Evolutionary families of peptidases

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The available amino acid sequences of peptidases have been examined, and the enzymes have been allocated to evolutionary families. Some of the families can be grouped together in 'clans' that show signs of distant relationship, but nevertheless, it appears that there may be as many as 60 evolutionary lines of

peptidases with separate origins. Some of these contain members with quite diverse peptidase activities, and yet there are some striking examples of convergence. We suggest that the classification by families could be used as an extension of the current classification by catalytic type.

INTRODUCTION

Amino acid sequence data are now available for over 600 peptidases (endopeptidases, exopeptidases and omega peptidases), and we have examined these in an attempt to establish what separate evolutionary lines exist. These take the form of families, or groups of related families ('clans'). The properties of the peptidases of each family have been considered from two main points of view. Firstly, we have asked how widely the enzymes have diverged in catalytic activity, and, secondly, we have asked to what extent peptidases from separate evolutionary lines have converged in properties. Finally, we have considered how compatible is a classification of peptidases based on their evolutionary relationships with the sort of classification that is currently in use, which depends upon the reaction catalysed by each enzyme and on the catalytic mechanism.

METHODS

Sources of data

Protein sequence data were obtained from the SwissProt database [1] (release 21), and the PIR-Protein database [2] (release 32), and nucleic acid sequence data from the EMBL database [1] (release 28 and daily updates). In addition, some sequences were obtained directly from the literature.

Detection of evolutionary relationships

The programs FASTP [3] and FASTA and TFASTA [4] were used to detect similarities between peptidases, and, on the basis of these, provisional assignments to a system of families was made. These assignments were refined by manual construction of optimized alignments. In many cases, the similarities between the sequences were so close that no further analysis was felt necessary, but whenever the similarity was questionable, the RDF program [3] was applied. This tests the statistical significance of a similarity between amino acid sequences by comparing the score for the alignment with those of random shuffles of the sequences. We took the value of six standard deviation units as that above which the similarity could be regarded as being significant. We assume that the significant similarities reflect evolutionary relationship, or homology as defined by Reeck et al. [5].

Definition of terms

The term type is used to refer to a set of peptidases distinguished according to the chemical groups responsible for catalysis, as in serine-type, cysteine-type, aspartic-type or metallo-type. The

term family is used to describe a group of enzymes in which each member shows evolutionary relationship to at least one other, either throughout the whole sequence or at least in the part of the sequence responsible for catalytic activity. As an example of the need for this, bone morphogenetic protein 1 is a chimaeric protein that contains a catalytic domain related to that of astacin, but also contains segments that are clearly homologous with non-catalytic parts of C1r and C1s in the chymotrypsin family [6]. We place bone morphogenetic protein 1 in the family of astacin and not in that of chymotrypsin.

A clan comprises a group of families for which there are indications of evolutionary relationship, despite the lack of statistically significant similarities in sequence. Such indications of distant relationship come primarily from the linear order of catalytic-site residues and the tertiary structure. Distinctive aspects of the catalytic activity such as specificity or inhibitor-sensitivity may also contribute occasionally.

The symbol '+' is used to indicate the scissile bond in a peptidase substrate.

RESULTS AND DISCUSSION

All of the amino acid sequences of peptidases that were available to us in July 1992 were examined for significant similarities as described in the Methods section, and grouped in families (Table 1). Some of the families show evidence of distant relationships to others, and these we group together in single 'clans'; others seem quite unrelated.

Serine peptidases

Most of the members of the chymotrypsin (S1) family are endopeptidases, which differ widely in specificity. No exopeptidase is known in this family, but it does contain several proteins that lack all peptidase activity: azurocidin, procarboxypeptidase A complex component III, the haptoglobins, apolipoprotein a, hepatocyte growth factor and protein Z. The family includes many enzymes of the coagulation, fibrinolysis and complement systems that are found in blood plasma, and these are mostly chimaeric proteins with modules, some of which are also found in other proteins, inserted N-terminally to the site of proteolytic activation [27].

Almost all of the known members of the chymotrypsin family have been found in animals, the only exceptions being two trypsins from actinomycetes. It is striking that no member of this otherwise very successful family has been encountered in protozoa, fungi or plants.

The linear order of catalytic triad residues in the polypeptide

Table 1 Evolutionary families of peptidases

The peptidases are allocated to families as described in the text. Clans and families are labelled with the prefix S for serine peptidases, C for cysteine, A for aspartic, M for metallo- and U for unknown, and listed in this order. It should be noted, however, that these labels are temporary, simply being assigned consecutively through the Table. 'EC' is the enzyme nomenclature number [7], but for peptidases the initial '3.4.' has been omitted; '-' indicates that no EC number has been assigned; 'n.a.' indicates that the protein is not known to be an enzyme. Literature references to the individual proteins are generally to be found in the database entries for which the codes are given. Most of the codes are from the Swiss-Prot database (release 21), but a code in parentheses is an EMBL database accession number and 'PIR' indicates a code from the PIR database. Numbers in square brackets are references to sequences from journal articles. For some viral sequences, the code given is that of the viral polyprotein. For some viruses, numerous variants with only minor differences exist, and only a single example of each has been included.

EC Database code

SERINE PEPTIDASES

Family S1: Chymotrypsin (Cla	an SA: Hi	is, Asp, Ser catalytic triad)
Trypsin (includes forms I, II, III, IV	21.4	TRYP_SACER, TRYP_STRGR, TRYP_ASTFL, TRYP_DROME,
Va and Vb)		TRYP_SQUAC, TRYP_XENLA, TRYP_BOVIN, TRY1_CANFA,
		TRY2_CANFA, TRY1_HUMAN, TRY2_HUMAN, TRY3_HUMAN,
		TRYP_MOUSE, TRYP_PIG, TRY1_RAT, TRY2_RAT, TRY3_RAT,
		TRY4_RAT, (M77814), (X59012), (X59013)
Cercarial elastase (Schistosoma)	-	CERC_SCHMA
Brachyurin	21.32	COGS_UCAPU
Factor C (Limulus)	-	(D90271)
Proclotting enzyme (Tachypleus)	-	PCE_TACTR
easter gene product (Drosophila)	-	EAST_DROME
snake gene product (Drosophila)	-	SNAK DROME
Vitellin-degrading endopeptidase Bombyx)	-	[8]
Hypodermin C	21.49	COGS_HYPLI
Serine proteases 1 and 2 (Drosophila)	-	SER1 DROME
Achelase (Lonomia)	-	ACH1 LONAC, ACH2 LONAC
Chymotrypsin (includes forms A, B, II and	2) 21.1	CTR2 VESCR, CTR2 VESOR, CTRA_BOVIN, CTRB_BOVIN,
, , , , ,		CTR2_CANFA, CTRB_HUMAN, CTRB_RAT
Proteinase RVV-V (Russell's viper)	-	RVVA_VIPRU, RVVG_VIPRU
(includes forms α and γ)		-
Flavoboxin (habu snake)	-	FLVB TRIFL
Venombin A	21.74	BATX_BOTAT, PTCA_AGKCO
Crotalase	21.74	-
Enteropeptidase	21.9	[10]
Acrosin	21.10	ACRO_HUMAN, ACRO_MOUSE, ACRO_PIG
Seminin	_	PROS HUMAN
Tissue kallikrein	21.35	KAG2_CAVPO, KAG1_HUMAN, KAG2_HUMAN, KAG_PIG,
		KAGP RAT
Renal kallikrein	21.35	KAGR_MOUSE, (X17352)
Submandibular kallikrein	21.35	KAG1_MOUSE, KAG2_MOUSE, KAG3_MOUSE, KAG5_MOUSE,
		KAGB_MOUSE, KAG1_RAT, KAG3_RAT
7S nerve growth factor (includes α and γ chains)	21.35	-
Epidermal growth factor-binding protein (includes forms 1, 2 and 3)	21.35	EGBA_MOUSE, EGBB_MOUSE, EGBC_MOUSE
Tonin	21.35	TONI RAT
Arginine esterase		ESTA CANFA
Pancreatic elastase I		EL1 PIG, EL1 RAT, (M27347)
Pancreatic elastase II (includes forms A	21.71	
and B)		EL2_RAT
Pancreatic endopeptidase E (includes	21.70	
forms A and B)		- · · -
Leukocyte elastase	21.37	ELNE, HUMAN
Medullasin	-	ELNE HUMAN
Azurocidin	n.a.	CAP7 HUMAN, CAP7_PIG
		<u> </u>

	Table 1 (contd.)					
	Cathepsin G	21.20	CATG_HUMAN			
	Proteinase 3 (myeloblastin)	•	MELB_HUMAN, PTN3_HUMAN			
	Chymase (includes forms I and II)	21.39	-			
	•		MCP1_RAT, MCP2_RAT, MCP4_MOUSE, (M69136), (M73759)			
	γ-Renin	21.54	RENG_MOUSE			
	Tryptase (includes forms 1, 2 and 3)	21.59	TRYT_CANFA, TRYA_HUMAN, TRYB_HUMAN, (M33493), (M30038),			
			MCP6_MOUSE			
	Hepsin	-	HEPS_HUMAN			
	Granzyme A	-	GRAA_HUMAN, GRAA_MOUSE, GRAX_MOUSE			
	Natural killer cell protease 1	-	NKP1_RAT			
	Granzymes B, C, D, E, F, G and Y	-	GRAB_MOUSE, GRAC_MOUSE, GRAD_MOUSE, GRAE_MOUSE,			
			GRAF_MOUSE, GRAG_MOUSE, GRAB_HUMAN, GRAY_HUMAN			
	Carboxypeptidase A complex component III		CAC3_BOVIN			
	Complement factor D	21.46	CFAD_HUMAN, ADIP_MOUSE			
	Complement factor B		CFAB_HUMAN, CFAB_MOUSE			
	Complement factor I		CFAI_HUMAN			
	Complement component CTr		CO1R_HUMAN			
	Complement component CTs	21.42	C1S_HUMAN			
	Calcium-dependent serine proteinase	-	CASP_MESAU			
	Complement component C2	21.43	CO2_HUMAN, CO2_MOUSE			
	Haptoglobin (includes forms 1 and 2)	n.a.	HPT1_HUMAN, HPT2_HUMAN			
	Haptoglobin-related protein	n.a.	HPTR_HUMAN			
	Plasmin	21.7	PLMN_BOVIN, PLMN_HUMAN, PLMN_MACMU, PLMN_MOUSE,			
	A matter a man A atta (a)		PLMN_PIG, (M62832)			
	Apolipoprotein(a)	n.a.	APOA_HUMAN, APOA_MACMU			
	Hepatocyte growth factor	n.a.	HGF_HUMAN, HGF_RAT			
	Thrombin	21.5	THRB_BOVIN, THRB_HUMAN, THRB_MOUSE, THRB_RAT			
	t-Plasminogen activator	21.68	UROT_HUMAN, UROT_MOUSE, UROT_RAT			
	u-Plasminogen activator	21.73	UROK_CHICK, UROK_HUMAN, UROK_MOUSE, UROK_PAPCY,			
	Coliner, planning on a stirle	04.00	UROK_PIG			
	Salivary plasminogen activator	21.68	UROT_DESRO			
	(vampire bat)	04.04	IZAL ANDRAAN IZAL DAT (NATORO)			
	Plasma kallikrein		KAL_HUMAN, KAL_RAT, (M58588)			
	Coagulation factor VII		FA7_BOVIN, FA7_HUMAN			
	Coagulation factor IX	21.22	FA9_BOVIN, FA9_CANFA, FA9_HUMAN, FA9_MOUSE			
	Coagulation factor X	21.6	FA10_BOVIN, FA10_HUMAN			
	Coagulation factor XI Coagulation factor XII	21.27	FA11_HUMAN			
	Protein C	21.38	FA12_HUMAN			
	Protein Z	21.69	PRTC_BOVIN, PRTC_HUMAN			
		n.a.	PTRZ_BOVIN, PRTZ_HUMAN			
1			is, Asp, Ser catalytic triad)			
	α-Lytic endopeptidase	21.12	PRLA_LYSEN			
	Proteases A and B (Streptomyces griseus)	-	PRTA_STRGR, PRTB_STRGR			
	Glutamyl endopeptidase (Strep. griseus)	<u>-</u>	[11]			
ı		n SA: Hi	is, Asp, Ser catalytic triad)			
	Polyprotein peptidase	-	POLS_EEEV, POLS_RRVN, POLS_SFV, POLS_SINDV, POLS_WEEV			
1	Family S4: Glutarnyl endopeptidase					
	Glutamyl endopeptidase (Staphylococcus)	21.19	_			
	Epidermolytic toxins A and B	-	ETA_STAAU, ETB_STAAU			
	(Staphylococcus)					
	"Metalloprotease" (Bacillus subtilis)	•	[12]			
	Family S5: Lysyl endopeptidase					
	Lysyl endopeptidase (Achromobacter)	21.50	API_ACHLY			
	Family S6: IgA-specific endopeptidase					
	In A anacific carino andonantidasa	04 70	ICA NEICO (V64367)			

21.72 IGA_NEIGO, (X64357)

IgA-specific serine endopeptidase

Family S7: Flavivirus endopeptidase

Nonstructural protein NS3 - POLG_DEN2J, POLG_JAEVJ, POLG_KUNJM, POLG_MVEV, POLG TBEVS, POLG WNV, POLG YEFV1

Family S8: Subtilisin (Asp, His, Ser catalytic triad)
Tripeptidyl-peptidase II 14.10 (M73047)

Subtilisin 21.62 SUBT BACAM, SUBT BACLI, SUBT BACMS, SUBT_BACSA,

SUBT BACSD, SUBT BACSU

Alkaline elastase (*Bacillus*) - ELYA_BACSU
Serine endopeptidase (*Bac. subtilis*) - (PIR S11504)

Major intracellular endopeptidase (Bacillus) - ISP1 BACSU, (D00862), (D10730)

Bacillopeptidase F (Bac. subtilis) - SUBF_BACSU

Neutral endopeptidase (Bacillus) - NPRE BACAM, NPRE BACSU

Thermitase 21.66 THET_THEVU
C5a peptidase (Streptococcus) - SCPA_STRPY

Cell-wall associated endopeptidase - P1P_LACLA, P2P_LACLA, P3P_LACLA

(Lactococcus) (forms PI, PII, PIII)

Aqualysin I (*Thermus*) - AQL1_THEAQ

Extracellular endopeptidase (*Serratia*) - PRTS_SERMA

Calcium-dependent extracellular - PROA_VIBAL

endopeptidase A (Vibrio)

Extracellular endopeptidase (Xanthomonas) - PIR S11890
Endopeptidase K 21.64 PRTK_TRIAL
Endopeptidase R (Tritirachium) - PRTR_TRIAL
Endopeptidase T (Tritirachium) - PRTT_TRIAL
Cuticle-degrading protease (Metarhizium) - (M73795)

Oryzin 21.63 AEP ASPOR, AEP YARLI

Alkaline protease (*Aspergillus*) - (Z11580)

Cerevisin 21.48 PRTB_YEAST

Subtilisin-like protease III - (M77197)

(Saccharomyces)

Alkaline endopeptidase (*Acremonium*) - PIR JU0332
Calcium dependent endopeptidase - PRCA_ANAVA

(Anabaena)

Kexin 21.61 KEX2_YEAST, KEX1_KLULA

Furin - FURI_HUMAN, FURI_MOUSE, FURI_RAT, (M81431)
Pituitary convertase (includes PC1 and PC2) - NEC1_MOUSE, NEC2_HUMAN, NEC2_MOUSE

Family S9: Prolyl oligopeptidase (Asp, Ser, His or Ser, Asp, His catalytic triad)

Dipeptidyl-peptidase IV 14.5 DPP_RAT, (X60708)

Dipeptidyl aminopeptidase B (Saccharomyces)- DAP2_YEAST

Acylaminoacyl-peptidase 19.1 ACPH_PIG, ACPH_RAT

Protease II (Escherichia coli) - TLP_ECOLI

Prolyl oligopeptidase 21.26 PPCE_PIG, (M81461), (M61966)

DNF1552 protein (3p21 protein) n.a. DNF1_HUMAN

Family S10: Serine-type carboxypeptidase (Ser, Asp, His catalytic triad)

Serine-type carboxypeptidase 16.1 CBPY YEAST, (D10199)

(Saccharomyces)

Carboxypeptidase B-like peptidase 16.1 KEX1 YEAST, CBP2_HORVU, CBP2_WHEAT,

Serine-type carboxypeptidase (forms I 16.1 CBP1_HORVU, CBP3_HORVU, CBP3_WHEAT, (D10985)

and III)

Carboxypeptidase Y-like protein - (M81130)

(Arabidopsis)

Serine-type carboxypeptidase - (M75784)

(Caenorhabditis)

Serine-type carboxypeptidase (Aedes) - (M79452)

Lysosomal carboxypeptidase A 16.1 PRTP_HUMAN, PRTP_MOUSE

Table 1 (contd.) Family S11: D-Ala-D-Ala carboxypeptidase (gene daca) (Clan SB: Ser, Lys, Ser, Glu catalytic tetrad) Serine-type D-Ala-D-Ala carboxypeptidase 16.4 DACA BACSU, DACA ECOLI, DACC ECOLI, (X59965), (M37688) Family S12: D-Ala-D-Ala carboxypeptidase (gene dac) (Clan SB: Ser, Lys, Ser, Glu catalytic tetrad) Serine-type D-Ala-D-Ala carboxypeptidase 16.4 DAC STRSP D-Aminopeptidase (Ochrobactrum) (M84523) **B-lactamase** 3.5.2.6 AMPC CITFR, AMPC ECOLI, AMPC ENTCL, AMPC SERMA Protein FIMD (Bacteroides) FMDH BACNO, FMDD BACNO Family S13: Penicillin-binding protein 4 (Clan SE: Ser, Lys, Ser, Glu catalytic tetrad) Serine-type D-Ala-D-Ala carboxypeptidase 16.4 [13] Penicillin-binding protein 4 PBP4 ECOLI Family S14: Clop (Ser, His catalytic residues (Asp not known)) ATP-dependent endopeptidase (ClpP subunit)-CLPP ECOLI (Escherichia coli) Chloroplast ATP-dependent endopeptidase CLPP MARPO, CLPP_TOBAC, CLPP_ORYSA, CLPP_WHEAT Potato leaf roll luteovirus genomic RNA n.a. (D00530), (X14600) Family S15: Lactococcus dipeptidyl peptidase IV Dipeptidyl peptidase IV (Lactococcus) 14.5 DPP_LACLA, DPP_LACLC Family S16: Endopeptidase La 21.53 LON_ECOLI, (D00863) Endopeptidase La Family S17: Bacteroides endopeptidase Extracellular endopeptidase (Bacteroides) PRTE BACNO Family S18: Endopeptidase VII OMPT ECOLI Protease VII (Escherichia coli) Coagulase/fibrinolysin (Yersinia) COLY_YERPE Phosphoglycerate transport system activator -PGTE SALTY (Salmonella) Family S19: Coccidioides endopeptidase Chymotrypsin-like protease (Coccidioides) (X63114)Family S20: Protease Do Protease Do (Salmonella) (X54548)Family S21: Assemblin, herpesvirus Assemblin UL26_HSV11, VG33_VZVD, CP40_ILV, YEC3_EBV, UL80_HCMVA, (M64627) Family S22: Placental protein 11 Placental protein 11 PP11 HUMAN CYSTEINE PEPTIDASES Family C1: Papain (Clan CA: Gln, Cys, His, Asn active site residues) Dipeptidyl peptidase I 14.1 (D90404) Cysteine endopeptidases 1 (Haemonchus) CYS1 HAECO, Cysteine endopeptidases 1 (Haemonchus) (M80385) Surface protective protein (*Plasmodium*) n.a. [14] Circumsporozoite protein (*Plasmodium*) CSP PLACM Cysteine endopeptidase (Entamoeba) (M27307), (M64712), (M64721) Cysteine endopeptidase (Trypanosoma) CYSP TRYBR Cruzipain (Trypanosoma) (M90067) CYSP_THEPA, (M86659) Cysteine endopeptidase (Theileria) Cysteine endopeptidase (Leishmania) (X62163)Cysteine endopeptidases 1 and 2 CYS1 DICDI, CYS2 DICDI (Dictyostelium) Endopeptidase (baculovirus of Autographa) (M67451)

22.2

22.6

PAPA_CARPA
PAP2_CARPA

22.30 PAP3_CARPA

Papain

Chymopapain Caricain

		Table 1 (contd.)		
Glycyl endopeptidase	22.25	PAP4_CARPA		
Actinidain		ACTN ACTCH		
Cysteine endopeptidase (tomato)	-	CYSL_LYCES		
Thaumatopain (Thaumatococcus)	-	THPA_THADA		
Calotropin (Calotropis)	-	CAL1_CALGI		
Cysteine endopeptidase (Brassica napus)		[15]		
Cysteine endopeptidase (mung bean)	-	SHEP_VIGMU		
Endopeptidase EP-C1 (Phaseolus vulgari		(X63102)		
Protein P34 (soya bean)	n.a.	P34 SOYBN		
Clone 15a protein (garden pea)	-	[16]		
Stem bromelain	22.32	BROM ANACO		
Aleurain (barley)	-	ALEU_HORVU		
Cysteine endopeptidases 2 and 3 (barley)	-	[17]		
Oryzain (includes forms α , β and γ) (rice)	-	[18]		
Cysteine protease (Caenorhabditis)	-	(M74797)		
Cysteine endopeptidases 1, 2 and 3 (Homarus)	-	(X63567), (X63568), (X63569)		
Allergen (Dermatophagoides)	•	MMAL_DERPT		
Allergen (Euroglyphus)	-	(X60073)		
Cathepsin L	22.15	CATL_CHICK, CATL_HUMAN, CATL_MOUSE, CATL_RAT		
Cathepsin S	22.27	CATS_BOVIN, (M86553)		
Cathepsin H	22.16	CATH_HUMAN, CATH_RAT		
Cathepsin B	22.1	CATB_BOVIN, CATB_HUMAN, CATB_MOUSE, CATB_RAT, (M75822), (M21309)		
	lan CA: G	In, Cys, His, Asn active site residues)		
Sol gene product (Drosophila)	-	(M64084)		
Calpain (Schistosoma)		(M67499)		
Calpain I		CAP1_CHICK, CAP1_HUMAN, CAP1_RABIT		
Calpain II		CAP2_HUMAN, CAP2_RABIT		
Calpain P94		CAP3_HUMAN, CAP3_RAT		
Calcium-binding protein PMP41	22.17	CAP4_MOUSE		
Family C3: Picornain (C	lan CB: H	is, Asp or Glu, Cys catalytic triad)		
Picornain 2A		POLG_POL1M, POLG_COXA2, POLG_SVDVH, POLG_BOVEV,		
		POLG HRV14		
Picornain 3C	22.28	POLH_POL1M, POLG_COXA2, POLG_SVDVH, POLG_BOVEV,		
		POLG_HRV14, POLG_ECHO9, POLG_TMEVD		
Aphthovirus endopeptidase		POLG_FMDVD		
Cardiovirus endopeptidase		POLG_EMCV		
Comovirus endopeptidase		VGNB_CPMV, (D00657)		
Family C4: Potyvirus endopeptidase 1 (C	lan CB: H	is, Asp, Cys catalytic triad)		
48 kDa endopeptidase	-	POLG_PPVD, POLG_PPVRA, POLG_PPVYN, POLG_TEV,		
		POLG_TVMV, POLG_WMV2, POLG_OMV		
	lan CB: H	is, Cys catalytic triad)		
Endopeptidase adenovirus	•	VPRT_ADEB3, VPRT_ADEB7, VPRT_ADE02, VPRT_ADE03,		
		VPRT_ADE04, VPRT_ADE05, VPRT_ADE12, VPRT_ADE40,		
_		VPRT_ADE41, (M81056)		
Family C6: Potyvirus endopeptidase 2				
29 kDa endopeptidase	•	POLG_PPVD, POLG_PVYN, POLG_TEV, POLG_TVMV		
Family C7: Chestnut blight virus p29 endop		(1457020)		
p29 Endopeptidase (Chestnut blight virus) - (M57938)				
Family C8: Chestnut blight virus p48 endopeptidase p48 Endopeptidase (Chestnut blight virus) - (M57938)				
Family C9: Togavirus cysteine endopeptida	IS 0	DOLNI CINIDA DOLNI DOLNI CENTOLNI ONNIVO		
Togavirus cysteine endopeptidase	-	POLN_SINDV, POLN_RRVN, POLN_SFV,POLN_ONNVG,		

Family C10: Streptopain

Streptopain 22.10 STCP STRPY

Family C11: Clostripain

α-Clostripain 22.8 CLOL_CLOHi

Family C12: Ubiquitin hydrolase

Ubiquitin carboxyl-terminal hydrolase - UBL1 HUMAN, UBL3_HUMAN, [19]

Family C13: Haemoglobinase

Haemoglobinase (Schistosoma) - HGLB SCHMA

Family C14: Interleukin-1B converting enzyme

Interleukin-1ß converting enzyme - [20]

ASPARTIC PEPTIDASES

Family A1: Pepsin (Clan AA: Asp, Asp catalytic residues)

Aspergillopepsin I 23.18 PEPA_ASPAW Penicillopepsin 23.20 PENP_PENJA

Rhizopuspepsin 23.21 CARP_RHICH, CARP_RHINI,

Endothiapepsin 23.22 CARP_CRYPA

Mucorpepsin 23.23 CARP_RHIMI, CARP_RHIPU

Candidapepsin 23.24 CARP_CANAL, (X61438), (Z11918), (M83663), (X56867),

(Z11919)

Polyporopepsin 23.29 CARP_IRPLA

Saccharopepsin 23.25 CARP SACFI, CARP YEAST, (D10198)

"Barrier" protein (*Saccharomyces*) - BAR1_YEAST Aspartic proteinase (barley) - (X56136)

Pepsin A 23.1 PEPA_CHICK, PEPA_BOVIN, PEPA_HUMAN, PEPA_MACFU,

PEPA MACMU, PEPA_PIG,

Aspartic endopeptidase P111 - PIR JT0398

Gastricsin 23.3 PEPC HUMAN, PEPC MACFU, PEPC_RAT

Chymosin 23.4 CHYM BOVIN, CHYM SHEEP

Embryonic pepsin (chicken) - PEPE_CHICK
Renin, submandibular 23.15 RENS_MOUSE

Renin, renal 23.15 RENI HUMAN, RENI MOUSE, RENI_RAT

Cathepsin D 23.5 CATD HUMAN, CATD MOUSE, CATD_PIG, CATD_RAT

Cathepsin E 23.34 CATE_HUMAN

Family A2: Retropepsin (Clan AA: Asp, Asp catalytic residues)

Retropepsin 23.16 POL_HIV1A, POL_HIV2D, POL_SIVMK, POL_BIV06, POL_EIAV,

POL_VILV, VPRT_MPMV, VPRT_MMTVB, GAG_RSVP, VPRT_BLV, POL_FLV, POL_GALV, VPRT_HTL1A, POL_MLVAV, VPRT_SMRVH,

VPRT_SRV1

Retrovirus-related endopeptidase (human) - VPRT_HUMAN

Retropepsin-like protein (vaccinia virus) - (M25392)

METALLO-PEPTIDASES

Family M1: Alanyl aminopeptidase (Clan MA: Peptidases with HEXXH zinc-binding motif)

Membrane alanyl aminopeptidase 11.2 AMPN_ECOLI, AMPN_HUMAN, AMPN_PIG, AMPN_RAT, (X51508),

(M75750)

Lysyl aminopeptidase (*Lactococcus*) 11.15 (X61230)
Aminopeptidase yscll (*Saccharomyces*) - (X63998)

BP-1/6C3 antigen, mouse - BP1_MOUSE

Leukotriene A₄ hydrolase 3.3.2.6 LKHA_HUMAN, (M63848)

Family M2: Peptidyl-dipeptidase A (Clan MA: Peptidases with HEXXH zinc-binding motif)

Peptidyl-dipeptidase A 15.1 ACE_HUMAN, ACET_HUMAN, ACE_MOUSE, ACET_MOUSE,

ACE_RABIT, ACET_RABIT

		Table 1 (contd.)
Family M3: Thimet oligopeptidase	(Clan MA: Pe	eptidases with HEXXH zinc-binding motif)
Peptidyl-dipeptidase, bacterial	-	(X57947), (M84575)
Oligopeptidase (Salmonella)	-	(M84574)
Mitochondrial intermediate peptidase	-	(M96633)
Saccharolysin	24.37	(X59720 - orf YCL57w)
Thimet oligopeptidase		MEPD_RAT
Family M4: Thermolysin	(Clan MA: Pe	eptidases with HEXXH zinc-binding motif)
Thermolysin		THER_BACST, THER_BACTH
Pseudolysin	24.26	ELAS_PSEAE
Neutral endopeptidase (Bacillus	-	PIR B36706
stearothermophilus)		
Bacillolysin	24.28	THER_BACCE, THER_BACCL, NPRE_BACSU, (D00861), (K02497),
Matalla and an antida as (1 a sign alls)		(M64815), (X61380)
Metalloendopeptidase (Legionella)	-	PROA_LEGPN
Vibriolysin (Vibrio)	-	(M64809), (M59466)
Extracellular endopeptidase (<i>Erwinia</i>) Metalloendopeptidase (<i>Listeria</i>)	-	(M36651)
Coccolysin	- 24.30	PROL_LISMO (M37185)
Family M5: Mycolysin		eptidases with HEXXH zinc-binding motif)
Mycolysin	24.31	· · ·
Family M6: Immune inhibitor A		eptidases with HEXXH zinc-binding motif)
Immune inhibitor A (Bacillus	-	INA_BACTL
thuringiensis)		110 (_D) (0 12
=	protease(Clar	n MA: Peptidases with HEXXH zinc-binding motif)
Small neutral protease (Streptomyces	•	(M81703), (M86606), (Z11929)
Family M8: Leishmanolysin	•	eptidases with HEXXH zinc-binding motif)
Leishmanolysin		GP63_LEICH, GP63_LEIDO, GP63_LEIMA, (X64394)
Family M9: Microbial collagenase		eptidases with HEXXH zinc-binding motif)
Collagenase (Vibrio)	24.3	[21]
Family M10: Interstitial collagenase	(Clan MA: Po	eptidases with HEXXH zinc-binding motif)
Serralysin	24.40	PRTB_ERWCH, PRTC_ERWCH, PRTX_ERWCH, PRZN_SERSP
Envelysin		HE_PARLI
Matrilysin		COG7_HUMAN
Interstitial collagenase	24.7	COG1_HUMAN, COG1_PIG, COG1_RABIT
Neutrophil collagenase		COG8_HUMAN
Stromelysin 1		COG3_HUMAN, COG3_RABIT, COG3_RAT
Stromelysin 2	24.22	COGX_HUMAN, COGX_RAT
Stromelysin 3	24.24	COGY_HUMAN
Gelatinase A Gelatinase B		GOG2_HUMAN COG9_HUMAN
Family M11: Autolysin		eptidases with HEXXH zinc-binding motif)
Autolysin	24.38	
Family M12: Astacin		eptidases with HEXXH zinc-binding motif)
Metalloendopeptidase (Caenorhabditi	•	(M75746)
Blastula protease-10 (Paracentrotus)	-	(X56224)
Astacin	24.21	ASTA_ASTFL
tolloid gene product (Drosophila)	-	(M76976)
UVS.2 protein (Xenopus)	-	[23]
Ruberlysin	24.48	HRT2_CRORU
Atrolysin c		HRTD_CROAT
Trimerelysin II	24.53	HR2_TRIFL
HR2a-endopeptidase (habu snake)	-	HR2A_TRIFL
HR1B-endopeptidase (habu snake)	•	HR1B_TRIFL
Haemorrhagic factor LHFII (bushmaster snake)	-	HRL2_LACMU
Meprin A	24.18	(M74897)

PABA-peptide hydrolase 24.18 (M82962) Bone morphogenetic protein 1 BMP1 HUMAN Family M13: Neprilysin (Clan MA: Peptidases with HEXXH zinc-binding motif) Neprilysin 24.11 NEP_HUMAN, NEP_RABIT, NEP_RAT Kell blood group protein **KELL HUMAN** Family M14: Carboxypeptidase A (HXXE zinc-binding motif) Zinc-carboxypeptidase (Streptomyces) **CBPS STRGR** Carboxypeptidase T (Thermoactinomyces) (X56901) Carboxypeptidase B 17.2 CBPB ASTFL, CBPB BOVIN, CBPB RAT, (M75106) Carboxypeptidase A 17.1 CBPA_BOVIN, CBPC_HUMAN, CBPC_MOUSE, CBP1_RAT, CBP2_RAT, (A25833) Lysine carboxypeptidase 17.3 **CBPN HUMAN** Carboxypeptidase H 17.10 CBPH BOVIN, CBPH HUMAN, CBPH_RAT, (X61232), [24] Carboxypeptidase M 17.12 CBPM HUMAN Family M15: Muramoyl-pentapeptide carboxypeptidase (HXH zinc-binding motif) Muramoyl-pentapeptide carboxypeptidase 17.8 **CBPM STRGR** (HXXEH zinc-binding motif) Family M16: Pitrilysin Pitrilysin 99.44 PTR ECOLI pqqF gene product (Klebsiella) (X58778) Insulinase 99.45 IDE DROME, IDE HUMAN Mitochondrial processing peptidase 99.41 MPP1 NEUCR, MPP1 YEAST, MPP1 RAT MPP2 NEUCR, MPP2 YEAST Processing enhancing protein Ubiquinol-cytochrome c reductase 1.6.99.3 UCR1_YEAST, UCR2_YEAST, UCR2_HUMAN core proteins 1 and 2 Family M17: Leucyl aminopeptidase (Peptidases binding two zinc atoms: Lys, Glu, Asp, Asp, Glu) Leucyl aminopeptidase 11.1 AMPL BOVIN, (X63444) Aminopeptidase A (Escherichia coli) AMPA ECOLI, (M68966) Family M18: Aminopeptidase yscl Aminopeptidase yscl (Saccharomyces) AMPL_YEAST, LAP4_YEAST Family M19: Membrane dipeptidase Membrane dipeptidase 13.19 MDP4_HUMAN, MDP4_PIG Open reading frame X product (Klebsiella) (X58778)Gene R product (Acinetobacter) (X06452)Family M20: Carboxypeptidase G2 Carboxypeptidase G2 (Pseudomonas) CBPG PSES6 Peptidase T (Salmonella) (M62725)Family M21: Gly-X carboxypeptidase Gly-X carboxypeptidase (Saccharomyces) 17.4 (X57316)Family M22: A1 Glycoprotease A1 Glycoprotease (Pasteurella) (M62364)OrfX (Escherichia coli) YRUX ECOLI OrfX (Salmonella) (M14427) Family M23: 8-lytic endopeptidase B-Lytic endopeptidase 24.32 PRLB LYSEN, (M60896) LasA protein (Pseudomonas) LASA PSEAE Family M24: Methionyl aminopeptidase Methionyl aminopeptidase 11.18 AMPM_BACSU, AMPM_ECOLI, AMPM_SALTY Aminopeptidase P (Escherichia coli) AMPP ECOLI X-Pro dipeptidase 13.9 PEPQ_ECOLI, PEPD_HUMAN Family M25: X-His-dipeptidase X-His dipeptidase 13.3 PEPD_ECOLI

PEPTIDASES OF UNKNOWN CATALYTIC TYPE

Family U1: Aminopeptidase T

Aminopeptidase T (Thermus)

AMPT THEAQ

Table 1 (contd.) Family U2: Aminopeptidase IAP Alkaline phosphatase isozyme conversion IAP ECOLI protein (Escherichia coli) Family U3: Spore endopeptidase, Bacillus megaterium Spore endopeptidase (Bacillus megaterium) -(M55262) Family U4: Sporulation sigma factor processing peptidase Sporulation sigma factor processing SP2G BACSU peptidase (Bacillus subtilis) Family U5: Tail-specific protease Tail-specific protease (Esherichia coli) (M75634)Family U6: Murein endopeptidase Penicillin-insensitive murein endopeptidase MEPA ECOLI (Esherichia coli) Family U7: Endopeptidase IV Endopeptidase IV (Escherichia coli) SPPA_ECOLI, LICA_HAEIN sohB gene product (E. coli) (M73320) Minor capsid protein precursor C VCAC_LAMBD (bacteriophage lambda) Family U8: Bacteriophage endopeptidase ENPP BPPA2, ENPP BPP22, ENPP_BPT3, ENPP_BPT7, Endopeptidase (bacteriophage) ENPP_LAMBD Family U9: Prohead endopeptidase Prohead endopeptidase (bacteriophage T4) PCPP_BPT4 Family U10: Leader peptidase Leader peptidase 99.36 LEP ECOLI, LEP SALTY, (X56466), (Z11847) Mitochondrial inner membrane peptidase 1 [25] (Saccharomyces) Family U11: Premurein leader peptidase 99.35 LPSA ECOLI, LPSA ENTAE, LPSA PSEFL, (M83994), Premurein leader peptidase (M84707) Family U12: Prepilin leader peptidase Prepilin leader peptidase (Vibrio) (M74708) COMC BACSU Late competence protein (Bacillus) xpcA protein (Pseudomonas) PILD PSEAE Pullulanase secretion protein (Klebsiella) PULO KLEPN Family U13: Leader peptidase component 3-4 Leader peptidase 21 kDa subunit (dog) 99.36 SPC3 CANFA Leader peptidase 18 kD subunit (dog) 99.36 SPC4_CANFA Leader peptidase (sec11) (Saccharomyces) 99.36 Family U14: Leader peptidase component 2 Leader peptidase 22-23 kDa subunit (dog) 99.36 SPC2 CANFA Microsomal leader peptidase (chicken) 99.36 (X60795) Leader peptidase (Drosophila) 99.36 (M32022)Family U15: Multicatalytic endopeptidase complex Multicatalytic endopeptidase subunits 99.46 PRCA THEAC, (M83674), (J05358), PRC1_YEAST, PRC7_YEAST, (M63641), PRCD_YEAST, PRCB_YEAST, PRCX_YEAST,

PRCZ_YEAST, PR28_DROME, PR29_DROME, PR35_DROME, PRC3 XENLA, (X62709), PRC2_RAT, PRC5_RAT, PRC3_RAT, PRC8_RAT, PRC9_RAT, (M64992), (D00760), (D00761), (D00762),

(D00763), (D10729), (X64449)

SCL1 suppressor protein (Saccharomyces) 99.46 SCL1_YEAST

Family U16: Thermopsin

99.43 THPS_SULAC Thermopsin

Family U17: Ubiquitin-specific processing protease

Ubiquitin-specific processing protease I (M63484)

(Saccharomyces)

Family U18: Scytalidiapepsin

Scytalidiapepsin B 23.32 PRTB_SCYLI Scytalidiapepsin (Aspergillus) - PRTA ASPNG

Family U19: Pestivirus endopeptidase

Endopeptidase (cattle viral diarrhoea virus) - (M37795), (M62430)

Family U20: y-D-Glutarnyl-L-diamino acid endopeptidase II

γ-D-glutamyl-L-diamino acid endopeptidase - (X64809)

II (Bacillus sphaericus)

Family U21: Potyvirus endopeptidase 3

35 kDa endopeptidase - POLG_PPVD, POLG_PPVYN, POLG_TEV,

POLG TVMV

chains of the enzymically active members of family S1 is His, Asp, Ser. The same order of residues is seen in family S2 (α -lytic endopeptidase) and family S3 (togavirus endopeptidase), and members of these families also have tertiary structures similar to that of chymotrypsin [28,29]. This strongly suggests that they share a common evolutionary origin, despite the differences of sequence, and accordingly we group families S1, S2 and S3 in a single clan (SA). The evidence is less complete for families S4, S5, S6 and S7, but there are indications that these also may belong in this clan [30–33].

The enzymes of the subtilisin (S8) family have a different order of catalytic-site residues from chymotrypsin, namely Asp, His, Ser, and also have different tertiary structures. It is therefore quite clear that the family represents a separate evolutionary line of serine peptidases [34]. The family contains an exopeptidase (tripeptidyl peptidase II) as well as endopeptidases with various specificities. Most of the microbial members of the family have specificities somewhat like that of chymotrypsin, but the eukaryote enzymes include the proprotein convertases such as kexin and furin, which are specific for substrates containing paired basic residues [35].

We consider that the family of prolyl oligopeptidase (S9) reflects a further distinct evolutionary line of serine peptidases. In this family there is again a different order of catalytic residues, Ser⁵⁵⁴ and His⁸⁸⁰ being known for pig prolyl oligopeptidase [36]. We have suggested that if an Asp residue completes a catalytic triad, Asp⁵²⁹ is the most likely [37]. There is evidence that prolyl oligopeptidase differs significantly in catalytic mechanism from the enzymes of families S1 and S8 [38,39]. The family contains two endopeptidases with the restricted specificity for substrate size that makes them oligopeptidases [37]; one of these cleaves prolyl bonds, whereas the other acts on bonds with a basic residue in the P1 position. The family also contains a dipeptidyl peptidase and an omega peptidase [37].

The serine-type carboxypeptidases form family S10, in which the order of catalytic residues is Ser, Asp, His. The tertiary structure of these enzymes is unlike those known for other families, and they are unusual amongst serine-type hydrolases in being maximally active at about pH 5 [40]. There are similarities between the structures of the active sites of these enzymes and those lipases [40] and acetylcholinesterases.

There are three distinct families of serine-type D-Ala-D-Ala carboxypeptidases, S11, S12 and S13, all confined to bacteria. Their members are similar in catalytic mechanism and three-dimensional structure [13,41,42], and thus are grouped in a single clan (SB), which also contains other penicillin-binding proteins. Family S12 also contains a D-aminopeptidase, the only serine-type aminopeptidase reported to date.

The Clp endopeptidase is one of the ATP-dependent proteolytic enzymes of *Escherichia coli* and contains two subunits, ClpP and ClpA, ClpP being responsible for the peptidase activity. The active-site serine (Ser¹¹¹) and histidine (His¹³⁶) of ClpP are known, but no aspartic acid that might form the third member of a catalytic triad has been identified. Other members of this family (S14) occur in plant chloroplasts, which may reflect their endosymbiont origins. We report here that the 5' end of potatoleaf-roll-luteovirus genomic RNA, which has been described as a non-coding region [43], also is homologous with ClpP.

The active-site serine of endopeptidase La (S16) has been determined as Ser⁶⁷⁹ [44], but otherwise no catalytic-site residues have been identified in families S15–S22. Family S15 contains a dipeptidyl-peptidase specific for the cleavage of Xaa-Pro + bonds that is unrelated to the enzyme with similar specificity in family S9

Cysteine peptidases

In addition to many endopeptidases, the papain family (C1) contains an exopeptidase, dipeptidyl-peptidase I, and proteins that lack peptidase activity, in *Plasmodium* and soya bean. Unusually, the papain family contains a sequence from a baculovirus genome, which may have been acquired from a host [45]. With this exception, proteins of the papain family have been found only in eukaryotes.

The calpains (family C2) are heterodimeric enzymes, the larger (80 kDa) subunit containing the proteolytic domain and also calcium-binding, E-F-hand structures similar to those found in other proteins. No active-site histidine has yet been positively identified, but Cys¹⁰⁸ and His²⁶⁵ (in chicken calpain) occur in sequences that show some similarity to those around catalytic residues in the papain family [46], and accordingly the families of papain and calpain form a clan (CA). The homologous sol protein from *Drosophila* has a distinctive structure, being a much larger protein with zinc fingers but no calcium-binding sites [47].

Families C3—C9 comprise viral enzymes. We group families C3—C5 in a clan (CB) in which the order of catalytic residues is His, Cys. It is possible that these enzymes are related to those of the chymotrypsin family (S1), with interconversion of the essential serine and cysteine residues [48,49]. Such a relationship would represent the only known homology across catalytic types. In most members of the clan an aspartic residue is thought to form the third member of a catalytic triad, but in the picornains 3C, the essential Asp is replaced by Glu [50,51].

Essential cysteine and histidine residues occur in the order Cys, His in families C6-C10, but families C6-C14 cannot as yet be assigned to clans.

Aspartic peptidases

At present, it seems that all of the aspartic peptidases are endopeptidases, and the great majority of these are members of the pepsin (A1) family, which have been found only in eukaryotic organisms. The peptidases of this family have bilobed molecules resulting from gene duplication [52]. In contrast, the viral retropepsins (family A2) have a monomeric structure in which each molecule contains only half of the functional catalytic site and dimerization is needed to form the active enzyme. We place families A1 and A2 in a clan (AA) on the basis of similarities in tertiary structure [53] and sensitivity to inhibition by pepstatin [54].

The catalytic residues of acid proteinases in two other families have not been identified, so it is not possible at this stage to say whether these pepstatin-resistant enzymes also are aspartic endopeptidases. These are the families of thermopsin (U16) and scytalidopepsin (U18). For scytalidopepsin B it was suggested that Glu⁵³ is involved in catalysis [55], but this is not conserved in the second member of the family [56].

Metallopeptidases

The structures of metallopeptidases are exceptionally diverse, and we recognize 25 families. The majority of the enzymes contain zinc, and for several of them the residues involved in binding the zinc have been identified by X-ray crystallography.

Of the families of metallopeptidases, 13 contain the sequence HEXXH, which is known or suspected to provide two of the three ligands for the zinc atom. These are the families of alanyl aminopeptidase (M1), peptidyl-dipeptidase A (M2), thimet oligopeptidase (M3), thermolysin (M4), mycolysin (M5), immune inhibitor A (M6), Streptomyces protease (M7), leishmanolysin (M8), Vibrio collagenase (M9), interstitial collagenase (M10), autolysin (M11), astacin (M12), and neprilysin (M13). With some reservations, we group these in a clan (MA). In each of these families, the HEXXH sequence occurs in a nine-residue consensus sequence, bXHEbbHbc, in which b is an uncharged residue, c is hydrophobic, and X can be any amino acid. The third ligand of the zinc atom is glutamic acid in the families of alanyl aminopeptidase [57], thermolysin [58] and neprilysin [59], but is histidine in the astacin family [60], and presumably also in those of autolysin and interstitial collagenase, since the histidine is conserved. Family M1 contains aminopeptidases and an ether hydrolase (EC 3.3.2.6), and family M3 contains peptidyl-dipeptidases and oligopeptidases. The large thermolysin family (M4) appears to be confined to bacteria, whereas members of the astacin family (M12) have been found only in animals.

Families M10 and M12 each contain members of only 200–300 residues (astacin and matrilysin respectively), but also much larger chimaeric proteins. Our inclusion of the snake-venom metalloendopeptidases within the astacin family is based on statistically significant sequence relationships with the endopeptidase domain of human bone morphogenetic factor 1 for ruberlysin and atrolysin C.

Four further families of zinc metallopeptidases exhibit distinctive modes of binding of the metal. In the family of carboxypeptidase A (M14) the short-spaced ligands of zinc (in the terminology of Vallee [61]) are histidine and glutamic acid in the sequence HXXE, whereas they are two histidine residues in the sequence HXH in the family of muramoylpentapeptide carboxypeptidase (M15). In both families, the third ligand is histidine. Even more distinctive is the set of lysine, two aspartic and two glutamic residues that bind a pair of zinc atoms at the active site in the leucyl aminopeptidase (M17) family. The enzymes of the pitrilysin family (M16) are thought to bind zinc

at an HXXEH sequence [62], and in insulinase the sequence is HFCEH, which may account for the thiol-dependence of the enzyme. In pitrilysin, which is not thiol-dependent, this sequence is HYLEH. Several members of the family lack the HXXEH consensus altogether, and presumably are inactive; these include 'mitochondrial processing peptidase' and the reductase subunits. It has been suggested that the activity formerly attributed to mitochondrial processing peptidase is due to the associated processing-enhancing protein [62]. We report here that the kpqqF gene of Klebsiella pneumoniae [63] shows homology with members of the pitrilysin family; this sequence, which does contain the HXXEH consensus, seems too dissimilar to that of pitrilysin for the Klebsiella protein to be simply the species variant.

The groups responsible for zinc-binding are unknown for the other families of metallopeptidases (M18–M25). Family M24 contains aminopeptidases and a dipeptidase.

Unknown catalytic type

There are 21 families for which the catalytic type remains to be determined. The family of endopeptidase IV (U7) contains viral and bacterial enzymes, the only such family we know of, but the virus is a bacteriophage, so presumably may have acquired the gene from a host.

The leader peptidases form the five families U10-U14. The bacterial leader peptidases (U10-U12) are not homologous with eukaryote microsomal leader peptidases, but the bacterial leader peptidase of family U10 is related to the eukaryotic mitochondrial leader peptidase, which may reflect the endosymbiont origin of mitochondria. The eukaryotic microsomal leader peptidases are multisubunit proteins, and it is not clear which components are directly responsible for peptidase activity. The component subunits form at least two families, U13 and U14. We have noticed that the 3'-terminal portion of a sequence that includes the sexspecific gene msP316 from Drosophila [64] is homologous with the glycoprotein component 2 of microsomal leader peptidase.

The molecules of multicatalytic endopeptidase complex (U15) contain two or more kinds of subunit, which are nevertheless homologous. It has yet to be established which of the three or more distinct peptidase activities of the enzyme are attributable to which subunits [65].

Conclusions

Until recently it has seemed that the vast majority of endopeptidases belonged to just a few evolutionary families, those of chymotrypsin, subtilisin, papain, pepsin and thermolysin [66]. By analogy, it might have been expected that the exopeptidases also would prove to belong to just a few families, which would have been separate from those of the endopeptidases. It would thus have been natural to assume that modern peptidases reflect a small number of independent evolutionary origins, perhaps a dozen or so. However, our analysis of the hundreds of amino acid sequences now available for peptidases points to different conclusions. Using rigorous standards for relatedness, we have had to recognize no fewer than 84 distinct families of peptidases. A number of families show signs of distant relationship to others, and accordingly have been placed in clans. Even so, we have 60 groups of sequences among which we can see no relationship, and there may therefore have been this many separate evolutionary origins of peptidases.

The origins of many of the modern families of peptidases clearly were very early, since members are present in modern prokaryotic micro-organisms. Families that appear in prokarvotes, but also are found in eukarvotic organisms, are those of chymotrypsin (S1), subtilisin (S8), prolyl oligopeptidase (S9), ClpP (S14), alanyl aminopeptidase (M1), thimet oligopeptidase (M3), interstitial collagenase (M10), carboxypeptidase A (M14), pitrilysin (M16), leucyl aminopeptidase (M17), methionyl aminopeptidase (M24), and multicatalytic endopeptidase complex (U15). No examples are known for cysteine- or aspartic-type peptidases. Three large families of endopeptidases have few or no known examples among prokaryotes, but underwent major expansion in the eukaryotes; these are the families of chymotrypsin (S1), papain (C1) and pepsin (A1). Possibly they developed in connection with the acquisition of the capacity for endocytosis by eukaryotic cells, to function in membrane-limited organelles, sometimes at acidic pH. All the known viral peptidases are endopeptidases, and the great majority of these show no relationship to the peptidases of other organisms, although exceptions are listed in families S3, C1, A2 and U7 of Table 1. No viral metallopeptidase has been described.

We have seen striking examples of the amount of divergent evolution that can occur in a family of peptidases. Six of the families contain proteins that are not peptidases (S1, S12, S14, C1. M1 and M16). Also, four families contain both exopeptidases and endopeptidases (S8, S9, C1 and M3). Family S12 contains both aminopeptidases and carboxypeptidases, and family M24 contains aminopeptidases and a dipeptidase. In most of the families there are peptidases differing greatly in specificity for amino acids around the scissile bond. Conversely, there has been convergence such that a number of peptidase specificities are exhibited by enzymes of more than one family; examples would be the activities of glutamyl endopeptidase (S2 and S4), Xaa-Pro + dipeptidyl peptidase (S9 and S15), peptidyl-dipeptidase (M2 and M3), carboxypeptidase specific for basic residues (S10 and M14) and D-Ala-D-Ala carboxypeptidases (S11, S12, S13 and M15).

Having constructed a classification of peptidases that is based on structural and evolutionary relationships, we have naturally considered whether this has anything to add to the currently accepted methods of classification by reaction catalysed and by catalytic mechanism. The evolutionary scheme is clearly not compatible with classification of the enzymes by the reactions they catalyse, since many families contain enzymes with quite different kinds of peptidase activities, and some specificities are found in several families. The evolutionary scheme does fit well within the system of classification by catalytic type, however, as can be seen in Table 1, and it tends to bring together the enzymes that resemble each other most closely in structure and catalytic mechanism. We therefore suggest that it deserves serious consideration for use in future schemes for the classification of these enzymes, as an extension of the classification by catalytic type.

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