3, mitochondrial\ protein-transporting$

[EC 3.6.3.51 created 2000, deleted 2018]

[3.6.3.52 Transferred entry. chloroplast protein-transporting ATPase. Now EC 7.4.2.4, chloroplast protein-transporting ATPase]

[EC 3.6.3.52 created 2000, deleted 2018]

[3.6.3.53 Transferred entry. Ag<sup>+</sup>-exporting ATPase. Now EC 7.2.2.15, Ag<sup>+</sup>-exporting ATPase]

[EC 3.6.3.53 created 2000, deleted 2018]

[3.6.3.54 Transferred entry. Cu<sup>+</sup>-exporting ATPase. Now EC 7.2.2.8, Cu<sup>+</sup>-exporting ATPase]

[EC 3.6.3.54 created 2013, deleted 2018]

[3.6.3.55 Transferred entry. tungstate-importing ATPase. Now EC 7.3.2.6, tungstate-importing ATPase]

[EC 3.6.3.55 created 2013, deleted 2018]

## EC 3.6.4 Acting on acid anhydrides to facilitate cellular and subcellular movement

[3.6.4.1 Transferred entry. myosin ATPase. Now EC 5.6.1.8, myosin ATPase]

[EC 3.6.4.1 created 1984 as EC 3.6.1.32, transferred 2000 to EC 3.6.4.1, deleted 2018]

[3.6.4.2 Transferred entry. dynein ATPase. Now EC 5.6.1.2, dynein ATPase]

[EC 3.6.4.2 created 1984 as EC 3.6.1.33, transferred 2000 to EC 3.6.4.2, deleted 2018]

[3.6.4.3 Transferred entry, microtubule-severing ATPase, Now EC 5.6.1.1, microtubule-severing ATPase]

[EC 3.6.4.3 created 2000 as 3.6.4.3, deleted 2018]

[3.6.4.4 Transferred entry. plus-end-directed kinesin ATPase. Now EC 5.6.1.3, plus-end-directed kinesin ATPase]

[EC 3.6.4.4 created 2000, deleted 2018]

[3.6.4.5 Transferred entry. minus-end-directed kinesin ATPase. Now EC 5.6.1.4, minus-end-directed kinesin ATPase]

[EC 3.6.4.5 created 2000, deleted 2018]

## EC 3.6.4.6

Accepted name: vesicle-fusing ATPase

**Reaction:** ATP +  $H_2O = ADP + phosphate$ **Systematic name:** ATP phosphohydrolase (vesicle-fusing)

**Comments:** A large family of ATP-hydrolysing enzymes involved in the heterotypic fusion of membrane vesicles

with target membranes and the homotypic fusion of various membrane compartments. They belong to the AAA-type ( $\underline{A}$ TPase  $\underline{a}$ ssociated with a variety of cell  $\underline{a}$ ctivities) ATPase superfamily. They include

peroxin, which apparently is involved in Zellweger's syndrome.

**References:** [472, 1239, 112]

[EC 3.6.4.6 created 2000]

## EC 3.6.4.7

**Accepted name:** peroxisome-assembly ATPase **Reaction:** ATP +  $H_2O = ADP + phosphate$ **Other name(s):** peroxisome assembly factor-2

**Systematic name:** ATP phosphohydrolase (peroxisome-assembling)

Comments: An extremely diversified group of enzymes that use the energy of ATP hydrolysis to import and as-

semble peroxisome components into the organelle. Their molecular masses range from 25 to 600

kDa.

**References:** [1638, 2982, 3240]

[EC 3.6.4.7 created 2000]

[3.6.4.8 Transferred entry, proteasome ATPase, Now EC 5.6.1.5, proteasome ATPase]

[EC 3.6.4.8 created 2000, deleted 2018]

[3.6.4.9 Transferred entry. chaperonin ATPase. Now EC 5.6.1.7, chaperonin ATPase]

[EC 3.6.4.9 created 2000, deleted 2018]

EC 3.6.4.10

**Accepted name:** non-chaperonin molecular chaperone ATPase

**Reaction:** ATP +  $H_2O = ADP + phosphate$ **Other name(s):** molecular chaperone Hsc70 ATPase

**Systematic name:** ATP phosphohydrolase (polypeptide-polymerizing)

**Comments:** This is a highly diverse group of enzymes that perform many functions that are similar to those of

chaperonins. They comprise a number of heat-shock-cognate proteins. They are also active in clathrin

uncoating and in the oligomerization of actin.

**References:** [2487, 244, 3140, 2747, 1673]

[EC 3.6.4.10 created 2000]

[3.6.4.11 Deleted entry. nucleoplasmin ATPase. The activity has been shown not to take place.]

[EC 3.6.4.11 created 2000, deleted 2018]

EC 3.6.4.12

Accepted name: DNA helicase

**Reaction:** ATP +  $H_2O = ADP + phosphate$ 

Other name(s): 3' to 5' DNA helicase; 3'-5' DNA helicase; 3'-5' PfDH; 5' to 3' DNA helicase; AvDH1; BACH1 he-

licase; BcMCM; BLM protein; BRCA1-associated C-terminal helicase; CeWR*N*-1; Dbp9p; Dm-RECQ5; DNA helicase 120; DNA helicase A; DNA helicase E; DNA helicase II; DNA helicase III; DNA helicase RECQL5β; DNA helicase VI; *dnaB*; DnaB helicase E1; helicase HDH IV; Hel E; helicase DnaB; helicase domain of bacteriophage T<sub>7</sub> gene 4 protein helicase; PcrA helicase; UvrD; hHcsA; Hmi1p; hPif1; MCM helicase; MCM protein; MER3 helicase; MER3 protein; MPH1; PcrA;

PcrA helicase; PDH120; PfDH A; Pfh1p; PIF1

**Systematic name:** ATP phosphohydrolase (DNA helix unwinding)

**Comments:** DNA helicases utilize the energy from ATP hydrolysis to unwind double-stranded DNA. Some of

them unwind duplex DNA with a 3' to 5' polarity [1,3,5,8], others show 5' to 3' polarity [10,11,12,13] or unwind DNA in both directions [2056, 2476]. Some helicases unwind DNA as well as RNA [814,

1279]. May be identical with EC 3.6.4.13 (RNA helicase).

**References:** [2219, 2885, 2034, 1617, 2273, 202, 2283, 521, 814, 1279, 1280, 3336, 890, 2056, 2476]

[EC 3.6.4.12 created 2009]

EC 3.6.4.13

**Accepted name:** RNA helicase

**Reaction:** ATP +  $H_2O = ADP + phosphate$ 

Other name(s): CSFV NS3 helicase; DBP2; DbpA; DDX17; DDX25; DDX3; DDX3X; DDX3Y; DDX4; DDX5;

DEAD-box protein DED1; DEAD-box RNA helicase; DEAH-box protein 2; DEAH-box RNA helicase; DED1; Dex(H/D) RNA helicase; EhDEAD1; EhDEAD1 RNA helicase; eIF4A helicase; KOKV helicase; Mtr4p; nonstructural protein 3 helicase; NPH-II; RHA; RNA helicase A; RNA helicase DDX3; RNA helicase Hera; RNA-dependent ATPase; TGBp1 NTPase/helicase domain; VRH1;

GRTH/DDX25

**Systematic name:** ATP phosphohydrolase (RNA helix unwinding)

**Comments:** RNA helicases utilize the energy from ATP hydrolysis to unwind RNA. Some of them unwind RNA

with a 3' to 5' polarity [1618], other show 5' to 3' polarity [?]. Some helicases unwind DNA as well as

RNA [814, ?]. May be identical with EC 3.6.4.12 (DNA helicase).

**References:** [487, 2445, 1618, 1670, 3218, 992, 814, ?]

[EC 3.6.4.13 created 2009]

## EC 3.6.5 Acting on GTP to facilitate cellular and subcellular movement

EC 3.6.5.1

**Accepted name:** heterotrimeric G-protein GTPase **Reaction:**  $GTP + H_2O = GDP + phosphate$  **Systematic name:** GTP phosphohydrolase (signalling)

**Comments:** This group comprises GTP-hydrolysing systems, where GTP and GDP alternate in binding. This

group includes stimulatory and inhibitory G-proteins such as  $G_s$ ,  $G_i$ ,  $G_o$  and  $G_{olf}$ , targetting adenylate cyclase and/or  $K^+$  and  $Ca^{2+}$  channels;  $G_q$  stimulating phospholipase C; transducin activating cGMP phosphodiesterase; gustducin activating cAMP phosphodiesterase.  $G_{olf}$  is instrumental in odour perception, transducin in vision and gustducin in taste recognition. At least 16 different  $\alpha$  subunits (39-

52 kDa), 5  $\beta$  subunits (36 kDa) and 12  $\gamma$  subunits (6-9 kDa) are known.

**References:** [2064, 2743, 261, 1924]

[EC 3.6.5.1 created 2000 as EC 3.6.1.46, transferred 2003 to EC 3.6.5.1]

EC 3.6.5.2

Accepted name: small monomeric GTPase

Reaction:  $GTP + H_2O = GDP + phosphate$ Systematic name: GTP + phosphohydrolase (cell-regulating)

Comments: A family of about 50 enzymes with a molecular mass of 21 kDa that are distantly related to the  $\alpha$ -

subunit of heterotrimeric G-protein GTPase (EC 3.6.5.1). They are involved in cell-growth regulation (Ras subfamily), membrane vesicle traffic and uncoating (Rab and ARF subfamilies), nuclear protein

import (Ran subfamily) and organization of the cytoskeleton (Rho and Rac subfamilies).

**References:** [270, 1026, 899, 3071]

[EC 3.6.5.2 created 2000 as EC 3.6.1.47, transferred 2003 to EC 3.6.5.2]

EC 3.6.5.3

**Accepted name:** protein-synthesizing GTPase **Reaction:** GTP +  $H_2O = GDP + phosphate$ 

Other name(s): elongation factor (EF); initiation factor (IF); peptide-release or termination factor

**Systematic name:** GTP phosphohydrolase (mRNA-translation-assisting)

Comments: This enzyme comprises a family of proteins involved in prokaryotic as well as eukaryotic protein syn-

thesis. In the initiation factor complex, it is IF-2b (98 kDa) that binds GTP and subsequently hydrolyses it in prokaryotes. In eukaryotes, it is eIF-2 (150 kDa) that binds GTP. In the elongation phase, the GTP-hydrolysing proteins are the EF-Tu polypeptide of the prokaryotic transfer factor (43 kDa), the eukaryotic elongation factor EF-1 $\alpha$  (53 kDa), the prokaryotic EF-G (77 kDa), the eukaryotic EF-2 (70-110 kDa) and the signal recognition particle that play a role in endoplasmic reticulum protein synthesis (325 kDa). EF-Tu and EF-1 $\alpha$  catalyse binding of aminoacyl-tRNA to the ribosomal A-site, while EF-G and EF-2 catalyse the translocation of peptidyl-tRNA from the A-site to the P-site. GT-Pase activity is also involved in polypeptide release from the ribosome with the aid of the pRFs and

eRFs.

**References:** [1576, 1469, 2447, 805, 1526]

[EC 3.6.5.3 created 2000 as EC 3.6.1.48, transferred 2003 to EC 3.6.5.3]

#### EC 3.6.5.4

Accepted name: signal-recognition-particle GTPase

**Reaction:**  $GTP + H_2O = GDP + phosphate$ 

**Systematic name:** GTP phosphohydrolase (protein-synthesis-assisting)

Comments: Activity is associated with the signal-recognition particle (a protein- and RNA-containing structure

involved in endoplasmic-reticulum-associated protein synthesis).

**References:** [479, 480, 1913, 813]

[EC 3.6.5.4 created 2000 as EC 3.6.1.49, transferred 2003 to EC 3.6.5.4]

#### EC 3.6.5.5

Accepted name: dynamin GTPase

**Reaction:**  $GTP + H_2O = GDP + phosphate$ 

**Systematic name:** GTP phosphohydrolase (vesicle-releasing)

Comments: An enzyme with a molecular mass of about 100 kDa that is involved in endocytosis and is instrumen-

tal in pinching off membrane vesicles.

**References:** [3130, 1863, 2154]

[EC 3.6.5.5 created 2000 as EC 3.6.1.50, transferred 2003 to EC 3.6.5.5]

#### EC 3.6.5.6

Accepted name: tubulin GTPase

**Reaction:**  $GTP + H_2O = GDP + phosphate$ 

**Systematic name:** GTP phosphohydrolase (microtubule-releasing)

**Comments:** An intrinsic activity of  $\alpha$ -tubulin involved in tubulin folding, division plane formation in prokaryotic

cells and others.

**References:** [3311, 2921, 2473]

[EC 3.6.5.6 created 2000 as EC 3.6.1.51, transferred 2003 to EC 3.6.5.6]

# EC 3.7 Acting on carbon-carbon bonds

This subclass contains a single sub-subclass for those enzymes that act on carbon-carbon bonds in ketonic substances (EC 3.7.1). There are relatively few carbon-carbon hydrolases and they mostly catalyse the hydrolysis of 3-oxo-carboxylic acids.

## EC 3.7.1 In ketonic substances

## EC 3.7.1.1

**Accepted name:** oxaloacetase

**Reaction:** oxaloacetate +  $H_2O$  = oxalate + acetate

**Other name(s):** oxalacetic hydrolase

Systematic name: oxaloacetate acetylhydrolase

**References:** [1085]

[EC 3.7.1.1 created 1961]

## EC 3.7.1.2

Accepted name: fumarylacetoacetase

**Reaction:** 4-fumarylacetoacetate +  $H_2O$  = acetoacetate + fumarate

Other name(s): β-diketonase; fumarylacetoacetate hydrolase
Systematic name: 4-fumarylacetoacetate fumarylhydrolase
Comments: Also acts on other 3,5- and 2,4-dioxo acids.

**References:** [482, 678, 1880]

[EC 3.7.1.2 created 1961]

EC 3.7.1.3

Accepted name: kynureninase

**Reaction:** L-kynurenine +  $H_2O$  = anthranilate + L-alanine

**Systematic name:** L-kynurenine hydrolase

**Comments:** A pyridoxal-phosphate protein. Also acts on 3'-hydroxy-L-kynurenine and some other (3-

arylcarbonyl)-alanines.

**References:** [1300, 1299, 1483, 3201]

[EC 3.7.1.3 created 1965]

EC 3.7.1.4

**Accepted name:** phloretin hydrolase

**Reaction:** phloretin +  $H_2O$  = phloretate + phloroglucinol

**Other name(s):** ErPhy; lactase-phlorerin hydrolase; *C*-acylphenol hydrolase; 2',4,4',6'-tetrahydroxydehydrochalcone

1,3,5-trihydroxybenzenehydrolase (incorrect)

**Systematic name:** phloretin acylhydrolase (phloroglucinol forming)

**Comments:** Also hydrolyses other *C*-acylated phenols related to phloretin. Isolated from the fungus *Aspergillus* 

niger and the bacteria Pantoea agglomerans and Eubacterium ramulus.

**References:** [403, 1923, 2580]

[EC 3.7.1.4 created 1972, modified 2018]

EC 3.7.1.5

Accepted name: acylpyruvate hydrolase

**Reaction:** a 3-acylpyruvate +  $H_2O$  = a carboxylate + pyruvate

**Systematic name:** 3-acylpyruvate acylhydrolase

**Comments:** Acts on formylpyruvate, 2,4-dioxopentanoate, 2,4-dioxohexanoate and 2,4-dioxoheptanoate.

**References:** [3137]

[EC 3.7.1.5 created 1976]

EC 3.7.1.6

**Accepted name:** acetylpyruvate hydrolase

**Reaction:** acetylpyruvate +  $H_2O$  = acetate + pyruvate **Systematic name:** 2,4-dioxopentanoate acetylhydrolase

**Comments:** Highly specific; does not act on pyruvate, oxaloacetate, maleylpyruvate, fumarylpyruvate or acety-

lacetone.

**References:** [542]

[EC 3.7.1.6 created 1984]

EC 3.7.1.7

**Accepted name:** β-diketone hydrolase

**Reaction:** nonane-4,6-dione +  $H_2O$  = pentan-2-one + butanoate

**Other name(s):** oxidized PVA hydrolase

**Systematic name:** nonane-4,6-dione acylhydrolase

Comments: Also acts on the product of the action of EC 1.1.3.18 secondary-alcohol oxidase, on polyvinyl alco-

hols; involved in the bacterial degradation of polyvinyl alcohol.

**References:** [2501, 2502]

[EC 3.7.1.7 created 1989]

EC 3.7.1.8

**Accepted name:** 2,6-dioxo-6-phenylhexa-3-enoate hydrolase

**Reaction:** 2,6-dioxo-6-phenylhexa-3-enoate +  $H_2O$  = benzoate + 2-oxopent-4-enoate

**Other name(s):** HOHPDA hydrolase

**Systematic name:** 2,6-dioxo-6-phenylhexa-3-enoate benzoylhydrolase

**Comments:** Cleaves the products from biphenol, 3-isopropylcatechol and 3-methylcatechol produced by EC

1.13.11.39 biphenyl-2,3-diol 1,2-dioxygenase, by ring-fission at a -CO-C bond. Involved in the break-

down of biphenyl-related compounds by Pseudomonas sp.

**References:** [2196]

[EC 3.7.1.8 created 1989]

EC 3.7.1.9

**Accepted name:** 2-hydroxymuconate-6-semialdehyde hydrolase

**Reaction:** 2-hydroxymuconate-6-semialdehyde +  $H_2O$  = formate + 2-oxopent-4-enoate

**Other name(s):** 2-hydroxy-6-oxohepta-2,4-dienoate hydrolase; 2-hydroxymuconic semialdehyde hydrolase;

HMSH; HOD hydrolase; xylF (gene name); 2-hydroxymuconate-semialdehyde formylhydrolase; 2-

hydroxymuconate-semialdehyde hydrolase

**Systematic name:** 2-hydroxymuconate-6-semialdehyde formylhydrolase

**Comments:** The enzyme is involved in the degradation of catechols.

**References:** [2508, 1050, 603]

[EC 3.7.1.9 created 1990, modified 2013]

EC 3.7.1.10

**Accepted name:** cyclohexane-1,3-dione hydrolase

**Reaction:** cyclohexane-1,3-dione +  $H_2O = 5$ -oxohexanoate

Other name(s): 1,3-cyclohexanedione hydrolase; cyclohexane-1,3-dione acylhydrolase (decyclizing)

**Systematic name:** cyclohexane-1,3-dione acylhydrolase (ring-opening)

**Comments:** Highly specific; does not act on other dione derivatives of cyclohexane, cyclopentane or cycloheptane.

**References:** [534]

[EC 3.7.1.10 created 1992]

EC 3.7.1.11

**Accepted name:** cyclohexane-1,2-dione hydrolase

**Reaction:** cyclohexane-1,2-dione + H<sub>2</sub>O = 6-oxohexanoate **Other name(s):** cyclohexane-1,2-dione acylhydrolase (decyclizing) **Systematic name:** cyclohexane-1,2-dione acylhydrolase (ring-opening)

**Comments:** Highly specific; does not act on cyclohexanone or cyclohexane-1,3-dione as substrate.

**References:** [1051, 800]

[EC 3.7.1.11 created 2009]

## EC 3.7.1.12

Accepted name: cobalt-precorrin 5A hydrolase

**Reaction:** cobalt-precorrin-5A +  $H_2O$  = cobalt-precorrin-5B + acetaldehyde +  $2 H^+$ 

Other name(s): CbiG

Systematic name: cobalt-precorrin 5A acylhydrolase

Comments: This enzyme hydrolyses the ring A acetate  $\delta$ -lactone of cobalt-precorrin-5A resulting in the loss of the

C-20 carbon and its attached methyl group in the form of acetaldehyde. This is a key reaction in the contraction of the porphyrin-type tetrapyrrole ring and its conversion to a corrin ring in the anaerobic

(early cobalt insertion) adenosylcobalamin biosynthesis pathway.

**References:** [1357, 1958]

[EC 3.7.1.12 created 2010]

## EC 3.7.1.13

**Accepted name:** 2-hydroxy-6-oxo-6-(2-aminophenyl)hexa-2,4-dienoate hydrolase

**Reaction:** (2E,4E)-6-(2-aminophenyl)-2-hydroxy-6-oxohexa-2,4-dienoate +  $H_2O$  = anthranilate + (2E)-2-

hydroxypenta-2,4-dienoate

Other name(s): CarC

**Systematic name:** (2E,4E)-6-(2-aminophenyl)-2-hydroxy-6-oxohexa-2,4-dienoate acylhydrolase

**Comments:** This enzyme catalyses the third step in the aerobic degradation pathway of carbazole. The effect of

the presence of an amino group or hydroxyl group at the 2-position of the substrate is small. The en-

zyme has no cofactor requirement [2424].

**References:** [2105, 2424]

[EC 3.7.1.13 created 2010]

#### EC 3.7.1.14

**Accepted name:** 2-hydroxy-6-oxonona-2,4-dienedioate hydrolase

**Reaction:** (1) (2Z,4E)-2-hydroxy-6-oxonona-2,4-diene-1,9-dioate + H<sub>2</sub>O = (2Z)-2-hydroxypenta-2,4-dienoate +

succinate

(2) (2Z,4E,7E)-2-hydroxy-6-oxonona-2,4,7-triene-1,9-dioate + H<sub>2</sub>O = (2Z)-2-hydroxypenta-2,4-

dienoate + fumarate

Other name(s): mhpC (gene name)

**Systematic name:** (2Z,4E)-2-hydroxy-6-oxona-2,4-dienedioate succinylhydrolase

Comments: This enzyme catalyses a step in a pathway of phenylpropanoid compounds degradation. The first step

of the enzyme mechanism involves a reversible keto-enol tautomerization [1593].

**References:** [328, 329, 1592, 1593, 757, 602]

[EC 3.7.1.14 created 2011, modified 2012]

[3.7.1.15 Transferred entry. (+)-caryolan-1-ol synthase. Now EC 4.2.1.138, (+)-caryolan-1-ol synthase]

[EC 3.7.1.15 created 2011, deleted 2013]

[3.7.1.16 Transferred entry. oxepin-CoA hydrolase. Now EC 3.3.2.12, oxepin-CoA hydrolase]

[EC 3.7.1.16 created 2011, deleted 2013]

## EC 3.7.1.17

**Accepted name:** 4,5:9,10-diseco-3-hydroxy-5,9,17-trioxoandrosta-1(10),2-diene-4-oate hydrolase

**Reaction:** (1E,2Z)-3-hydroxy-5,9,17-trioxo-4,5:9,10-disecoandrosta-1(10),2-dien-4-oate + H<sub>2</sub>O = 3-

[(3aS,4S,7aS)-7a-methyl-1,5-dioxo-octahydro-1H-inden-4-yl]propanoate + (2Z,4Z)-2-hydroxyhexa-

2,4-dienoate

**Other name(s):** *tesD* (gene name); *hsaD* (gene name)

Systematic name: 4,5:9,10-diseco-3-hydroxy-5,9,17-trioxoandrosta-1(10),2-diene-4-oate hydrolase ((2Z,4Z)-2-

hydroxyhexa-2,4-dienoate-forming)

**Comments:** The enzyme is involved in the bacterial degradation of the steroid ring structure, and is involved in

degradation of multiple steroids, such as testosterone [1179], cholesterol [586], and sitosterol.

**References:** [1179, 586, 1585, 1586]

[EC 3.7.1.17 created 2012]

EC 3.7.1.18

**Accepted name:** 6-oxocamphor hydrolase

**Reaction:** bornane-2,6-dione +  $H_2O = [(1S)-4-hydroxy-2,2,3-trimethylcyclopent-3-enyl]acetate$ 

Other name(s): OCH; *camK* (gene name)

Systematic name: bornane-2,6-dione hydrolase

**Comments:** Isolated from *Rhodococcus* sp. The bornane ring system is cleaved by a retro-Claisen reaction to

give the enol of  $\alpha$ -campholonate. When separate from the enzyme the enol is tautomerised to the keto form as a 6:1 mixture of [(1S,3R)-2,2,3-trimethyl-4-oxocyclopentyl]acetate and [(1S,3S)-2,2,3-trimethyl-4-oxocyclopentyl]acetate acetate acetate and [(1S,3S)-2,2,3-trimethyl-4-oxocyclopentyl]acetate acetate acetat

trimethyl-4-oxocyclopentyl]acetate.

**References:** [990, 3175, 1648]

[EC 3.7.1.18 created 2012]

EC 3.7.1.19

**Accepted name:** 2,6-dihydroxypseudooxynicotine hydrolase

**Reaction:** 1-(2,6-dihydroxypyridin-3-yl)-4-(methylamino)butan- $1-\text{one} + \text{H}_2\text{O} = 2,6-\text{dihydroxypyridine} + 4-\text{one}$ 

methylaminobutanoate

**Systematic name:** 1-(2,6-dihydroxypyridin-3-yl)-4-(methylamino)butan-1-one hydrolase

**Comments:** The enzyme, characterized from the soil bacterium *Arthrobacter nicotinovorans*, participates in nico-

tine degradation.

**References:** [902, 2485]

[EC 3.7.1.19 created 2012]

EC 3.7.1.20

**Accepted name:** 3-fumarylpyruvate hydrolase

**Reaction:** 3-fumarylpyruvate +  $H_2O$  = fumarate + pyruvate

**Other name(s):** *nagK* (gene name); *naaD* (gene name)

**Systematic name:** 3-fumarylpyruvate hydrolase

**Comments:** The enzyme is involved in bacterial degradation of 5-substituted salicylates, including gentisate (5-

hydroxysalicylate), 5-nitrosalicylate and 5-halosalicylates.

**References:** [3337, 2340]

[EC 3.7.1.20 created 2012]

EC 3.7.1.21

Accepted name: 6-oxocyclohex-1-ene-1-carbonyl-CoA hydratase

**Reaction:** 6-oxocyclohex-1-ene-1-carbonyl-CoA +  $\mathbf{2}$  H<sub>2</sub>O = 3-hydroxypimeloyl-CoA (overall reaction)

(1a) 6-oxocyclohex-1-ene-1-carbonyl-CoA +  $H_2O$  = 2-hydroxy-6-oxocyclohexane-1-carbonyl-CoA

(1b) 2-hydroxy-6-oxocyclohexane-1-carbonyl-CoA +  $H_2O$  = 3-hydroxypimeloyl-CoA

Other name(s): 6-oxocyclohex-1-ene-1-carbonyl-CoA hydrolase; 6-oxocyclohex-1-ene-1-carbonyl-CoA hydrolase

(decyclizing)

**Systematic name:** 6-oxocyclohex-1-ene-1-carbonyl-CoA hydrolase (ring-opening)

Comments: The enzyme, which participates in the anaerobic benzoyl-CoA degradation pathway in certain organ-

isms, catalyses the addition of one molecule of water to the double bound of 6-oxocyclohex-1-ene-1-

carbonyl-CoA followed by the hydrolytic C-C cleavage of the alicyclic ring.

**References:** [1589, 1557]

## [EC 3.7.1.21 created 2014]

EC 3.7.1.22

**Accepted name:** 3D-(3,5/4)-trihydroxycyclohexane-1,2-dione acylhydrolase (ring-opening) **Reaction:** 3D-3,5/4-trihydroxycyclohexa-1,2-dione +  $H_2O = 5$ -deoxy-D-glucuronate

Other name(s): IoID; THcHDO hydrolase; 3D-3,5/4-trihydroxycyclohexa-1,2-dione hydrolase (decyclizing); 3D-

(3,5/4)-trihydroxycyclohexane-1, 2-dione acylhydrolase (decyclizing)

**Systematic name:** 3D-3,5/4-trihydroxycyclohexa-1,2-dione hydrolase (ring-opening)

**Comments:** The enzyme, found in the bacterium *Bacillus subtilis*, is part of the *myo*-inositol degradation pathway

leading to acetyl-CoA.

**References:** [3299]

[EC 3.7.1.22 created 2014, modified 2014]

EC 3.7.1.23

Accepted name: maleylpyruvate hydrolase

**Reaction:** 3-maleylpyruvate +  $H_2O$  = maleate + pyruvate

Other name(s): hbzF (gene name)

**Systematic name:** (2Z)-4,6-dioxohept-2-enedioate acylhydrolase

Comments: The enzyme, characterized from the bacterium *Pseudomonas alcaligenes* NCIMB 9867, catalyses the

hydrolysis of 3-maleylpyruvate, the ring-cleavage product of gentisate. The enzyme can also act on a number of maleylpyruvate derivatives, such as (2E)-2-methyl-4,6-dioxohept-2-enedioate and (2E)-3-methyl-4,6-dioxohept-2-enedioate. Activated by  $\mathrm{Mn}^{2+}$ . May be identical to EC 3.7.1.5, acylpyruvate

hydrolase.

**References:** [1175, 178, 1722]

[EC 3.7.1.23 created 2016]

# EC 3.8 Acting on halide bonds

This subclass contains enzymes that hydrolyse carbon-halide compounds in a single sub-subclass (EC 3.8.1).

### EC 3.8.1 In carbon-halide compounds

EC 3.8.1.1

Accepted name: alkylhalidase

**Reaction:** bromochloromethane  $+ H_2O =$  formaldehyde + bromide + chloride **Other name(s):** halogenase; haloalkane halidohydrolase; haloalkane dehalogenase

**Systematic name:** alkyl-halide halidohydrolase

**References:** [1125]

[EC 3.8.1.1 created 1961]

EC 3.8.1.2

**Accepted name:** (S)-2-haloacid dehalogenase

**Reaction:** (S)-2-haloacid +  $H_2O = (R)$ -2-hydroxyacid + halide

**Other name(s):** 2-haloacid dehalogenase[ambiguous]; 2-haloacid halidohydrolase [ambiguous][ambiguous]; 2-

haloalkanoic acid dehalogenase; 2-haloalkanoid acid halidohydrolase; 2-halocarboxylic acid dehalo-

genase II; DL-2-haloacid dehalogenase[ambiguous]; L-2-haloacid dehalogenase; L-DEX

**Systematic name:** (S)-2-haloacid halidohydrolase

**Comments:** Acts on acids of short chain lengths,  $C_2$  to  $C_4$ , with inversion of configuration at C-2. [See also EC

3.8.1.9 (R)-2-haloacid dehalogenase, EC 3.8.1.10 2-haloacid dehalogenase (configuration-inverting)

and EC 3.8.1.11 2-haloacid dehalogenase (configuration-retaining)]

**References:** [949, 1990, 1471, 611, 1983, 1495, 1987, 1567, 2707]

[EC 3.8.1.2 created 1972, modified 2003]

EC 3.8.1.3

Accepted name: haloacetate dehalogenase

**Reaction:** haloacetate +  $H_2O$  = glycolate + halide

Other name(s): monohaloacetate dehalogenase
Systematic name: haloacetate halidohydrolase

**References:** [946, 948]

[EC 3.8.1.3 created 1972]

[3.8.1.4 Transferred entry. thyroxine deiodinase. Now EC 1.97.1.10, thyroxine 5'-deiodinase]

[EC 3.8.1.4 created 1984, deleted 2003]

EC 3.8.1.5

Accepted name: haloalkane dehalogenase

**Reaction:** 1-haloalkane +  $H_2O$  = a primary alcohol + halide

Other name(s): 1-chlorohexane halidohydrolase; 1-haloalkane dehalogenase

**Systematic name:** 1-haloalkane halidohydrolase

**Comments:** Acts on a wide range of 1-haloalkanes, haloalcohols, haloalkenes and some haloaromatic compounds.

**References:** [1431, 2583, 3287]

[EC 3.8.1.5 created 1989]

EC 3.8.1.6

**Accepted name:** 4-chlorobenzoate dehalogenase

**Reaction:** 4-chlorobenzoate +  $H_2O = 4$ -hydroxybenzoate + chloride

Other name(s): halobenzoate dehalogenase

Systematic name: 4-chlorobenzoate chlorohydrolase

Comments: Catalyses the first step in the degradation of chlorobenzoate in *Pseudomonas*. In many microor-

ganisms, this activity comprises three separate enzymes, EC 6.2.1.33 (4-chlorobenzoate—CoA ligase), EC 3.8.1.7 (4-chlorobenzoyl-CoA dehalogenase) and EC 3.1.2.23 (4-hydroxybenzoyl-CoA

thioesterase).

**References:** [1998, 1125]

[EC 3.8.1.6 created 1989, modified 1999]

EC 3.8.1.7

**Accepted name:** 4-chlorobenzoyl-CoA dehalogenase

**Reaction:** 4-chlorobenzoyl-CoA +  $H_2O$  = 4-hydroxybenzoyl CoA + chloride

**Systematic name:** 4-chlorobenzoyl CoA chlorohydrolase

Comments: Specific for dehalogenation at the 4-position. Can dehalogenate substrates bearing fluorine, chlorine,

bromine and iodine in the 4-position. This enzyme is part of the bacterial 2,4-dichlorobenzoate degra-

dation pathway.

**References:** [394, 508]

[EC 3.8.1.7 created 1999]

### EC 3.8.1.8

Accepted name: atrazine chlorohydrolase

**Reaction:** atrazine +  $H_2O$  = hydroxyatrazine + chloride

Other name(s): AtzA

**Systematic name:** atrazine chlorohydrolase

**Comments:** Involved in the degradation of the herbicide atrazine, 2-chloro-4-(ethylamino)-6-(isopropylamino)-

1,3,5-triazine, in bacteria.

**References:** [562, 561]

[EC 3.8.1.8 created 2000, modified 2011]

### EC 3.8.1.9

**Accepted name:** (R)-2-haloacid dehalogenase

**Reaction:** (*R*)-2-haloacid +  $H_2O = (S)$ -2-hydroxyacid + halide

Other name(s): 2-haloalkanoic acid dehalogenase[ambiguous]; 2-haloalkanoid acid halidohydrolase[ambiguous]; D-

2-haloacid dehalogenase; D-DEX

**Systematic name:** (*R*)-2-haloacid halidohydrolase

**Comments:** Acts on acids of short chain lengths, C<sub>2</sub> to C<sub>4</sub>, with inversion of configuration at C-2. [See also EC

3.8.1.2 (S)-2-haloacid dehalogenase, EC 3.8.1.10 2-haloacid dehalogenase (configuration-inverting)

and EC 3.8.1.11 2-haloacid dehalogenase (configuration-retaining)]

**References:** [2698, 1641, 2707]

[EC 3.8.1.9 created 2003]

### EC 3.8.1.10

**Accepted name:** 2-haloacid dehalogenase (configuration-inverting)

**Reaction:** (1) (S)-2-haloacid +  $H_2O = (R)$ -2-hydroxyacid + halide

(2) (R)-2-haloacid +  $H_2O = (S)$ -2-hydroxyacid + halide

Other name(s): 2-haloalkanoic acid dehalogenase; 2-haloalkanoid acid halidohydrolase; DL-2-haloacid dehaloge-

nase; DL-2-haloacid dehalogenase (inversion of configuration); DL-2-haloacid halidohydrolase (inversion of configuration); DL-DEXi; (*R*,*S*)-2-haloacid dehalogenase (configuration-inverting)

**Systematic name:** (S)-2-haloacid dehalogenase (configuration-inverting)

**Comments:** Dehalogenates both (S)- and (R)-2-haloalkanoic acids to the corresponding (R)- and (S)-

hydroxyalkanoic acids, respectively, with inversion of configuration at C-2. The enzyme from *Pseudomonas* sp. 113 acts on 2-haloalkanoic acids whose carbon chain lengths are five or less. [See also EC 3.8.1.2 (*S*)-2-haloacid dehalogenase, EC 3.8.1.9 (*R*)-2-haloacid dehalogenase and EC 3.8.1.11 2-

haloacid dehalogenase (configuration-retaining)]

**References:** [1987, 1989, 1988, 1567, 1721, 354, 1641, 3149, 2707]

[EC 3.8.1.10 created 2003]

### EC 3.8.1.11

Accepted name: 2-haloacid dehalogenase (configuration-retaining)

**Reaction:** (1) (S)-2-haloacid +  $H_2O = (S)$ -2-hydroxyacid + halide

(2) (R)-2-haloacid +  $H_2O = (R)$ -2-hydroxyacid + halide

Other name(s): 2-haloalkanoic acid dehalogenase; 2-haloalkanoid acid halidohydrolase; DL-2-haloacid dehaloge-

nase; DL-DEXr

**Systematic name:** (S)-2-haloacid dehalogenase (configuration-retaining)

**Comments:** Dehalogenates both (S)- and (R)-2-haloalkanoic acids to the corresponding (S)- and (R)-

hydroxyalkanoic acids, respectively, with retention of configuration at C-2. [See also EC 3.8.1.2 (*S*)-2-haloacid dehalogenase, EC 3.8.1.9 (*R*)-2-haloacid dehalogenase and EC 3.8.1.10 2-haloacid dehalogenase

genase (configuration-inverting)]

**References:** [3149, 2707]

## EC 3.8.2 In phosphorus-halide compounds (deleted sub-subclass)

[3.8.2.1 Transferred entry. di-isopropyl-fluorophosphatase. Now EC 3.1.8.2, diisopropyl-fluorophosphatase]

[EC 3.8.2.1 created 1961, modified 1976, deleted 1992]

# EC 3.9 Acting on phosphorus-nitrogen bonds

This subclass contains a single sub-subclass for enzymes that act on phosphorus-nitrogen bonds (EC 3.9.1).

## EC 3.9.1 Acting on phosphorus-nitrogen bonds (only sub-subclass identified to date)

#### EC 3.9.1.1

Accepted name: phosphoamidase

**Reaction:** N-phosphocreatine +  $H_2O$  = creatine + phosphate

Other name(s): creatine phosphatase

Systematic name: phosphamide hydrolase

**Comments:** Also acts on N-phospho-arginine and other phosphoamides. Possibly identical with EC 3.1.3.9

(glucose-6-phosphatase) or EC 3.1.3.16 (protein-serine/threonine phosphatase).

**References:** [2242, 2668, 2801]

[EC 3.9.1.1 created 1961]

## EC 3.9.1.2

**Accepted name:** protein arginine phosphatase

**Reaction:** a [protein]- $N^{0}$ -phospho-L-arginine + H<sub>2</sub>O = a [protein]-L-arginine + phosphate

Other name(s): YwlE

**Systematic name:** [protein]- $N^{\omega}$ -phospho-L-arginine phosphohydrolase

Comments: The enzyme, characterized from Gram-positive bacteria, hydrolyses the phosphoramidate (P-N) bond

of  $N^{\omega}$ -phospho-L-arginine residues in proteins and peptides that were phosphorylated by EC 2.7.14.1,

protein-arginine-kinase.

**References:** [831, 2962, 695]

[EC 3.9.1.2 created 2014]

## EC 3.9.1.3

**Accepted name:** phosphohistidine phosphatase

**Reaction:** a [protein]-N-phospho-L-histidine +  $H_2O$  = a [protein]-L-histidine + phosphate

**Other name(s):** PHPT1 (gene name); protein histidine phosphatase; PHP **Systematic name:** [protein]-*N*-phospho-L-histidine phosphohydrolase

Comments: This eukaryotic enzyme dephosphorylates phosphorylated histidine residues within proteins and pep-

tides. The enzyme acts on phosphate groups attached to both the *pros*- and *tele*-nitrogen atoms, but the *pros*- position is somewhat preferred (by a factor of two at the most) [95]. The substrate specificity depends on the amino acid sequence or structural context of the phosphohistidine in a phosphoprotein. The enzyme is also active on free phosphoramidate [689, 95] and peptide-bound phospholysine [688].

**References:** [689, 1478, 176, 95, 688]

[EC 3.9.1.3 created 2016]

# EC 3.10 Acting on sulfur-nitrogen bonds

This subclass contains a single sub-subclass for enzymes that act on sulfur-nitrogen bonds (EC 3.10.1).

## EC 3.10.1 Acting on sulfur-nitrogen bonds (only sub-subclass identified to date)

EC 3.10.1.1

**Accepted name:** *N*-sulfoglucosamine sulfohydrolase

**Reaction:** N-sulfo-D-glucosamine +  $H_2O = D$ -glucosamine + sulfate

Other name(s): sulfoglucosamine sulfamidase; heparin sulfamidase; 2-desoxy-D-glucoside-2-sulphamate sulphohy-

drolase (sulphamate sulphohydrolase)

**Systematic name:** *N*-sulfo-D-glucosamine sulfohydrolase

**References:** [609, 1775]

[EC 3.10.1.1 created 1972, modified 1981, modified 1982]

EC 3.10.1.2

Accepted name: cyclamate sulfohydrolase

**Reaction:** cyclohexylsulfamate  $+ H_2O =$  cyclohexylamine + sulfate

Other name(s): cyclamate sulfamatase; cyclamate sulfamidase; cyclohexylsulfamate sulfamidase

Systematic name: cyclohexylsulfamate sulfohydrolase

**Comments:** Also readily hydrolyses aliphatic sulfamates with 3 to 8 carbons.

References: [2084]

[EC 3.10.1.2 created 1976, modified 1981]

# EC 3.11 Acting on carbon-phosphorus bonds

This subclass contains a single sub-subclass for enzymes that hydrolyse C-phosphono-groups (EC 3.11.1).

## EC 3.11.1 Acting on carbon-phosphorus bonds (only sub-subclass identified to date)

EC 3.11.1.1

**Accepted name:** phosphonoacetaldehyde hydrolase

 $\begin{tabular}{ll} \textbf{Reaction:} & phosphonoacetal dehyde + $H_2O$ = acetal dehyde + phosphote \\ \textbf{Other name(s):} & phosphonoacetylal dehyde phosphonohydrolase \\ \end{tabular}$ 

**Systematic name:** 2-oxoethylphosphonate phosphonohydrolase

**Comments:** This enzyme destabilizes the C-P bond, by forming an imine between one of its lysine residues and

the carbonyl group of the substrate, thus allowing this, normally stable, bond to be broken. The mechanism is similar to that used by EC 4.1.2.13, fructose-bisphosphate aldolase, to break a C-C bond.

Belongs to the haloacetate dehalogenase family.

**References:** [2060, 2061, 2059, 2193, 124]

[EC 3.11.1.1 created 1972, modified 1976, modified 2001]

EC 3.11.1.2

Accepted name: phosphonoacetate hydrolase

**Reaction:** phosphonoacetate +  $H_2O$  = acetate + phosphate

Systematic name: phosphonoacetate phosphonohydrolase

Comments: A zinc-dependent enzyme. Belongs to the alkaline phosphatase superfamily of zinc-dependent hydro-

lases.

**References:** [1871]

[EC 3.11.1.2 created 1999]

EC 3.11.1.3

**Accepted name:** phosphonopyruvate hydrolase

**Reaction:** 3-phosphonopyruvate +  $H_2O$  = pyruvate + phosphate

Other name(s): PPH

**Comments:** Highly specific for phosphonopyruvate as substrate [1546]. The reaction is not inhibited by phosphate

but is inhibited by the phosphonates phosphonoformic acid, hydroxymethylphosphonic acid and 3-phosphonopropanoic acid [1546]. The enzyme is activated by the divalent cations Co<sup>2+</sup>, Mg<sup>2+</sup> and Mn<sup>2+</sup>. This enzyme is a member of the phospho*enol*pyruvate mutase/isocitrate lyase superfamily

[411].

**References:** [2904, 1546, 411]

[EC 3.11.1.3 created 2007]

# EC 3.12 Acting on sulfur-sulfur bonds

This subclass contains a single sub-subclass for enzymes that act on sulfur-sulfur bonds (EC 3.12.1).

## EC 3.12.1 Acting on sulfur-sulfur bonds (only sub-subclass identified to date)

EC 3.12.1.1

Accepted name: trithionate hydrolase

**Reaction:** trithionate +  $H_2O$  = thiosulfate + sulfate +  $2 H^+$ 

**Systematic name:** trithionate thiosulfohydrolase

**References:** [1746, 2966]

[EC 3.12.1.1 created 1990]

# EC 3.13 Acting on carbon-sulfur bonds

This subclass contains a single sub-subclass for enzymes that act on carbon-sulfur bonds (EC 3.13.1).

## EC 3.13.1 Acting on carbon-sulfur bonds (only sub-subclass identified to date)

EC 3.13.1.1

Accepted name: UDP-sulfoquinovose synthase

**Reaction:** UDP- $\alpha$ -D-sulfoquinovopyranose + H<sub>2</sub>O = UDP- $\alpha$ -D-glucose + sulfite

Other name(s): sulfite:UDP-glucose sulfotransferase; UDPsulfoquinovose synthase; UDP-6-sulfo-6-deoxyglucose

sulfohydrolase

Systematic name: UDP-6-sulfo-6-deoxy- $\alpha$ -D-glucose sulfohydrolase

**Comments:** Requires NAD<sup>+</sup>, which appears to oxidize UDP-α-D-glucose to UDP-4-dehydroglucose, which dehy-

drates to UDP-4-dehydro-6-deoxygluc-5-enose, to which sulfite is added. The reaction is completed when the substrate is rehydrogenated at C-4. The enzyme from *Arabidopsis thaliana* is specific for

UDP-Glc and sulfite.

**References:** [719, 720, 1993, 2515]

[EC 3.13.1.1 created 2001, modified 2010]

[3.13.1.2 Deleted entry. 5-deoxyribos-5-ylhomocysteinase. The activity is most probably attributable to EC 4.4.1.21, S-ribosylhomocysteine lyase]

[EC 3.13.1.2 created 1972 as EC 3.3.1.3, transferred 2001 to EC 3.2.1.148, transferred 2004 to EC 3.13.1.2, deleted 2005]

### EC 3.13.1.3

**Accepted name:** 2'-hydroxybiphenyl-2-sulfinate desulfinase

**Reaction:** 2'-hydroxybiphenyl-2-sulfinate +  $H_2O = 2$ -hydroxybiphenyl + sulfite

Other name(s): gene dszB-encoded hydrolase; 2-(2-hydroxyphenyl) benzenesulfinate: H<sub>2</sub>O hydrolase; DszB;

HBPSi desulfinase; 2-(2-hydroxyphenyl) benzenesulfinate sulfohydrolase; HPBS desulfinase; 2-(2-hydroxyphenyl)benzenesulfinate hydrolase; 2-(2'-hydroxyphenyl)benzenesulfinate desulfinase; 2-(2-hydroxyphenyl)benzenesulfinate desulfinase; 2-(2-hydroxyphenyl)benzenesulfinase; 2-(2-hydroxyphenyl)benzenes

 $hydroxyphenyl) benzene sulfinate\ desulfinase$ 

**Systematic name:** 2'-hydroxybiphenyl-2-sulfinate sulfohydrolase

**Comments:** The enzyme from *Rhodococcus* sp. strain IGTS8 is encoded by the plasmid-encoded

dibenzothiophene-desulfurization (dsz) operon. The enzyme has a narrow substrate specificity with

biphenyl-2-sulfinate being the only other substrate known to date [2053].

**References:** [2192, 2053, 3136]

[EC 3.13.1.3 created 2000 as EC 3.1.2.24, transferred 2005 to EC 3.13.1.3]

### EC 3.13.1.4

**Accepted name:** 3-sulfinopropanoyl-CoA desulfinase

**Reaction:** 3-sulfinopropanoyl-CoA +  $H_2O$  = propanoyl-CoA + sulfite

Other name(s): 3SP-CoA desulfinase; AcdDPN7; 3-sulfinopropionyl-CoA desulfinase

**Systematic name:** 3-sulfinopropanoyl-CoA sulfinohydrolase

Comments: The enzyme from the  $\beta$ -proteobacterium *Advenella mimigardefordensis* contains one non-covalently

bound FAD per subunit.

**References:** [2595, 2594]

[EC 3.13.1.4 created 2014]

### EC 3.13.1.5

Accepted name: carbon disulfide hydrolase

**Reaction:** carbon disulfide  $+ 2 H_2O = CO_2 + 2$  hydrogen sulfide (overall reaction)

(1a) carbon disulfide +  $H_2O$  = carbonyl sulfide + hydrogen sulfide

(1b) carbonyl sulfide +  $H_2O = CO_2$  + hydrogen sulfide

Other name(s): CS2 hydrolase (misleading); carbon disulfide lyase; CS2-converting enzyme; carbon disulphide-lyase

(decarboxylating)

**Systematic name:** carbon-disulfide hydrogen-sulfide-lyase (decarboxylating)

**Comments:** The enzyme contains  $Zn^{2+}$ . The hyperthermophilic archaeon *Acidianus* sp. A1-3 obtains energy by

the conversion of carbon disulfide to hydrogen sulfide, with carbonyl sulfide as an intermediate.

References: [2695]

[EC 3.13.1.5 created 2013 as EC 4.4.1.27, transferred 2017 to EC 3.13.1.5]

### EC 3.13.1.6

**Accepted name:** [CysO sulfur-carrier protein]-S-L-cysteine hydrolase

**Reaction:** [CysO sulfur-carrier protein]-Gly-NH-CH<sub>2</sub>-C(O)-S-L-cysteine +  $H_2O$  = [CysO sulfur-carrier protein]-

Gly-NH-CH<sub>2</sub>-COOH + L-cysteine

**Other name(s):** *mec* (gene name)

**Systematic name:** [CysO sulfur-carrier protein]-S-L-cysteine sulfohydrolase

**Comments:** Requires  $Zn^{2+}$ . The enzyme, characterized from the bacterium *Mycobacterium tuberculosis*, partic-

ipates in an L-cysteine biosynthesis pathway. It acts on the product of EC 2.5.1.113, [CysO sulfur-

carrier protein]-thiocarboxylate-dependent cysteine synthase.

**References:** [330]

[EC 3.13.1.6 created 2017]

### EC 3.13.1.7

Accepted name: carbonyl sulfide hydrolase

**Reaction:** carbonyl sulfide +  $H_2O$  = hydrogen sulfide +  $CO_2$ 

**Other name(s):** COSase; COS hydrolase; cos (gene name)

**Systematic name:** carbonyl sulfide hydrogen-sulfide-lyase (decarboxylating)

Comments: The enzyme, characterized from the bacterium *Thiobacillus thioparus*, catalyses a step in the degrada-

tion pathway of thiocyanate. This activity is also catalysed by the archaeal EC 3.13.1.5, carbon disul-

fide lyase.

**References:** [2151]

[EC 3.13.1.7 created 2018]

#### EC 3.13.1.8

**Accepted name:** S-adenosyl-L-methionine hydrolase (adenosine-forming) **Reaction:** S-adenosyl-L-methionine  $+ H_2O = adenosine + L-methionine$ 

**Other name(s):** SAM hydroxide adenosyltransferase

**Systematic name:** S-adenosyl-L-methionine hydrolase (adenosine-forming)

Comments: The enzyme, found in bacteria and archaea, catalyses a nucleophilic attack of water at the C5' carbon

of *S*-adenosyl-L-methionine to generate adenosine and L-methionine. May be involved in regulating SAM levels in the cell. *cf.* EC 3.3.1.2, *S*-adenosyl-L-methionine hydrolase (L-homoserine-forming).

**References:** [722, 583]

[EC 3.13.1.8 created 2018]

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