Tunicates illuminate the enigmatic evolution of chordate metallothioneins by gene gains and losses, independent modular expansions and functional convergences

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ABSTRACT
To investigate novel patterns and processes of protein evolution, we have focused in the metallothioneins (MTs), a singular group of metal-binding, cysteine-rich proteins that, due to their high degree of sequence diversity, still represents a black hole in Evolutionary Biology. We have identified and analyzed more than 160 new MTs in non-vertebrate chordates (especially in 37 species of ascidians, 4 thaliaceans and 3 appendicularians) showing that prototypic tunicate MTs are mono-modular proteins with a pervasive preference for cadmium ions, whereas vertebrate and cephalochordate MTs are bi-modular proteins with diverse metal preferences. These structural and functional differences imply a complex evolutionary history of chordate MTs—including de novo emergence of genes and domains, processes of convergent evolution, events of gene gains and losses, and recurrent amplifications of functional domains—, that would stand for an unprecedented case in the field of protein evolution.
1. INTRODUCTION

Heavy metals such as zinc (Zn) and copper (Cu) are integral components of numerous proteins involved in multiple cellular functions (reviewed in (Vallee and Falchuk 1993; Maret 2013; Vest, et al. 2013)), while other metals such as cadmium (Cd) have no known biological functions and are highly toxic for living beings. Organisms have developed different molecular mechanisms that regulate the homeostasis of the physiological metals and counteract the harmful effects of the toxic ones. One of such mechanisms is based on an extremely diverse group of metal-binding proteins known as metallothioneins (MTs), which are engaged in the physiological control of metals operating as ion reservoirs, metal transporters and/or metal deliverers to target metalloproteins, but also in radical scavenging, oxidative stress protection and anti-apoptotic defense (reviewed in (Capdevila, et al. 2012; Ziller and Fraissinet-Tachet 2018)).

Most MTs are low molecular weight (<100 amino acids) and cysteine-rich (≈15%-30%) proteins. Their cysteine (Cys, C) residues are arranged in CxC, CC and CCC motifs, whose number and distribution define functional domains. Many MTs have a bi-modular organization made of two functional domains, in which each domain enables the coordination of a number of metal ions through metal–thiolate bonds (reviewed in (Blindauer 2014)). MTs have been classified in a stepwise gradation, from extreme Zn/Cd-thioneins to extreme Cu-thioneins depending on their metal-binding preference (Bofill, et al. 2009; Palacios, et al. 2011a). There have been attempts to predict the metal preference of a given MT from the analysis of their non-Cys amino acids, and in the case of gastropod MTs, for instance, it has been proposed that the lysine-asparagine (K/N) ratio could be important for metal preference, so that a preponderance of K over N residues would be a feature of Cd selectivity, whereas a predominance of N over K would be typical of Cu-MTs (Perez-Rafael, et al. 2014; Pedrini-Martha, et al. 2020). Due to the high sequence and structural diversity of MTs, however, it is unclear whether the rule of K/N ratio applies to non-gastropod MTs, and thereby, metal-binding assays are needed to experimentally determine the metal preference of newly discovered MTs.

Comparative analyses of the MTs across diverse species have been useful to better understand the function and evolution of these proteins, and to illustrate unusual modes of molecular evolution of these genes and proteins. MT research has exposed, for instance, pervasive events of internal sequence duplications (Tanguy and Moraga 2001; Palacios, et al. 2014a; Iturbe-Espinoza, et al. 2016; Jenny, et al. 2016; Calatayud, et al. 2018), de novo emergence of domains (Calatayud, et al. 2021b), and a high bias in the amino acid composition probably as a result of the 3D structural
requirements for metal binding. MT research has also provided useful information for many other research fields. In the field of Evolutionary Ecology and Environmental Sciences, for instance, MTs are considered a good model to investigate the mechanisms by which different organisms—from protists and fungi to insects, mollusks and mammals—have adapted to diverse metal bioavailability and other environmental stress conditions. These mechanisms include duplications of MT genes (Maroni, et al. 1987; Adamo, et al. 2012; Luo, et al. 2020), increases in the levels of MT expression (Timmermans, et al. 2005; Janssens, et al. 2008; Janssens, et al. 2009; Costa, et al. 2012; Catalan, et al. 2016; de Francisco, et al. 2018) and changes in metal specificity (Tio, et al. 2004; Palacios, et al. 2011b; de Francisco, et al. 2017; Dallinger, et al. 2020; García-Risco, et al. 2021).

In the fields of Bioinorganic Chemistry and Metallomics, MTs have been used to analyze metal-protein interactions, analyzing the qualitative and quantitative characteristics of the metal-binding capacities of these proteins (Blindauer and Leszczyszyn 2010; Palacios, et al. 2011b; Beil, et al. 2019) and by extension, of a broad range of metalloproteins (Waldron, et al. 2009).

In this work, we aim to explore the MTs found in a large number of species of the three tunicate classes—Asciidae, Thaliaceae and Appendicularia—in order to investigate the intriguing origin and evolution of MTs in chordates. Tunicates (Tunicata subphylum, a.k.a. Urochordata) are the sister group of vertebrates, and together with basally branching cephalochordates form the phylum Chordata. Tunicates have undergone a rapid and ‘liberal’ pattern of evolution in comparison with the ‘conservative’ pattern of vertebrates and cephalochordates (Somorjai, et al. 2018; Ferrández-Roldán, et al. 2019a), yielding a group of ecologically diverse filter-feeding marine animals—including planktonic and benthic specimens as well as solitary and colonial forms—, which have adapted to different conditions of metal bioavailability. In addition, the filtering lifestyle makes tunicates prone to accumulate metals from the seawater ((Papadopoulou and Kaniás 1977; Tzafriri-Milo, et al. 2019) and references therein), and thereby, molecular mechanisms such as the MTs that regulate metal homeostasis and detoxification are physiologically relevant to them.

Our understanding of the MTs in tunicates is, however, scarce, limited to two MT sequences from the ascidian Ciona robusta (formerly C. intestinalis) and Herdmania curvata (Franchi, et al. 2011), and two sequences from the appendicularian Oikopleura dioica recently described by our group (Calatayud, et al. 2018). To overcome this situation, we have conducted an exhaustive survey of the MTs in tunicate databases, identifying 168 MTs in 37 species of ascidians, 4 of thaliaceans and 3 of appendicularians, representing planktonic and benthic species, as well as solitary and colonial forms. We have also functionally characterized the metal-binding properties of
eight tunicate MTs, which were selected on the basis of their taxon representativeness, sequence composition, degree of multiplicity or modular organization. Comparative analyses of tunicate MTs with those of cephalochordates and vertebrates have exposed a high degree of structural diversity between lineages, which would not have been accompanied, however, by differences in their metal preferences. This diversity seems the result of numerous and intricate evolutionary events, and therefore, chordate MTs would represent one of the most complex and enigmatic examples of protein evolution.

2. RESULTS

The biological interest for tunicates in the last decades has given rise to many genomic and transcriptomic resources, facilitating the in silico identification of virtually any gene in a wide set of species. This advantageous situation allowed us to perform an extensive survey of the MT system in tunicate databases.

2.1. MTs in ASCIDIANS.

Ascidians, commonly known as sea squirts, are the most studied group of tunicates. Adults are sessile, either solitary or colonial, usually found in shallow water worldwide. We surveyed ascidian databases and identified 145 MT sequences from 37 species distributed among Aplousobranchia (3 species), Phlebobranchia (8 species) and Stolidobranchia (3 molgulidae + 5 pyuridae + 18 styelidae species) orders (Figure 1; see Table S1 for the species list). Sequence comparison showed that most ascidian MTs are 38-45 amino acids long with 12 cysteines (31%-27%) organized in a single functional domain. In this domain, the C-motifs were distributed as \([CxC]_x[CxC]_x[CxC]_x[Cx3-6CC]_x\), which would represent the prototypical organization of the mono-modular ascidian MTs (Figure S1 and Table 1).

Twenty-seven out of the 37 analyzed species showed multiple MT genes: seven species with two sequences, six with three, two with four, three with six, two with seven, four with eight, one with nine, and two species with up to ten sequences. Despite in some cases, due to the nature of the data source –mainly sequence read archive (SRA) projects–, allelic variants of a particular gene could not be discriminated from gene duplicates, overestimating thereby the degree of MT multiplicity in our catalog, our results suggested that, overall, the MT multiplicity in ascidians is pervasive, although a definitive evaluation of the degree of multiplicity must await more complete and deeper genomic databases.

Interestingly, there were some exceptions to the prototypical organization, ranging from MTs that have lost or gained some cysteines or small protein fragments (Figure S1) to multi-modular forms containing tandem repeats of the prototypical
domain (Figure S2 and S3). We were able to reconstruct 12 multi-modular MTs (five partial and seven full-length sequences), whose number of repeats ranged from two to nine (Figure 2). Noteworthy, all the species with multi-modular MTs belonged to the styelidae family in the stolidobranchia order (Figure 1), suggesting that multi-modular MTs arose by a lineage-specific evolutionary event.

2.2. Metal-binding features of mono-modular ascidian MTs

In order to analyze the metal-binding capacity and preference of the ascidian MTs, we selected three MTs of two species belonging to distinct orders within the Asciidiacea class: CroMT1 from C. robusta (phlebobranchia order) and HroMT1 and HroMT2 from Halocynthia roretzi (stolidobranchia order). These MTs were chosen because CroMT1 was the only MT in C. robusta, whereas HroMT1 and HroMT2 represented duplicate forms in H. roretzi. These MTs were also selected because of their K/N ratios, which in the CroMT1 was 3:5, opposite of that in HroMT1 and HroMT2 sequences, 6:3 and 8:2, respectively. HroMT2 showed indeed one of the highest K/N ratios among tunicate MTs (Table S1). The K/N ratio seems important for metal preference of gastropod MTs (Perez-Rafael, et al. 2014; Pedrini-Martha, et al. 2020), and the study of the metal selectivity of the CroMT1, HroMT1 and HroMT2 would allow us to experimentally evaluate the K/N rule in the ascidian system.

We characterized the formation of metal–MT complexes of CroMT1, HroMT1 and HroMT2 proteins heterologously expressed in E. coli cultures supplemented with Cu, Cd or Zn salts by inductively-coupled plasma atomic emission spectroscopy (ICP-AES) and electrospray ionization mass spectrometry (ESI-MS). ICP-AES was used for protein quantification and metal-to-protein stoichiometry determination through the measurement of element composition of the samples (S, Zn, Cd and Cu) (Bongers, et al. 1988), and ESI-MS allowed us to determine the molecular mass of the species formed, i.e. the speciation of the samples (Capdevila, et al. 2012). Single well-folded Cd₄-MT species were recovered from CroMT1, HroMT1 and HroMT2 Cd-productions (Figure 3A-C), and a major Zn₄-MT species were found in their recombinant Zn productions (Figure S4A, S4C and S4E), which contrasted with the mixtures of homometallic Cu-MT species purified from Cu-supplemented E. coli cultures (Figure S4B, S4D and S4F). It is also noteworthy the resemblance of the CD spectra of the Cd-MT complexes of all ascidian MTs, denoting structurally equivalent metal clusters for these proteins (Figure 4). The exemplary exciton coupling centered at ca. 250 nm exhibited in all the CD envelopes confirms high robustness and compactness of the Cd₄S(Cys)₁₂ clusters and, therefore, high Cd specificity. Altogether, these results showed that Zn₄- or Cd₄-MT complexes were energetically the most favorable forms of these proteins, suggesting that their function would be related to binding divalent metal...
ions, either as Zn- or Cd-thioneins [8]. Genuine Zn-thioneins, however, render heterometallic Cu,Zn-MT complexes when synthesized under Cu surplus, meaning that the ascidian MTs deviated towards a more specific Cd-thionein behavior (Palacios, et al. 2011a). Regarding the K/N rule used to predict the metal-binding preference in gastropod MTs, our results demonstrated that this rule did not apply to tunicate MTs, since the three of them displayed contrasted K/N ratios and their metal preferences were exactly the same.

2.3. METAL-BINDING FEATURES OF MULTI-MODULAR ASCIDIAN MTs

In order to determine the metal-binding features of multi-modular MTs, we selected BscMT1 (ID: ATSW01006707.1; chromosome 6) of Botryllus schlosseri (stolidobranchia order) because this MT was not only identified in an accurately assembled genome (Voskoboynik, et al. 2013), but also because it was supported by 3 EST sequences (Table S1). BscMT1 was a multi-modular MT of 365 amino acids with 105 cysteines (29%) and a K/N ratio of 1:1 (31K vs. 31N) (Figure S2 and Table S1). BscMT1 was one of the longest MT described so far, and a detailed analysis of its sequence revealed that it was made of nine tandem repeats (R1-R9, the last one only partially complete) of a 12-Cys domain, each one resembling a prototypical ascidian form (Figure S3).

The metal–BscMT1 complexes purified from the heterologous expression in E. coli cultures supplemented with Cu, Cd or Zn salts were characterized by the IPC-AES and ESI-MS. The concentrations of metal–BscMT1 productions, despite the several attempts, were always low, probably due to the high molecular weight of BscMT1 and its structural complexity made of repeated domains. Only the samples recovered from the Zn(II)-enriched culture media rendered informative data through these techniques. The IPC-AES and ESI-MS analyses showed that this preparation contained Zn36-MT complexes (Figure 3D). The binding of 36 metal ions agreed with its multi-modular organization made of 9 repeated domains, each one similar to a prototypical Mt4-MT: 4 metal ions x 9 domains = 36 metal ions. To further verify the functional correspondence between one of the 12-Cys tandem repeats of BscMT1 and the mono-modular ascidian MTs, we analyzed the ability of the R4 domain (R4-BscMT1) to form metal complexes when it was expressed alone. The election of the R4 domain was due to the fact that it was highly similar to the prototypical ascidian MTs, including some additional amino acids after the carboxyl-terminal CC doublet (Figure S3). We heterologously expressed the R4 domain in E. coli grown in media supplemented with Cu, Cd or Zn salts. The ICP-AES and ESI-MS analyses of the recovered samples confirmed the ability of this fragment to autonomously form metal-complexes by itself, and revealed that the biochemical features of R4 domain were highly similar to the mono-modular
ascidian forms (compare Figure 3A-C and 3E). Thus, this domain yielded major M₄-R₄ complexes with Zn(II) and Cd(II), being unique species only in the latter, while rendered a homometallic mixture of species in the presence of copper, as the ascidian MTs discussed above (Figure S4G and S4H). Based on this R₄ fragment as a representation of the whole protein, the multi-domain BscMT1 would be a Cd-thionein capable to bind up to 36 divalent ions, and its stoichiometry —i.e. M₃₆-BscMT1— would nicely match with what would be expected from the repetition of nine domains, i.e. M₉-R₁ to M₄-R₉. These results supported a direct relationship between the binding capacity and the number of the repeated domains in multi-modular MTs.

2.4. MTs in Thaliaceans

We analyzed the thaliacean MTs by surveying the databases of the five available species. We identified 14 MTs from four thaliacean species, three from Salpida (Iasis cylindrica, Salpa fusiformis and Salpa thompsoni) and one from Pyrosomata (Pyrosomella verticillata) orders. Sequence comparison showed that thaliacean MTs were 36-41 amino acids long, with 12 cysteines (33%-29%) organized in a single functional domain with the C-motifs distributed as [CxC]ₓ₃[CxC]ₓ₃[CxC]ₓ₃[CxC]ₓ₃-C(3-4)CC (noteeworthy, we did not find multi-modular MTs in thaliacean databases) (Table 1). This motif arrangement was identical to that of the prototypical ascidian MTs, denoting a common origin of that thaliacean and ascidian MTs, and supporting the recently suggested phylogenetic relationship of these classes (Delsuc, et al. 2018; Kocot, et al. 2018) (Figure 1).

Importantly, MT multiplicity were found in the four thaliacean species although as in ascidians, it was challenging to determine whether the multiple sequences represented different genes or allelic variants. In S. thompsoni, for instance, we identified six MT sequences, five full-length (SthMT1, MT2, MT3, MT5 and MT6) and one partial (SthMT4), which coded for proteins 48% to 95% identical at amino acid level (Table S2). The highest identities were between SthMT1 and SthMT5 (88%) and SthMT2 and SthMT6 (95%) sequences, and therefore, these pairs were the most plausible cases of allelic variants. To investigate this possibility, we analyzed the conservation of 5’ and 3’ flanking regions, as well as of intron sequences in PCR-amplified genomic regions from two S. thompsoni specimens. Nucleotide identity (excluding gaps) in SthMT1-SthMT5 and SthMT2-SthMT6 alignments was 91%-100%, 87%-82% and 83%-86% at 5’ regions, introns and 3’ regions, respectively (Table S3). These results would be compatible with that SthMT5 and SthMT6 were allelic variants of SthMT1 and SthMT2, respectively, although the possibility they may be recently duplicated genes cannot be fully discarded.
2.5. METAL-BINDING FEATURES OF THALIACEAN MTs

We analyzed the metal-binding capacity and metal preference of the thaliacean MTs, and compared them with those of ascidian MTs in order to investigate whether the differences in lifestyles – planktonic vs. benthic—might have been key for MT evolution. We characterized the metal–MT complexes rendered for the four MTs from *S. thompsoni*, SthMT1 to SthMT4, heterologously expressed in Zn-, Cd- or Cu-supplemented *E. coli* cultures. The clear Cd-thionein character of *S. thompsoni* MTs was reflected, once again, by the ICP-AES and ESI-MS analyses of the recovered samples. SthMTs yielded single Cd₄-MT species under Cd(II) surplus (Figure 5), as well as major M₄-MT complexes with Zn(II) (Figure S5A-S5D). A mixture of species (for SthMT1 and SthMT3) or not detection of the metallated species (for SthMT2 and StMT4) were obtained in the presence of copper (Figure S5E-S5F), evidencing that these proteins struggle to build metal clusters with Cu. Our results demonstrated that the four SthMTs were functionally very similar among them (compare panels A to D in Figure 5), as well as to the mono-modular ascidian MT (compare Figure 3 and Figure 5). The fact that thaliacean (SthMTs) and ascidian (CroMT1, HroMT1, HroMT2 and BscMT1) MTs exhibited a similar Cd-thionein character strongly supported that the prototypical ascidian/thaliacean MT (and most likely the ancestral tunicate MT, see below) was a mono-modular protein with a 12C domain capable to bind four metal ions with preference for Cd.

2.6. MTs IN APPENDICULARIANS

We investigated the MTs in the Appendicularia (a.k.a. Larvacea) class. In a previous work, we had characterized the *OdiMT1* and *OdiMT2* genes (formerly *OdMT1* and *OdMT2*) of *O. dioica* (Calatayud, et al. 2018). We showed that both genes encoded MTs very different from all MTs described so far because their cysteines motifs were arranged in a novel pattern forming a “repeat unit” (RU) made of two C7 subunits (C7a and C7b) plus a carboxyl-terminal tail with 5 cysteines (C5) (Calatayud, et al. 2018) (Figure S6A). We concluded that OdiMT1 was made of a single RU, while OdiMT2 was a multi-modular protein made of six direct tandem RU (Calatayud, et al. 2018). We wondered whether these unconventional MTs were specific of *O. dioica*—gene and protein sequences are known to have considerably diverged during *O. dioica* evolution (reviewed in (Ferrández-Roldán, et al. 2019b))—or they were widespread in the appendicularian lineage. We took advantage of recent sequence projects of several appendicularian species (Naville, et al. 2019) and identified seven new MT sequences in two additional Oikopleura species: *O. albicans* (3 sequences) and *O. vanhoeffeni* (4 sequences) (Table S1). We did not identify any putative MT in databases of other oikopleurids such as *O. longicauda, Bathochordaeus sp.* and *Mesochordaeus*
erythrocephalus, nor in the fritillarid species Fritillaria borealis, although any conclusion about the loss of the MTs in these lineages must await deeper sequence projects.

Regarding Oikopleura MTs, although most of the sequences were partial, they were clearly similar to O. dioica MTs. Sequence analyses showed that these new MTs were multi-modular proteins made of repeats of a conserved 12-Cys domain with C-motifs distributed as CX₃Cx₂Cx₃Cx₂{Cx₃}x₂Cx₄{Cx₃}xCC (Table 1 and Figure S6B). This 12-Cys domain corresponded to the previously described C7b+C5 subunits (Calatayud, et al. 2018). OdiMT1 consist, therefore, of two 12-Cys domains, one full-length (12C=C7b+C5) and one 12-Cys domain that has been trimmed (t-12C, formerly known as C7a) and lacks the five last cysteines (C5) (Figure S6A and Figure 7). Interestingly, the trimmed domain when expressed alone did not efficiently bind divalent metals by itself, suggesting that it has lost its capacity to independently work (Calatayud, et al. 2021a). OdiMT2 would be a multi-modular MT made of twelve 12-Cys domains, whose odd domains would have been also trimmed as t-12C. The identification of this conserved 12-Cys domain in all Oikopleura MTs suggested that this domain appeared in the ancestor of the Oikopleura genus, and that it was duplicated generating multi-modular MTs independently in the diverse appendicularian species (Figure S6B).

2.7. MTs in Other Chordates

To provide a broad evolutionary perspective to our analyses we compared the tunicate MTs with those from the other chordate subphyla: cephalochordates and vertebrates.

In cephalochordates, MTs had been characterized in two amphioxus species, Branchiostoma floridæ and Branchiostoma lanceolatum (Guirola, et al. 2012). We extended the surveys of MTs to two additional amphioxus species, Branchiostoma belcheri (BbeMT1 and BbeMT2) and Branchiostoma japonicum (BjaMT1 and BjaMT2), and to Asymmetron lucayanum (AluMT1 and AluMT2), a cephalochordate distantly related to the other Branchiostoma species (Yue, et al. 2014). We identified two MTs, MT1 and MT2, in all the cephalochordate species. B. lanceolatum MT2 showed two alternatively spliced forms, MT2-L (for the long form, with 67-70 amino acids) and MT2-S (for the short form, with 46-47 amino acids) (Figure S7). Splice variants were also identified in B. floridæ, B. japonicum and A. lucayanum cDNA databases (i.e. BflMT2L and S, BjaMT2L and S, and AluMT2L and S), and alternative splicing was predicted from genomic sequence in B. belcheri species (BbeMT2L and S).

Sequence comparisons revealed that cephalochordate MT2s were multi-modular proteins made of two (MT2S) or three (MT2L) 9-Cys domains (CCx₂Cx₂Cx₃Cx{Cx₃}xCC) depending on an alternative splicing on exon 2 (Figure S7.)
and Figure S8B). In contrast, MT1s were bi-modular proteins with two different domains: a 11-Cys, N-terminal domain with the C-motifs distributed as \([\text{CxC}]_4[\text{CxC}]_3[\text{CxC}]_2\) (Table 1 and Figure S8A) linked by 9-11 residues to a domain similar to the 9-Cys domain of MT2 (we named this domain 9-Cys\text{Like} because an Ile substitutes the fourth Cys: \(\text{CCx}_2\text{Cx}_1\text{Cx}_3[\text{CxC}]\text{xCxCC}\)) followed by a carboxy-terminal tail of 16-17 amino acids devoid of Cys residues. Interestingly, MT2 9-Cys and the MT1 9-Cys\text{Like} domains showed a clear sequence similarity (Figure S8C), which suggested that cephalochordate MT2s might have arisen from the duplication of an ancestral MT1 that lost the 11-Cys domain but amplified its 9-Cys domain.

In vertebrates, MTs had been classified in four different MT types: MT1, MT2, MT3 and MT4. Vertebrate MTs had been characterized in many species with significant differences of multiplicity in a species-dependent manner (Serén, et al. 2014). Phylogenetic and synteny analyses had suggested the existence of mammalian MT1s and MT2s, amniote MT4s, tetrapod MT3s, and fish MTs (Figure 1) (Serén, et al. 2014). The appearance of these four types would not be related with the two rounds of whole-genome duplication events that occurred during early vertebrate evolution since they are not localized in paralogous chromosomes but are located in tandem on a same chromosome (Serén, et al. 2014). At protein level, all vertebrate MTs (even an MT we identified in the early divergent group of hagfish \(\text{Eptatretus burgeri}\)), showed a conserved bi-modular organization made of a 9-Cys N-terminal domain (the so-called \(\beta\)-domain: \([\text{CxC}]_4[\text{CxC}]_3[\text{CxC}]_2[\text{CxC}]\text{xCx}_3\)) linked to a 11-Cys domain (the so-called \(\alpha\)-domain: \(\text{CCxCx}_3\text{Cx}_2\text{Cx}_3[\text{CxC}]\text{xCx}_3\text{CxCC}\)) at the carboxyl-terminal end (Table S1 and Figure S9). The arrangement of the C-motifs in the \(\alpha\) and \(\beta\) domains did not show any obvious resemblance between them that might suggest of a common ancestry of these domains, though vertebrate \(\beta\)-domain appeared somewhat similar to cephalochordate and tunicate 11/12-Cys domains.

3. Discussion
3.1. Origin and Convergent Evolution of Tunicate MTs

Ascidian and thaliacean MTs share the same arrangement of their C-motifs (Table 1) and are clearly similar in their amino acid sequence, also in the non-cysteine residues: ascidian MTs are at least \(\approx 40\%\) identical in their non-cysteine residues to any of the thaliacean MTs, and \(\approx 60\%\) identical when considering the full sequence. In contrast, their similarity with appendicularian MTs is negligible at the arrangement of the C-motifs (Table 1) and in the composition of the non-cysteine residues. For instance, OdIMT1 only is, at most, 13\% identical to ascidian (e.g. CroMT1 and HroMT1) or thaliacean (e.g. SthMT1) MTs in non-cysteine residues (notice that 25%-
30% sequence identity is generally taken as the minimal threshold for presumption of homology (Rost 1999). Therefore, although it cannot be ruled out that appendicularian MTs derive from ascidian and thaliacean MTs that have diverged too much for orthology to be recognized, their sequence and the C-motifs are so divergent that they seem to have had independent evolutionary origins. This possibility agrees with the idea that de novo emergence of MTs might be more habitual than previously thought because the only requirement for a peptide to function as a metal ion chelator would be a high content of coordinating residues (i.e. cysteines) and a relative small length that would favor its proper folding (Capdevila and Atrian 2011; Calatayud, et al. 2021b). The independent emergence of MTs in distant animal groups have been, indeed, implicitly suggested in diverse evolutionary studies (Capdevila and Atrian 2011; Blindauer 2014; Isani and Carpene 2014; Ziller and Fraissinet-Tachet 2018) we have recently showed de novo evolution of MT domains within the Mollusca phylum (Calatayud, et al. 2021b).

Intriguingly, ascidian/thaliacean MTs and appendicularian MTs seem to have converged in the number of cysteines –twelve– of their basic functional organizations: the prototypical 12-Cys MTs of ascidians and thaliaceans, and the repeated 12-Cys domains of appendicularian MTs. Functional domains with 11/12 cysteines organized in different patterns of C-motifs but converging to similar 3D architectures (Beil, et al. 2019) appear a recurrent evolutionary solution in MTs of phylogenetically distant animal clades, from vertebrates and tunicates to mollusks and insects (Table 1). 3D structural constraints to efficiently coordinate four metal ions through the formation of metal–thiolate bonds might have led to the convergent evolution of 11/12-Cys domains in MTs of different animal phyla ((Stillman, et al. 1987; Egli, et al. 2006; Digilio, et al. 2009; Tomas, et al. 2013) and this work).

3.2. EVOLUTION OF THE METAL BINDING PREFERENCE

Ascidian/thaliacean MTs and appendicularian MTs also share the Cd-thionein character. The evolution of Cd-MTs seems, indeed, a common adaptive event in marine animals, including mollusk (Calatayud, et al. 2021b), crustacean (Narula, et al. 1995; Valls, et al. 2001; Munoz, et al. 2002), echinoderm (Riek, et al. 1999), and chordate species (Guirola, et al. 2012; Calatayud, et al. 2018). The preference of tunicate MTs for Cd further support the idea that ancestral MTs arose and evolved probably driven by the Cd(II) concentrations in the seawater because Cd is highly toxic by interfering in Zn-dependent cellular processes (Dallinger, et al. 2020; Calatayud, et al. 2021b). In addition to sporadic episodes of increases in Cd emissions throughout geological eras (from Paleozoic to Mesozoic and Cenozoic, (Dallinger, et al. 2020) and references therein), Cd is frequently found with Zn in ore deposits of the earth crust.
The natural Cd(II) concentrations are, therefore, higher where seawater comes into contact with the continental earth crust (≈0.8 nmol/kg) than in surface pelagic waters (≈0.002 nmol/kg) (Bruland 1980). Cd(II) concentrations are also higher in eutrophic coastal regions of the littoral and its close neritic zone (≈0.16 nmol/kg) (Bruland 1980), which are usual habitats for tunicates. It is noteworthy that all tunicate MTs analyzed in this work show a similar sequence and metal preference, either they are from benthic ascidians or from planktonic thaliaceans, or from solitary (C. intestinalis and H. roretzi) or colonial species (B. schlosseri). This similarity suggests that the different lifestyles of tunicates might have not been a significant factor in the evolution of metal binding preference.

3.3. EVOLUTION OF MT MULTIPLICITY AND MULTI-MODULAR MTs

Increasing the metal binding capacity enhance the physiological capabilities of the organisms to adapt to diverse conditions of metal bioavailability and other environmental stress situations (Jenny, et al. 2016; Baumann, et al. 2017). In a simplified view, such increments might be attained through two strategies: increasing the number of MT genes (i.e. augmenting MT multiplicity) (Maroni, et al. 1987; Adamo, et al. 2012; Luo, et al. 2020) or increasing the number of metal-binding domains in a given MT (i.e. generating multi-modular MTs) (Jenny, et al. 2016; Baumann, et al. 2017). Tunicates have used both strategies.

3.3.1. Evolution of MT multiplicity

Multiplicity in the MT system has been associated to neofunctionalization or subfunctionalization processes, in which different paralogs have distinct metal-binding preferences and/or expression patterns. For instance, in mollusks, several MTs can be found in a given species, each one with distinct metal selectivity –i.e. Cd-MT, Cd/Cu-MT and Cu-MT– and differential transcription patterns (Palacios, et al. 2014b; Dvorak, et al. 2018; Dallinger, et al. 2020). Similarly, in the mammalian system, ubiquitous metal-induced forms MT1 and MT2 have preference for Zn(II), whereas restrictedly expressed MT3 and MT4 have intermediate Zn/Cu- and Cu-thionein characters (Capdevila and Atrian 2011; Vasak and Meloni 2011; Artells, et al. 2013). MT multiplicity has been observed in 36 tunicate species, patchily distributed among the three tunicate classes. However, in contrast with the observed in the MT system of other animal groups, the analyzed tunicate MT paralogs do not seem to have essentially diverged in their metal-binding selectivity. The two paralogs of the ascidian H. roretzi (HroMT1 and HroMT2), the four paralogs of the thaliacean S. thompsoni (SthMT1 to SthMT4), and the two paralogs of the appendicularian O. dioica (OdiMT1 and OdiMT2 (Calatayud, et al. 2018)) have a conserved Cd selectivity. These results suggest that tunicate paralogs have undergone a subfunctionalization process.
partitioning the expression of the predecessor, rather than a neofunctionalization event acquiring new metal preferences. Future detailed spatiotemporal analyses of the transcription patterns of each paralog will provide a definitive answer to this question, and will allow us to understand the ecophysiological needs that led the evolution of multiple MTs in these species.

3.3.2. Evolution of multi-modular MTs

Evolution of multi-modular MTs seems a recurrent solution for creating large MTs with high cysteine content and a high capacity of metal binding ((Niederwanger, et al. 2017; Palacios, et al. 2017; Calatayud, et al. 2018) and this work). Multi-modular MTs have been found in very diverse organisms, from fungi (Palacios, et al. 2014a; Iturbe-Espinoza, et al. 2016) and mollusks (Calatayud, et al. 2021b) and references therein), to chordates (Guirola, et al. 2012; Calatayud, et al. 2018). Here, we have identified multi-modular MTs in 20 species of ascidian and appendicularian classes (thaliaceans appear devoid of multi-modular MTs). These multi-modular MTs are made of tandem repeats of a prototypical organization within each tunicate class – [CxC]x5[CxC]x2Cx2CxCj[CxC]x3j[CxC]x3;6CC for ascidian and CX3Cx2Cx3Cxj3CxCj2Cx4j[CxC]x2CC for appendicularian MTs –, implying independent evolutionary origins for ascidian and appendicularian multi-modular MTs. In ascidians, the multi-modular MTs arose during the evolution of the Styelidae family in the Stolidobranchia order, after its split from the Pyuridae family, 277 million years ago (Delsuc, et al. 2018). In appendicularians, multi-modular MTs arose after its divergence from the other tunicate classes, 447 million years ago (Delsuc, et al. 2018). The adaptive causes that favored the evolution of large MT in some benthic ascidian species but not in others, or in some planktonic tunicates such as appendicularians but not in thaliaceans, are still an unsolved question.

3.4. THE ENIGMATIC ORIGIN OF THE MTs IN CHORDATES

The extensive catalog of tunicate MTs collected in this work together with the analysis of cephalochordate and vertebrate MTs has allowed us to investigate the origin and evolution of the MTs in the chordate phylum. This has been a particularly difficult challenge because when comparing distant taxonomic groups (e.g. at phylum, subphylum or class level), one has to be cautious with the MT sequence alignments. The general cysteine richness of the MTs distorts the comparisons by forcing the alignment of Cys positions and inserting many gaps in the aligned sequences. This distortion results in a false impression of similarity between non-homologous MT sequences, and if the alignments are used for phylogenetic reconstructions, they can lead to wrong evolutionary inferences. To circumvent this drawback, we focused our analyses on the patterns of the cysteine motifs (CxC, CC and CCC) in the MT domains.
that, as it has been proved for other molecular markers (Rokas and Holland 2000), might have enormous potential as a tool for phylogenetic inferences.

Regarding the ancestral chordate MT, it is generally accepted that due to their substantial genomic stasis (Cañestro and Albalat 2012; Somorjai, et al. 2018; Coppola, et al. 2019), cephalochordate genes and genomes are more similar to those of the chordate ancestor than those of tunicates and vertebrates (Louis, et al. 2012). It can be argued, therefore, that the MT of the chordate ancestor was a bi-modular protein with an amino-terminal 11/12-Cys domain and a carboxyl-terminal 9-Cys domain (Figure 6), resembling the organization of the current cephalochordate MT1s. The similarity of cephalochordate and vertebrate domains (Figure 7) supports the bi-modular organization of the ancestral chordate MT. In addition, the fact that cephalochordate MT1s are also similar to the α and β1 domains of mollusks (Calatayud, et al. 2021b) would push back in evolutionary time this MT organization to the origins of the bilaterian animals.

The evolutionary scenario of a chordate MT ancestor similar to the current cephalochordate MT1 implies a number of evolutionary events independently shaping the MTs in each chordate subphylum: (I) In cephalochordates, an ancient tandem duplication of the ancestral MT followed by the loss of the 11-Cys domain and the amplification of the 9-Cys domain in one of the copies would have led the MT2 forms of the subphylum (Figure 6); (II) In vertebrates, the ancestral MT would have readjusted its Cys motifs, leading to the current amino-terminal β- and the carboxyl-terminal α-domains (Figure 6). Successive lineage-specific tandem duplications would have led to a variable degree of the MT multiplicity in the different vertebrate species (Serén, et al. 2014); (III) In tunicates, the ancestral MT would have lost the 9-Cys domain in the ascidian/thaliacean lineage. This mono-modular 12-Cys MT was patchily duplicated, giving rise to a variable number of MTs in different ascidian/thaliacean species, or internally repeated in the multi-modular MTs of styelidae species (Figure 6); (IV) Finally, in the fast–evolving appendicularians, the ancestral chordate MT would have been lost, and a new MT with a different pattern of Cys motifs seems to have evolved de novo (Figure 6). The primeval appendicularian MT would have been a bi-modular form made of two 12-Cys domains, apparently unrelated with the other 12-Cys domains of chordates. This ancestral appendicularian MT lost five Cys in the N-terminal domain in the O. dioica lineage (but remained intact in other Oikopleura lineages), yielding the t-12C+12C organization of the current OdiMT1. This MT was later duplicated, and internal duplications in one of the copies of a repeat unit (RU) made of the t-12C+12C domains gave rise to the large multi-modular OdiMT2 (Calatayud, et al. 2018).
In summary, our analyses have shown (1) de novo emergence of MT genes and domains during chordate diversification; (2) convergent evolution to 12-Cys domains in chordate MTs, but also in other animals, probably favored by essential constraints of the metal coordination; (3) convergent evolution in the metal preference for Cd ions, probably triggered by an ancestral requirement for cadmium-selective forms in many marine animals; (4) lineage-specific events of gene duplications yielding significant levels of MT multiplicity; and (5) independent amplifications of domains that generated multi-modular MTs with high metal-binding capacities. All these results have led us to reconstruct an intricate evolutionary history for the chordate MTs, from a bi-modal, cadmium-selective form in the last common ancestor, to a high structural and functional MT diversity currently found in the diverse species of the phylum. To discover the adaptive causes that shaped this history and the molecular mechanisms that made it possible will certainly be a fascinating challenge.

4. MATERIAL AND METHODS

4.1. CHORDATE METALLOTHIONEIN SEARCH IN DATABASES

Tunicate and cephalochordate MT sequences were retrieved from public databases (NCBI database: https://www.ncbi.nlm.nih.gov/; and Octopus database: http://octopus.obs-vlfr.fr/public/botryllus/blastbotryllus.php) by Blastp and TBLastn searches using known chordate MTs as queries: tunicate CroMT1 (ACN32211), HcuMTa (AY314949), OdiMT1 (AYN64372) and OdiMT2 (AYN64376); cephalochordate BflMT1 (XP_035672826), BflMT2 (BW841405), BlaMT1 (BL11229) and BlaMT2 (JT872034); and vertebrate HsaMT1A (NP_005937), HsaMT2A (NP_005944), HsaMT3 (NP_005945) and HsaMT4 (NP_116324). In addition, RNA-SRA for each tunicate species deposited in NCBI were Blast searched using as queries MT sequences from the nearest phylogenetic species as well as from different species covering the main tunicate clades. Raw sequence data was retrieved and assembled using SeqMan 8.0.2 (Pro Assembler) software from the DNASTAR Lasergene package, and manually inspected in order to reconstruct new MT sequences. The MT nature of each new identified sequence was evaluated by Blast searches against metazoan NCBI non-redundant protein sequence database. The amino acid sequences and the accession numbers of the retrieved MTs are provided in Table S1. Sequence alignments were generated with Aliview program (Larsson 2014) and reviewed manually.

4.2. CHARACTERIZATION OF S. THOMPSONI GENES.

Salpa thompsoni specimens were kindly provided by Nora-Charlotte Pauli from the Bettina Meyer’s group at Alfred Wegener Institute for Polar and Marine Research
Genomic DNA was extracted using Quick-DNA/RNA™ Microprep Plus Kit Protocol (Zymo Research, CA USA) using 5 mg of tissue and following the manufacturer’s instructions. The DNA concentration, purity and integrity were checked using Tecan Infinite M200 (Tecan Group Ltd, Switzerland) measuring the absorbance at 260 and 280 nm.

Based on S. thompsoni genome and transcriptome projects (PRJNA318929 and GFCC00000000.1, respectively) and the raw data from S. thompsoni SRA projects, several pairs of primers were designed in order to PCR amplify the 5’ and 3’ flanking regions, as well as of intron sequences of the different SthMTs (Table S4).

For SthMT1 we used the primer pairs FwMT1/MT5 with RvMT1/MT2; for SthMT5, FwMT1/MT5 with RvMT5, and finally, for SthMT6 we used FwMT2/MT6 with RvMT6.

For each PCR reaction, 1 ng of genomic DNA was amplified using selected pairs of primers and the Phusion High-Fidelity DNA Polymerase (Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA) in a final 25 µL reaction. PCR conditions were 98°C 30 seconds (s); 35 cycles of 98°C 10 s, 50-54°C 30 s (Table S4) and 72°C 3 minutes (min); and 72°C 10 min. PCR products were visualized in 0.8% agarose gels, isolated with the GelElute Plasmid Miniprep Kit (Sigma-Aldrich, Merck KGaA, Darmstadt, Germany), and cloned with TOPO® TA Cloning® Kit (Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA). Plasmid DNA was purified from bacteria using the GeneElute Plasmid Miniprep Kit (Sigma-Aldrich), screened for insert presence by digestion with Fast Digest EcoRI (Invitrogen, Thermo Fisher Scientific, Waltham, MA, USA), and sequenced at the Scientific and Technological Centers of the University of Barcelona using the Big Dye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems) in an automatic sequencer (ABIPRISM 310, Applied Biosystems). Sequences were analyzed with Unipro UGENE software (Okonechnikov, et al. 2012).

### 4.3. Production and Purification of Recombinant Metal-MT Complexes

Synthetic cDNAs codifying the selected MTs (i.e CroMT1, HroMT1, HroMT2, BscMT1, R2BscMT1, SthMT1, SthMT2, SthMT3 and SthMT4) were provided by Synbio Technologies (Monmouth Junction, NJ, USA), cloned in the pGEX-4T-1 expression vector (GE Healthcare, Chicago, IL, USA) and transformed in protease-deficient E. coli BL21 strain for heterologous expression. For heterologous protein production, 500 mL of Luria-Bertani (LB) medium with 100 µg/mL ampicillin was inoculated with E. coli BL21 cells transformed with the corresponding recombinant plasmid. After overnight growth at 37°C/250 rpm, the cultures were used to inoculate 5 L of fresh LB-100 µg/mL ampicillin medium. Gene expression was induced with 100 µM isopropyl-β-D-thiogalactopyranoside (IPTG) for 3 hours (h). After the first 30 min of induction, cultures were supplemented with ZnCl2 (300 µM), CdCl2 (300 µM) or CuSO4
(500 µM) in order to generate metal-MT complexes. Cells were harvested by centrifugation for 5 min at 9,100 g (7,700 rpm), and bacterial pellets were suspended in 125 mL of ice-cold phosphate-buffered saline (PBS: 1.4 M NaCl, 27 mM KCl, 101 mM Na₂HPO₄, 18 mM KH₂PO₄ and 0.5% v/v β-mercaptoethanol). Resuspended cells were sonicated (Sonifier Ultrasonic Cell Disruptor) 8 min at voltage 6 with pulses of 0.6 sec, and then centrifuged for 40 min at 17,200 g (12,000 rpm) and 4°C.

Soluble protein extracts containing GST-MT fusion proteins were incubated with glutathione sepharose beads (GE Healthcare) for 1 h at room temperature with gentle rotation. GST-MT fusion proteins bound to the sepharose beads were washed with 30 mL of cold 1xPBS bubbled with argon to prevent oxidation. After three washes, GST-MT fusion proteins were digested with thrombin (SERVA Electrophoresis GmbH Heidelberg, Germany, 25U/L of culture) overnight at 17°C, thus enabling separation of the metal-MT complexes from the GST that remained bound to the sepharose matrix. The eluted metal-MT complexes were concentrated with a 3 kDa Centripep Low Concentrator (Amicon, Merck-Millipore, Darmstadt, Germany), and fractionated on a Superdex-75 FPLC column (GE Healthcare) equilibrated with 20 mM Tris-HCl, pH 7.0. The protein-containing fractions, identified by their absorbance at 254 nm, were pooled and stored at -80°C until use.

4.4. ANALYSIS OF METAL-MT COMPLEXES.

Protein quantification and element composition of all the samples were achieved by ICP-AES measurements performed in a Optima 4300DV apparatus (Perkin-Elmer, MA, USA) (S, 182.040 nm; Zn, 213.856 nm; Cd, 228.802 nm; Cu, 324.803 nm) under conventional conditions following an already stablished method (Bongers, et al. 1988). Molecular weights were determined by ESI-MS, in a MicroTof-Q instrument (Bruker Daltonics GmbH, Bremen, Germany) connected to a Series 1100 HPLC pump (Agilent Technologies) controlled by the Compass Software. The instrument was calibrated with ESI-L Low Concentration Turning Mix (Agilent Technologies, Santa Clara, CA, USA). Metallated forms were detected under native conditions: 20 µL of sample injected through a PEEK tube at 30-50 µL·min⁻¹ in a 3.5-5.0 kV capillary-counter voltage, at 90-110°C of desolvation temperature, and with dry gas at 6 L·min⁻¹. The spectra were recorded between a m/z range from 800 to 3000. The liquid carrier was a 90:10 mixture of 15 mM ammonium acetate and acetonitrile at pH 7.0. All molecular masses were calculated according to the bibliography (Fabris, et al. 1996).

5. SUPPLEMENTARY MATERIAL

Supplementary data are available at Molecular Biology and Evolution online.
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7. DATA AVAILABILITY

The data underlying this article are available in NCBI (https://www.ncbi.nlm.nih.gov/) and Octopus (http://octopus.obs-vf.fr/public/botryllus/blastbotryllus.php) databases. The amino acid sequences and the accession numbers of the MTs used in this work are available in supplementary Table S1, Supplementary Material online.

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Table 1. Cysteine motifs in 11/12-Cys domains from different animal taxa

| Taxon               | 11/12-Cys domains | Cysteine motifs | Metal ions |
|---------------------|-------------------|----------------|------------|
| ascidians/thaliaceans| full length MT    | [CxC]x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CC) | 4 (this work) |
| appendicularians    | 12C domain        | CxCx(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CC) | nd^a |
| vertebrates         | α-domain^b        | CxCx(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CC) | 4 (Stillman, et al. 1987) |
| cephalochordates    | 11C domain        | [CxC]x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CC) | nd |
| mollusks            | α-domain^c        | [CxC]x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CC) | 4 (Digilio, et al. 2009) |
| echinoderms         | α-domain^d        | CxCx(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CC) | 4 (Tomas, et al. 2013) |
| insects             | MTnB/C/D          | CxCx(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CxC)x(CC) | 4 (Egli, et al. 2006) |

^a nd: not determined
^b Notice that although vertebrate, echinoderm and mollusk MTs have α-domains, the origin and evolutionary relationship of these α-domains remains uncertain.

FIGURE LEGENDS

**Figure 1. Phylogenetic relationships in the Chordata phylum.** Regarding tunicates, traditional classifications divided this subphylum in three classes, Appendicularia (green background), Asciidiacea (yellow background) and Thaliacea (blue background), but recent analyses propose Thaliacea species (salps, doliolids and pyrosomes) are nested within the Asciidiacea class, closer to the Aplousobranchia and Phlebobranchia orders than to the Stolidobranchia order (Delsuc, et al. 2018; Kocot, et al. 2018). A schematic representation of the MTs of each taxonomic group, with a mono-, bi- or multi-modular organization with distinct domains (color coded) containing seven, nine, eleven or twelve cysteines, is showed at the right. See Figure 7 and text for additional details.

**Figure 2. Schematic representation of multimodular Stelydae MTs.** Internal duplications of a 12-Cys domain (grey box) generated multimodular MTs with variable number of repeats (R), from 2 to 9. The number of MTs with different number of repeats (R1-R9) is indicated (DgrMT7 =2 repeats; EtiMT8part ≥4 repeats; ScaMT6part and SsoMT8part ≥6 repeats; PmiMT10part, BleMT1 and BleMT2 ≥7 repeats; PauMT10part and PspMT7 ≥8 repeats; BscMT1, PmamiMT3 and PpoMT7 =9 repeats). See Figure S2 for further details.

**Figure 3.** Deconvoluted ESI-MS spectra of the Cd(II)-productions of ascidian CroMT1 (A), HroMT1(B), HroMT2(C) and +14 charge state ESI-MS spectra of the Zn(II)-production of BscMT1 (D), as well as the deconvoluted ESI-MS spectra of the Cd(II)-production of its fragment R4 BscMT1 (E). Notice that BscMT1 is capable to bind up to 36 divalent ions, which nicely match with a multi-modular organization made
of nine repeated domains \((12C)_9\), each one binding four divalent ions alike the mono-modular \((12C)\) MTs (A-C) or the \(R_4\) \((12C)\) domain (E).

**Figure 4.** Circular dichroism spectra of the Cd(II)-productions of CroMT1, HroMT1, HroMT2, BscMT1 and \(R_4\) BscMT1, all exhibiting very similar CD signals. The exciton coupling centered at ca. 250 nm confirms high robustness and compactness of the clusters formed.

**Figure 5.** Deconvoluted ESI-MS of the Cd(II)-production of *S. thompsoni* SthMT1 (A), SthMT2 (B), SthMT3 (C) and to SthMT4 (D). Thaliacean MTs are mono-modular proteins made of a single 12C domain capable to bind four divalent ions, identically to ascidian MTs.

**Figure 6.** Reconstruction of MT evolution in the Chordata phyla. The most parsimonious scenario would be that the ancestral chordate MT was a bimodular MT made of an N-terminal 11/12-Cys domain (in blue) and a C-terminal 9-Cys domain (in red), similar to the current cephalochordate MT1. In cephalochordates, this ancestral form was tandem duplicated. One copy lost the 11/12-Cys domain and expanded the 9-Cys domain, yielding the current bi/trimodular MT2 form. In vertebrates, the number of cysteines in each domain was readjusted to the current 9 Cys of the N-terminal \(\beta\)-domains, and 11 Cys of the C-terminal \(\alpha\) domain. Successive lineage-specific tandem duplications of this primeval vertebrate MT led to different MT types (MT1, MT2, MT3 and MT4) with a variable degree of multiplicity in different species. In tunicates, the ancestral form lost the C-terminal 9-Cys domain in ascidian/thaliacean lineages, leading a mono-modular. This MT was frequently duplicated generating a high degree of MT multiplicity in many ascidian and thaliacean species. In contrast, internal duplications led to the emergence of multi-modular MTs in the stelydae lineage. Finally, in the appendicularian clade, the ancestral MT appears to have been lost, and a new MT would have emerged de novo by the duplication of a 12-Cys domain (in green). During *O. dioica* evolution, the new bi-modular MT lost five Cys in the N-terminal domain, yielding a 7-Cys/12-Cys arrangement (as it is found in current OdiMT1). Next, this 7-Cys/12-Cys MT was internal duplicated generating multi-modular MTs (as it is OdiMT2).

**Figure 7.** Comparison of the arrangement of Cys motifs in different MT domains. The arrangement of the Cys motifs in the 11-Cys amino-terminal domain of
cephalochordate MT1s are somewhat similar to that in the vertebrate 9-Cys β domains and in the ascidian/thaliacean 12-Cys MTs (blue boxes), while the Cys motifs in the carboxyl-terminal domains of cephalochordate MT1s and MT2s (9C and 9C\textsubscript{Like} domains) are vaguely similar to those in vertebrate 11-Cys α domains (pink boxes). In contrast, the arrangement of the Cys motifs in the appendicularian MT1 seems totally different (green boxes).
Figure 2

| STELYDAE | 1 MT | 12C-R_1 | 12C-R_2 |
|----------|------|---------|---------|
| 1 MT     | 12C-R_1 | 12C-R_2 | 12C-R_3 | 12C-R_4 |
| 2 MTs    | 12C-R_1 | 12C-R_2 | 12C-R_3 | 12C-R_4 | 12C-R_5 | 12C-R_6 |
| 3 MTs    | 12C-R_1 | 12C-R_2 | 12C-R_3 | 12C-R_4 | 12C-R_5 | 12C-R_6 | 12C-R_7 |
| 2 MTs    | 12C-R_1 | 12C-R_2 | 12C-R_3 | 12C-R_4 | 12C-R_5 | 12C-R_6 | 12C-R_7 | 12C-R_8 |
| 3 MTs    | 12C-R_1 | 12C-R_2 | 12C-R_3 | 12C-R_4 | 12C-R_5 | 12C-R_6 | 12C-R_7 | 12C-R_8 | 12C-R_9 |
Figure 4

The graph shows the molar ellipticity ($\Delta \varepsilon$ in M$^{-1}$cm$^{-1}$) as a function of wavelength ($\lambda$ in nm). The different lines represent the following MT proteins:

- **Cd-CroMT1**
- **Cd-HroMT1**
- **Cd-HroMT2**
- **Cd-BscMT1**
- **Cd-$R_4$ BscMT1**

The data points and curves vary across different wavelengths, indicating distinct spectral characteristics for each protein variant.
| Domain Description | Formula | Image |
|--------------------|---------|-------|
| 9C domain          | $\text{CCx}_n\text{Cx}_n\text{Cx}_n\text{C}_x\text{n}[\text{C x C}]\text{xCC}$ | ![9C domain image] |
| 11C domain         | $\text{[CxC]}_n\text{[CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{Cx}]$ | ![11C domain image] |
| 9C-like domain     | $\text{CCx}_n\text{Cx}_n\text{Ix}_n\text{C} \times [\text{C x C}]\text{xCC}$ | ![9C-like domain image] |
| β-9C domain        | $\text{[CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{Cx}]$ | ![β-9C domain image] |
| α-11C domain       | $\text{CCxCCx}_n\text{Cx}_n\text{Cx}_n\text{[C x C]}\text{xn}[\text{Cx}]\text{xn}[\text{Cx}]\text{xCC}$ | ![α-11C domain image] |
| 12C domain         | $\text{C}_x\text{n}[\text{CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{Cx}]\text{xn}[\text{Cx}]\text{xn}[\text{Cx}]\text{xn}[\text{Cx}]\text{[Cx]}$ | ![12C domain image] |
| t-12C domain       | $\text{C}_x\text{n}[\text{CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{Cx}]\text{xn}[\text{Cx}]\text{xn}[\text{Cx}]\text{xn}[\text{Cx}]\text{xCC}$ | ![t-12C domain image] |
| 12C domain         | $\text{C}_x\text{n}[\text{CxC]}_x\text{n}[\text{CxC]}_x\text{n}[\text{Cx}]\text{xn}[\text{Cx}]\text{xn}[\text{Cx}]\text{xn}[\text{Cx}]\text{[Cx]}$ | ![12C domain image] |

**Figure 7**

- Cephalochordate MT1
- Cephalochordate MT2
- Vertebrate MT
- Ascidian/thaliaceous MT
- Appendicularian MT1