

Evolution of Disintegrin Cysteine-Rich and Mammalian Matrix-Degrading Metalloproteinases: Gene Duplication and Divergence of a Common Ancestor Rather Than Convergent Evolution

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Abstract. The evolution of the Metalloproteinase Disintegrin Cysteine-rich (MDC) gene family and that of the mammalian Matrix-degrading Metalloproteinases (MMPs) are compared. The alignment of snake venom and mammalian MDC and MMP precursor sequences generated a phylogenetic tree that grouped these proteins mainly according to their function. Based on this observation, a common ancestry is suggested for mammalian and snake venom MDCs; it is also possible that gene duplication of the already-assembled domain structure, followed by divergence of the copies, may have significantly contributed to the evolution of the functionally diverse MDC proteins. The data also suggest that the structural resemblance of the zinc-binding motif of venom MDCs and MMPs may best be explained by common ancestry and conservation of the proteolytic motifs during the divergence of the proteins rather than through convergent evolution.

Key words: Metalloproteinase — Disintegrin — Evolution — Venom — Phylogeny — Gene duplication

Introduction

Matrix-degrading metalloproteinases (MMPs) are a group of related enzymes involved in extracellular deg-

radation of the protein components of the connective tissue. They are involved in the tissue remodeling which occurs during embryogenesis, development, and wound healing. They may also be involved in certain diseases such as arthritis, periodontitis, and tumor metastasis. The proteolytic activity of these enzymes is dependent on the presence of a consensus zinc-binding motif in their catalytic domain (Blundell 1994). The same zinc-binding motif is present in certain toxins of viper venoms which induce hemorrhage in the prey or victims of snake bite. They act on similar substrates to MMPs present in the basement membrane of the endothelium (Bjarnason and Fox 1994). Both snake venom and matrix metalloproteinases are classified in the Astacin family of zinc metalloproteinases based on the structure of the catalytic domain and zinc-binding motif; however, the sequence and structure of the surrounding areas are variable (Blundell 1994).

Other components of viper venoms that interfere with hemostasis are the disintegrins. These comprise a family of RGD-containing peptides with high affinity for the platelet glycoprotein GP IIb/IIIa integrin receptor (Gould et al. 1990). They prevent platelet aggregation by inhibiting the binding of fibrinogen and von Willebrand factor to platelets (Huang et al. 1989).

Interestingly, like MMPs both hemorrhagic toxins and disintegrins are coded by cDNAs that predict zymogen molecules containing a large precursor pro-peptide domain followed by a metalloproteinase domain. These precursor proteins differ from MMPs in having a car-

boxy-terminal disintegrin domain of varying length which replaces the hemopexin domain present in MMPs (Paine et al. 1992). The pro-peptide domain is apparently involved in the inactivation of the proteolytic activity prior to the secretion of the enzymes by venom gland cells. This process is thought to occur through a cysteine switch mechanism, as also proposed for the MMPs (Woessner 1991), and is correlated with the sequence PRCGV in MMPs, and with PKMCGV in the snake toxins. The metalloproteinase domain has a zinc-binding sequence, HEXXHXXGXXH, characteristic of the Astacin family described above (Blundell 1994). The disintegrin domain is capable of binding to the platelet integrins through the RGD motif that is classically represented in venom disintegrins (Huang et al. 1989). Large hemorrhagins contain a region of sequence similarity with the disintegrins, with a substitution of the RGD motif with E/DCD at the same position, and a cysteine-rich extension of the carboxy-terminus (Paine et al. 1994).

It has recently become apparent that a number of cysteine-rich proteins with similar structural organizations occur in the mammalian male reproductive tract. The best characterized of these are Fertilin (formerly PH-30), a guinea pig sperm surface protein involved in sperm-egg fusion (Blobel et al. 1992), and the androgen-regulated epididymal apical protein-I, EAP-I (Perry et al. 1992). These proteins include a carboxy-terminal extension linked to a venom-like long hemorrhagin sequence comprising an epidermal growth factor (EGF) repeat, a transmembrane domain, and a cytoplasmic tail (Weskamp and Blobel 1994). Representatives of this protein family were also detected in other tissues, including the MS-2 antigen present on the surface of certain lineages of macrophages (Yoshida et al. 1990) and the protein predicted by the MDC gene, which has been associated with tumor suppression (Emi et al. 1993).

All the mammalian proteins possess a non-RGD disintegrin domain, characteristic of the large hemorrhagins. The zinc-binding motif is present only in EAPs, Fertilin α , and the monocyte surface antigen metalloproteinase domains, and is not yet fully correlated with the function of these proteins. The cysteine switch motif is absent in all the mammalian pro-domains. These cysteine-rich mammalian proteins plus the snake venom hemorrhagins and disintegrins make up the metalloproteinase, disintegrin, cysteine-rich (MDC) family (Fig. 1).

The evolutionary aspects of the complex MDC gene family are still poorly understood. Paine et al. (1994) suggested that accelerated evolution may apply to the snake toxins, specifically with regard to their high variability occurring mainly in the metalloproteinase domain. They also suggested that the generation of the RGD disintegrins might be associated with a deletion in the DNA region that codes for the cysteine-rich domain in long hemorrhagins. Recently, Wolfsberg et al. (1993)

suggested, based on phylogenetic trees, that the individual domains of the MDCs may have been assembled before the divergence of the members of the family. This last report does not, however, adequately explain the close functional similarity between the venom metalloproteinases and MMPs as opposed to the mammalian MDCs. We now report an analysis of MMP and MDC sequences undertaken precisely to address this issue. The aim was to study the evolutionary aspects of the MDCs in comparison to MMPs in order to provide new insights into the evolution of these gene families.

Materials and Methods

The sequences used in this paper were obtained by scanning GenBank, Swiss-Prot, and PIR databases using the BLAST program (Altschul et al. 1990). The sequences of the following proteins, predicted from cDNA sequences, were used: Human Matrilysin (gb L22524), collagenase 3 (gb X75308), and tumor suppressor gene product MDC (gb D17390); mouse cyritestin (pir S18968) and monocyte surface antigen (pir A60385); rat Stromelysin-2, MMP-10 (sp P07152), and epididymal protein, EAP (pir S28259); guinea pig Fertilin α (gb Z11719) and β (gb Z11720); monkey epididymal protein, EAP (pir S28258), tMDC-I (gb X76637) and II (gb X77619); Rhodostomin (pir S33792) from *Calloselasma rhodostoma* snake venom; Atrolysin E (pir A43296), B (pir S41608), C (pir S41609), and Catrocollastatin (gb U21003) from *Crotalus atrox* venom; Trigramin (pir A30065) from *Trimeresurus gramineus* venom; Jararhagin (pir S24789) from *Bothrops jararaca* venom; Trimucin (pir S43125) from *Trimeresurus mucrosquamatus* venom; a hemorrhagin (gb U18234) from *Agkistrodon contortrix* venom; Halystatin (gb D28870) from *Agkistrodon halys* venom; EcH I (gb X78970), and EcH II (gb X78971) from *Echis pyramidum leakeyi* venom. Because of recent taxonomic reorganisation, it is likely that *Tr. gramineus* and *A. halys* referred to here now correspond to *Tr. stejnegeri* and *A. blomhoffi breviceaudus*, respectively (D.A. Warrell, personal communication).

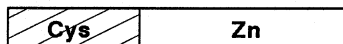
Percentage similarity and frequency of codon usage have been calculated using the GCG software, bestfit, and codon usage program (Genetics Computer Group 1991). Phylogenies were constructed using the Clustal V program (Higgins et al. 1992). Progressive alignments used the multiple alignment algorithms described by Higgins and Sharp (1989), with a fixed gap penalty of 10 and the Dayhoff PAM 250 protein weight matrix (Dayhoff et al. 1978). Alignments were finally refined by eye and differ very slightly from those generated by computer. The phylogenetic trees according to distances were generated from the above alignments using a neighbor-joining method (Saitou and Nei 1987) and rooted using the Thermolysin sequence (pir M21663) from *Bacillus stearothermophilus*. The degree of error was calculated for each branch by bootstrapping (Felsenstein 1985) and the values are shown in the tree. Trees were constructed using the whole protein sequence, as well as for the pro-protein, metalloproteinase, and disintegrin domains. In all cases phylogenies constructed with sites where gaps occurred in any one sequence of the alignment being deleted and not deleted were compared.

Results and Discussion

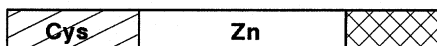
Sequence Similarity Between MDCs and MMPs Appears to Be Restricted to the Functional Motifs

This study has been carried out using the precursor sequences predicted by the cDNAs coding for three MMPs,

Mammalian MMPs



Matrilysin (MAT)

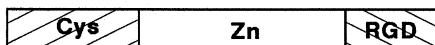


Human Collagenase-3 (COL)

Stromelysin-2 (STR)

Snake venom MDCs

Disintegrins



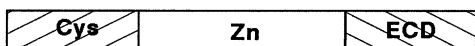
Rhodostomin (RHO)

Trigramin (TRG)

Trimucin (TRM)

Halystatin (HAL)

Long-chain haemorrhagins



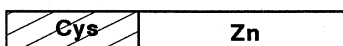
Jararhagin (JAR)

Catocollastatin (CAT)

EcH-I (ECHI)

EcH-II (ECHII)

Short-chain haemorrhagins



Atrolysin E (ATRE)

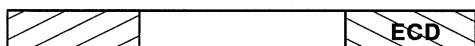
Atrolysin B (ATRB)

Atrolysin C (ATRC)

Agkistrodon Haemorrhagin (AGH)

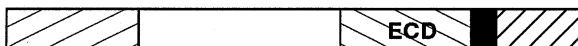
Mammalian MDCs

Soluble



Tumour Suppressor gene (TSG)

Without Zn motif



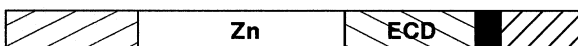
Cyritestin (CYR)

tMDC-I (MDCI)

tMDC-II (MDCII)

Fertilin β (FER β)

With Zn motif



Monocyte Surface Antigen (MS2)

Rat Epididymal Protein (EAPR)

Monkey Epididymal Protein (EAPM)

Fertilin α (FER α)

Fig. 1. Schematic representation of mammalian matrix-degrading metalloproteinases (MMPs) and snake venom and mammalian proteins belonging to the metalloproteinase-like, disintegrin-like, cysteine-rich (MDC) gene family: (▨) Pro-peptide domain; (□) metalloproteinase domain; (▩) disintegrin domain; (▧) hemopexin; (■) EGF repeats; (▨) transmembrane domain. The presence of the cysteine-switch and zinc-binding motifs is represented by Cys and Zn, respectively. RGD and ECD represent the potential integrin binding peptides in the disintegrin domain.

nine mammalian MDCs, and 12 snake venom MDCs. The criteria used to select these sequences was based on a wider representation of the functional diversity among these proteins and also a number of representatives from different genera/species of snakes. Figure 2 shows two representative fragments extracted from the alignment of the whole precursor sequences, containing, respectively, the cysteine-switch and zinc-binding motifs. The similarity between snake and mammalian MDCs is recognized by the conservation of most cysteines and other

residues throughout all the MDC sequences (Fig. 2, stars). Some conserved positions among snake toxins are also shared with few, but not all, mammalian MDCs (Fig. 2, dots). Therefore, MDCs may be considered a family of proteins with high sequence similarity. However, the comparison of MDCs with MMPs reveals little sequence similarity (below 45%), this being confined to the zinc-binding and cysteine switch motifs, the latter being observed only in MMPs and snake venom MDCs. These observations also apply to the conformation of

Table 1. Frequency of codon usage for the essential residues for the catalytic activity in cDNAs regions coding for the zinc-binding motif (Zn) and the whole precursor molecules (WP) of MMPs and proteolytic MDCs^a

		His			Glu			Gly	
		WP	Zn		WP	Zn		WP	Zn
AGH	cat	0.87	1.0	gag	0.5	1.0	ggg	0.16	—
	cac	0.13	—	gaa	0.5	—	gga	0.44	—
							ggt	0.28	—
							ggc	0.12	1.0
JAR	cat	0.68	1.0	gag	0.36	1.0	ggg	0.09	—
	cac	0.32	—	gaa	0.64	—	gga	0.51	—
							ggt	0.19	—
							ggc	0.21	1.0
TRG	cat	0.82	1.0	gag	0.33	1.0	ggg	0.13	—
	cac	0.18	—	gaa	0.67	—	gga	0.48	—
							ggt	0.16	—
							ggc	0.23	1.0
ECHI	cat	0.69	1.0	gag	0.38	1.0	ggg	0.13	—
	cac	0.31	—	gaa	0.62	—	gga	0.49	—
							ggt	0.15	—
							ggc	0.23	1.0
EAPM	cat	0.69	1.0	gag	0.32	—	ggg	0.16	1.0
	cac	0.31	—	gaa	0.68	—	gga	0.56	—
							ggt	0.14	—
							ggc	0.14	—
FER α	cat	0.27	0.33	gag	0.46	1.0	ggg	0.24	—
	cac	0.73	0.67	gaa	0.54	—	gga	0.31	—
							ggt	0.27	1.0
							ggc	0.18	—
COL	cat	0.67	0.33	gag	0.41	1.0	ggg	0.14	—
	cac	0.33	0.67	gaa	0.59	—	gga	0.29	—
							ggt	0.31	1.0
							ggc	0.26	—
MAT	cat	0.60	0.33	gag	0.22	—	ggg	0.19	—
	cac	0.40	0.67	gaa	0.78	1.0	gga	0.33	—
							ggt	0.14	1.0
							ggc	0.33	—

^a Abbreviations as in Fig. 1

these proteins. A comparison of the crystal structure of Adamalysin II, a snake venom metalloproteinase, with the crayfish Astacin reveals some topologically equivalent residues (Gomis-Ruth et al. 1993). However, only the active site environment, comprising the zinc-binding consensus region and the active site basement, appears to exhibit identical conformation (Table 1).

The Evolutionary History of MDCs

Evolutionary trees have been constructed using the alignments of the complete precursor proteins (Fig. 3) or the regions comprising the distinct domains of the sequences described above (data not shown). The tree constructed using the complete sequences (Fig. 3) shows two primary divergent groups comprising the MMPs and MDCs, respectively. The MDC cluster is apparently monophyletic and the sequences are distributed mainly according to their function. The first group contains the sperm proteins related to Fertilin, and the second group includes the EAPs. The tumor suppressor gene and the monocyte surface antigen are located in the first and second groups,

respectively. Snake venom sequences are also distributed according to function, the first group representing the long-chain hemorrhagic toxins, the second clustering the RGD-disintegrins, and the third enclosing the short hemorrhagins. Clearly, some of these sequences are quite dissimilar, and it is important to note that the clustering of the different molecules in the tree remains essentially the same even when all sites containing alignment gaps are removed from the data set during tree construction. In this situation, the analysis to some extent favors the combined pro- and metalloproteinase domains of the molecules.

Trees corresponding to the pro-, metalloproteinase or disintegrin domains were also constructed using the alignments corresponding to the relevant fragment of each toxin. The same clustering characteristics are suggested by the trees generated using whole sequences (Fig. 3) or independent domains (data not shown). The only exceptions were the long-chain hemorrhagins, Jararhagin and Catrocollastatin, from pit vipers, which appear to cluster preferentially with other pit viper se-

...	156			248...
MAT	GKLSFYIMEIMQKPRCGV-----PDVAEYSLMP--NSPKW-----HSRIVTYRIVSYTSDLPRIIVVDQIV-----			
STR	GKLDNTVEMMHKPRCGV-----PDVGGFTFP--GSPKW-----RKNHISYRIVNYTLDLPRESVDSAI-----			
COL	GKLDNTVDMVKKPRCGV-----PDVGEYVFP--RTLKW-----SKMNLTYRIVNYTLDPMTHSEVEKAF-----			
TSG	GKLRGNPHSFAALSTCQGLHGVFSDGNLTYVEPQEVAGPWGAPQGPLPHLYRTPLLPDLCREPGCLFAVPAQASAPPNRPRLRRKRQVR			
FER α	GYIEGASSFSVVSACSLRGLILKENTSYGIEPLLSQR-----FEHVLTYT--MARQAPVSCR--ASAKDSQAVSTSWQOQSRKPHSVQ			
FER β	GHIIEGFPSTLASISTCAGLRGLLQFETVSYGIEPLKSSIG-----FEHVIYP--V-KHDNEKSO-YLKKSINV--K--NVVYKI-----			
MDC1	GYVADIPKSAVTLRSTCGLRGLLQDNIYSYIEPLESSPT-----YEHVVYR--I-KNDAIGHF-SFQENYVP--AQYIDQSYRILVKS			
MDC1I	GYVAGIPNSLVTLSVCSGLRGTMLKNIYSYIEPEMAVSG-----FIHKIYE--E-KFADTNI--LLEENDT--YSWFNSEYQVRKSS			
CYR	GHAAEIPVSTVTLSTCGLRGLLQLENITGYIEPLESSAT-----FEHILYE--I-KNNKIDYS-PLKENFAN--SEQESQSYRILVVKP			
MS2	GHVEGYEGSAASISTCAGLRGFFRVGTVHLIEPLDADDE-----GQHAMYQAKHLQOQAGT--CGVKDNTNLNDLGP--RALEIYRAQ			
EAPR	GSIIHEFDSAAISISTCAGLRGFFRVNDQRYLIEPLKYSDE-----GDHLVFKY-NVKAPYATNS-CBGLNFTKKSSTLIDAKIIE-----E			
EAPM	GSIVHEYDSAAISISTCAGLRGFFRVNDQRYLIEPLKYSDE-----GEHLVFKY-NPRVPYVANY-S-CTELNFTRKTVPDGTESRG----D			
EChI	GRIQNDADSTASMSACNLKGYFMLRGETYLIIEPLKIPDS-----EAHAVYKYENVEKEDEAPKM-CGVTTQTNWES-DELKASQLVATS			
EChII	GRVQNDADHSSASISACNLKGFHKLQGETYFIEPLKIPDS-----EAHAVYKYENIEKEDQAPKM-CGVTHTNWESDEPIKEASRLVASS			
JAR	GRIENDADSTASISACNLKGYFKLQRETYFIEPLKLPDS-----EAHAVFKYENVEKEDEAPKM-CGVTTQ-NWKSYPEIKKASQLAFTA			
CAT	GRIENDADSTASISACNLKGFHKLQEGEMYLIEPLKLPDS-----EAHAVYKYENVEKEDEAPKM-CGVTTQ-NWESYEP1KKASQLVTTA			
RHO	GRIHNDADSTASISACNLKGFHKLQGETYFIEPMKLPDS-----EAHAVFKYENIEKEDESPKM-CGVTTQTNWESDEPIKKSQNLNHN			
TRM	GRIHNDADSTASISACDGLKGFHKLQGETYFIEPLKIPDS-----EAHAVFKYENVEKEDEAPKM-CGVTTQ-NWESDES1KKASQLVLT			
TRG	GRIENDADSTASISACDGLKGFHKLQEGEMYLIEPLELSDS-----EAHAVFKYENVEKEDEPPKM-CGVTTQ-NWESYESTKKASQLNVTP			
HAL	GRVENDADSTASISACDGLKAHFKIQEGEMYLIEPLEVSDT-----DAHAVFKYENVEKEDEPPKM-CGVTTQ-NWESYESTKKASQLNVSP			
AHL	GRIQNDADSTASISACNLKGFHKLQEGEMYLIEPLELSDS-----EAHAVFKYENVEKEDEAPKI-CGVTTQ-NWESYEP1KKASQLNLNY			
ATRe	GRIENDADSTASISACNLKGFHKLQEGEMYLIEPLKLPDS-----EAHAVFKLNVEKEDEAPKM-CGVTTQ-NWESYEP1KKASDLNLNP			
ARTB	GRIENDADSTASISACNLKGFHKLQEGEMYLIEPLELSDS-----EAHAVFKLENVEKEDEAPKM-CGVTTQ-NWESYEP1KKASDLNLNP			
ATRC	GRIENDADSTASISACNLKGFHKLQEGELYLIEPLELSDS-----EAHAVFKLENVEKEDEAPKM-CGVTTQ-NWESYEP1KKASDLNLNP			
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...	363			441...
MATR	GPFLGG--DAHFDDKEYWTDDEAGVNFLEAAT	HEFGHSLGL--SHS	SVPGTV-----	
STRM	GPFGYV--DAHFDDDEKWSLSSG-GTNLFLVAA	HELGHSLGL--FHS	NNKESL-----	
COLG	GNVYGG--DAHFDDDETSSSK-GYNLFLVAA	HEFGHSLGL--DHS	KDPGAL-----	
TSG	GAAYVGGICSLSHGGGVNEYGN--MGAMAVTLA	QTLGQNLGMMWNKHR	SSAGDKCCKPDIWLG--CIMEDTGFY--LPR	
FER α	GQAFNLGACSSGFAAAVEAFPHBEDALLS-AALLV	HELGHNLGI--RHD	-HS-ACVCRDKH--SCLMQENITEESG--	
FER β	GAIFQGMICNTSYGGGIALHSKTITLDSFGVILV	QLLSVSMGI--AYD	-NADLCRCRGA--ICLMSPEAVFSSGMK	
MDC1	GATYHGMACDPKFATGIALYPKK1TVEAFSVVMA	QLLGINLGL--TYD	-DIYNCPGPG--TCIMNPDAIRSHGMK	
MDC2	GATFPQVCKDFAAVAALYPEGLSLESYTVIIV	QLLGLNLGL--TYD	-KTDTCCHSGD--VCTMTPKAVYSGGVK	
CYRT	GATYHGMACNPNFTAGIALHPKTLAVEGFAIVLS	QLLGINLGL--AYD	-DVYNCFCPGS--TCIMNPASAIRSQGIK	
MS-2	GLAKVSALC-SRHSAGVNDHNSKNSIGV-ASTMA	HELGHNLGM--SHD	EDIPGCYCEPREGGGC1MTE-SIGSKFPR	
EAPr	GIAYPGGICQTLRSCSVKDLLPDVNI1-GNRMA	HQLGHSLGM--RHD	D-FP-CTCPL--GK-CVMGA--GSIPAI	
EAPm	G1SYPAGMCLPYSTSI1KDLLPDTNI1-ANRMA	HQLGHNLGM--QHD	E-FP-CTCPS--GK-CVMDS-D-GSIPAL	
ECh1	GIARNRGMCSPPNSVGV1QDYCKNYLLV-AITMA	HELGHNLGM--DHD	N-GNCCPD--TS-CIMSA-VAGPEPVF	
ECh2	GLRDVSSMCQATRSVGVVQDHSPTVRAV-AVTMA	HEMGNLGM--SHD	G-NHCNC-G--ANSCIMAA-VLRNPAPPE	
JARH	GYAYIGSMCHPKRSVGIQDYSINLVV-AVIMA	HEMGNLGI--HHD	T-GSCSC-G--DYPCIMGP-TISNEPSK	
CATR	GLAVVGSMPKPRSTGI1QDYSBINLVV-AVIMA	HEMGNLGI--NHD	S-GYCSG-G--DYACIMRP-EISPEPST	
RHOD	GKAYLDSTCDPERSVGIQNYHG1ITLNV-AAIMA	HEMGNLGV--RHD	G-EYCTCYG--SSBCIMSS-HISDPPSK	
TRIM	GWAVVGRMCDEKYSVAVVKDHSKVFVAV-AVTMT	HELGHNLGM--EHD	D-KDKCK-D--T--CIMS-VISDQSK	
TRIG	GRAPVGGMCDPKRSVAIVRDHNAIVFVV-AVTMT	HEMGNLGM--HHD	E--DKCNC-N--T--CIMSK-VLSRQPSK	
HALS	GRAPVGGMCDPKRSVAIVRDHNAIFIV-AVTMT	HEMGNLGM--HHD	E--DKCNC-N--T--CIMSK-VLSRQPSY	
AGKH	GLAVVGTMCDPKLSSTGVVEDHSKINFLV-AVTMA	HEMGNLGM--RHD	T-GSCSC-G--GYSCIMSP-VISDPSK	
ATRe	GRAYIGGICDPKRSTGVVQDHSBINLV-AVTMT	HELGHNLGI--HHD	T-DSCSC-G--GYSCIMSP-VISDPSK	
ARTb	GRAYTSSMCPKRSVGIQDHSBINLLV-AVTMT	HELGHNLGM--NHD	G--DKCLR-G--ASLCIMRP-GLTPGRSY	
ATRC	GLAPLGTMCDPKLSIGIVQDHSBINLLM-GVTMA	HELGHNLGM--EHD	G--KDCLR-G--ASLCIMRP-GLTKGRSY	
	* . . . * . . . * . . . * . . . * . . . * . . . * . . . * . . . * . . . * . . . * . . . * . . . *			

Fig. 2. Segments extracted from the alignments of the MMP and MDC complete precursor sequences comprising the cysteine-switch and zinc-binding motifs (*bold*). The positions corresponding to residues conserved among all MDCs (*) or among snake toxins and at least two mammalian MDCs (•) are indicated. The zinc-binding motif and its homologue in nonproteolytic MDCs are *boxed*. The *numbers* correspond to the amino acid position in the consensus sequence of the original alignment. *Abbreviations* as in the legend to Fig. 1.

quences in metalloproteinase and pro-domain alignments, and Fertilin α , whose disintegrin domain does not cluster with other sperm proteins.

The apparent monophyletic distribution of MDC proteins and their independent domains suggests that both mammalian and snake venom proteins have evolved from a common ancestor gene (already assembled as the multidomain structure) both by speciation and by gene duplication. The evolution of a gene family with different functions is currently associated with rapid amino acid divergence among duplicated copies of the genes, thus increasing the functional diversity of the gene family (Ohta 1994). Certainly in the case of snakes, such diversity would seem to be of real benefit because it may result in rapid variation in venom toxicity which could broaden the spectrum of available prey (Daltry et al. 1996). In some cases, a single amino acid change in these

molecules could significantly alter their toxicity in venom, and an accumulation of such changes may account for a rapid amino acid divergence. An example of this is the functionally distinct members of venom phospholipases A₂ (Moura-da-Silva et al. 1995) and members of the serine proteinase family (Creighton and Darby 1989), which are thought to have arisen through gene duplication followed by divergence of the copies through positive Darwinian selection causing sequence hyper-variability. However, serine proteinases present a slightly different evolutionary history since the different domains of the molecules appear to have evolved as independent units rather than as an already-assembled block (Ikeo et al. 1995).

With regard to points in the tree at which gene duplications may have occurred, it seems reasonable to suggest that gene duplications generating the α and β chains

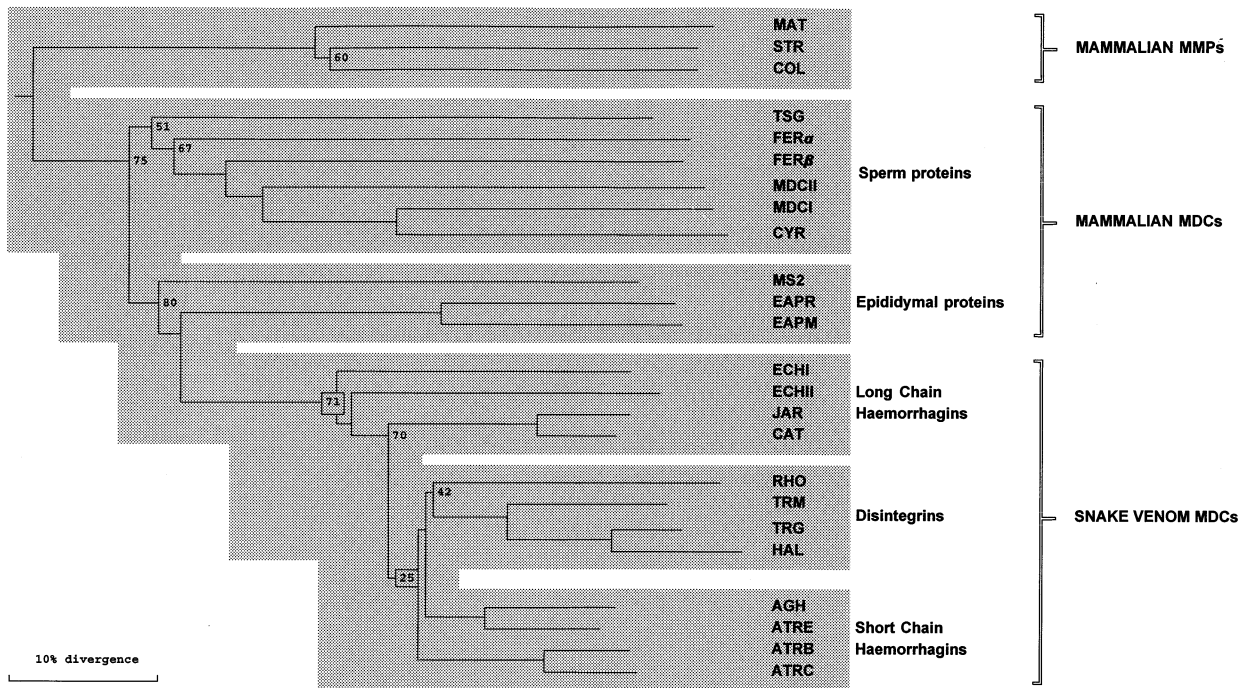


Fig. 3. Phylogenetic tree of the MDC proteins: All amino acid positions predicted by the cDNA regions coding for the whole protein have been included in the analysis. The *horizontal scale* is proportional to the calculated distance. The *numbers* indicated at nodes represent the percentage confidence limits by bootstrapping. Only those values less than 95% are indicated. *Abbreviations* as in the legend to Fig. 1.

of Fertilin and RGD and non-RGD venom disintegrins may have occurred. Gene duplications may also have taken place between EAPs and Fertilin and between short-chained hemorrhagins and RGD-disintegrins. The generation of the different venom MDC proteins has been correlated with post-translational proteolytic processing or RNA splicing (Bjarnason and Fox 1994). However, if these were the only mechanisms operating, a taxonomically related duplication would be expected, at least for the trees generated with the sequences comprising the metalloproteinase and pro-domains. The distribution of sequences showed by the trees does not eliminate the possible occurrence of different processing of RNA and translated proteins; however, if this is the case, alternative processing may be occurring on products generated by different gene copies.

The trees also suggest that the venom toxins arose relatively late during the evolution of MDCs. According to this data, venom metalloproteinases have appeared from a common ancestor gene only after mammals and reptiles diverged; copies of the gene having evolved in snakes to become venom toxins and, perhaps, proteins with some function in the male reproductive tract. However, no work has yet been carried out to identify MDC proteins in snake sperm cells or in the epididymis.

MDCs and MMPs: Evolutionary Conservation of the Proteolytic Motifs or Sequence Convergence?

With regard to the evolutionary history of the MDCs, how can one explain the similarity of the zinc-binding

and cysteine-switch motifs of venom MDCs with MMPs? The similarity mainly confined to the proteolytic motifs has already been shown for microbial and mammalian metalloproteinases (Jongeneel et al. 1989), and Woessner (1991) considered that convergent evolution could explain such similarity. Convergent evolution could also be applied to explain the structural resemblance of the proteolytic motifs between snake venom MDCs and mammalian MMPs. Considering this possibility, the zinc-binding motif may have arisen after a gene duplication, during the early divergence of the reproductive tract MDCs. One copy might have accumulated point mutations introducing the histidines, conferring a selective advantage and resulting in the proteolytic MDCs, which became fully functional in the venom proteins. The other unmutated copy evolved as the nonproteolytic MDCs. In favor of this hypothesis, two out of the three histidines critical for the zinc-binding replace very conserved residues in nonproteolytic MDCs, glutamine, and tyrosine. A single point mutation is needed to change the triplet codon that codes for both amino acids to histidines. The third histidine shares a very variable position in nonproteolytic MDCs, where substitutions might be expected. The cysteine-switch motif may have arisen later due to the need to regulate the proteolytic activity. The mammalian proteolytic MDCs containing a cysteine in the same position of the pro-domain that might possess functional activity, but lack other postulated consensus regions between MMPs and venom MDCs that consist of four residues, PXCGV (Fig. 2).

However, we have to consider a second possibility in which a common ancestor gene to MDCs and MMPs may have carried the proteolytic motifs which were later lost in the nonproteolytic MDCs. This hypothesis is supported by the similarity between microbial and mammalian metalloproteinases. De Souza and Brentani (1993) suggest that the similarity of the zinc-binding motifs arose in these molecules by conservation of sequences present in ancestral genes, while other domains of the molecule, such as the hemopexin-like domain, could be a recent acquisition of the eukaryotic metalloproteinases. The topology of the phylogenetic tree presented in this paper would support the hypothesis that MMPs and MDCs are derived from a common ancestor bearing the zinc-binding motif. The recent assembly of the disintegrin domain on MDCs generated a new functional possibility for the molecules, possibly independent of the proteolytic activity. The selective pressure on mammalian MDCs could therefore be attributed to a cell-matrix/cell-cell adhesion function, thus explaining the loss of the proteolytic motif. On the other hand, the main toxicity of certain viper venoms seems to be proteolytic. This observation would suggest a high selective pressure, and hence conservation, of the proteolytic motifs in the venom MDCs, despite the marked divergence of the remaining parts of the molecules, presumably enhanced by positive Darwinian selection. This hypothesis therefore explains why only the proteolytic motif appears to be shared between MMPs and MDCs in relation to both their sequence and conformation (Fig. 2).

In conclusion, the functional diversity of the MDC proteins may have been generated by gene duplication and divergence of common ancestor genes. The functional proximity of venom MDCs and MMPs can best be explained by evolutionary conservation of the proteolytic motifs rather than by sequence convergence.

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