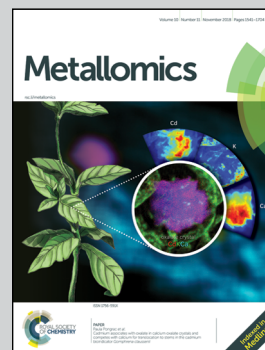


Showcasing research from Professor Ricard Albalat's laboratory, Department of Genetics, Microbiology and Statistics, University of Barcelona, Catalonia.

Metallothioneins of the urochordate *Oikopleura dioica* have Cys-rich tandem repeats, large size and cadmium-binding preference

We have characterized two novel metallothionein (MT) genes in the planktonic animal *Oikopleura dioica*, revealing a particular repeat structure that have led to one of them to encode for the longest MT reported to date for any living being. The image, provided by Alfonso Ferrández-Rodán, shows a living *O. dioica* specimen inside its mucous 'house'. The trunk of the animal is visible at the top-center of the image surrounded by the house in false green and grey.

As featured in:



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Metallothioneins of the urochordate *Oikopleura dioica* have Cys-rich tandem repeats, large size and cadmium-binding preference†

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The increasing levels of heavy metals derived from human activity are poisoning marine environments, threatening zooplankton and ocean food webs. To protect themselves from the harmful effects of heavy metals, living beings have different physiological mechanisms, one of which is based on metallothioneins (MTs), a group of small cysteine-rich proteins that can bind heavy metals counteracting their toxicity. The MT system of urochordate appendicularians, an ecologically relevant component of the zooplankton, remained, however, unknown. In this work, we have characterized the MTs of the appendicularian species *Oikopleura dioica*, revealing that *O. dioica* has two MT genes, named *OdMT1* and *OdMT2*, which encode for Cys-rich proteins, the former with 72 amino acids comparable with the small size MTs of other organisms, but the second with 399 amino acids representing the longest MT reported to date for any living being. Sequence analysis revealed that *OdMT2* gene arose from a duplication of an ancestral *OdMT1* gene followed by up to six tandem duplications of an ancestral repeat unit (RU) in the current *OdMT2* gene. Interestingly, each RU contained, in turn, an internal repeat of a 7-Cys subunit (X₃CX₃CX₂CX₂CX₃₋₆CX₂CXCX), which is repeated up to 12 times in *OdMT2*. Finally, ICP-AES analyses of heterologously expressed *OdMT* proteins showed that both MTs were capable to form metal-complexes, with preference for cadmium ions. Collectively, our results provide the first characterization of the MT system in an appendicularian species as an initial step to understand the zooplankton response to metal toxicity and other environmental stress situations.

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Significance to metallomics

The analysis of the metallothioneins of *Oikopleura dioica* is important for understanding the evolution and function of the detoxification mechanisms that this zooplankton species might use to counteract the harmful effects of heavy metal exposure. This knowledge is fundamental to estimate the potential impact that an increase of heavy metal amounts may have on marine ecosystems.

1. Introduction

Heavy metals such as zinc, iron or copper are essential for several biological processes, but toxic at high concentrations, while others such as cadmium, mercury or lead are highly poisonous even at low concentrations. Living beings have

different physiological mechanisms to control the homeostasis of essential metals as well as to counteract the harmful effects of the non-essential ones. One of these mechanisms is based on metallothioneins (MTs), a group of metal-binding proteins originally discovered in 1957 in the horse kidney cortex,¹ and classically considered a diverse family of Cys-rich ($\approx 30\%$) and low molecular weight (<60 amino acids) proteins found in almost all organisms (reviewed in ref. 2 and 3). The amino acid sequence of different MTs is highly heterogeneous, particularly when MTs of distantly related taxa are compared. In such comparisons, sequence similarity appears restricted to an overall “Cys abundance” with distinctive cysteine motifs (*i.e.* CXC, CC and CCC), whereas the length of the protein, the number

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and distribution of cysteines and the intercalating residues are largely variable from one taxon to another.² Vertebrate MT sequences, for instance, can be reliably aligned within the subphylum, but they are barely comparable with other MT sequences, even from their closest chordate relatives, amphioxus cephalochordates or ascidian urochordates.^{4,5}

From a structural perspective, MTs have been extensively analyzed showing that they are able to coordinate a number of heavy metal ions through the formation of metal–thiolate bonds.^{6,7} Depending on their metal-binding preferences, MTs can be classified in a stepwise gradation from extreme Zn/Cd-thioneins to extreme Cu-thioneins.^{8,9} The metal-binding preference of a given MT cannot be, however, directly predicted from its amino acid sequence, and metal-binding assays have to be performed to classify newly discovered forms.

From a physiological perspective, MTs are activated for controlling metal homeostasis and detoxification roles, but also for radical scavenging, oxidative stress protection or anti-apoptotic defense (reviewed in ref. 6). The presence of Metal Responsive Elements (MRE) and different response elements for transcription factors (TF) involved in stress response (e.g. AREs, HSE, StRE) in the promoter regions has been associated to the transcriptional activation of MT genes in response to heavy metals and other stress situations (reviewed in ref. 10).

In the last century, what has been called the Anthropocene, human activities such as those related to the industries of mining, metal plating, fertilizer, paper, pesticides or batteries, are increasing the amount of heavy metals in aquatic environments,^{11,12} becoming a serious problem for coastal and marine ecosystems.¹³ In marine ecosystems, Appendicularians (a.k.a. Larvaceans; phylum Chordata, subphylum Urochordata) represent the second most abundant group of mesozooplankton grazers and an important component of food for fish and zooplankton species.¹⁴ Appendicularians are also ecologically relevant because they contribute to the vertical transport of carbon to deep ocean through the rapid sinking of fecal pellets and discarded houses.^{15–17} Among Appendicularians, *Oikopleura dioica* is the most studied species, and it is becoming a new laboratory model for comparative genetic and genomic analyses,^{18–22} developmental biology studies,^{23–30} as well as ecological and toxicological investigations.^{31–34}

The ecological relevance of Appendicularians together with the potential of *O. dioica* as an experimental model^{35–38} prompted us to survey the MTs of *O. dioica* to characterize the MT system of an Appendicularian species. Here we report the identification of two MT genes in *O. dioica*, named *OdMT1* and *OdMT2*, and the analysis of their gene structures and promoter regions. *OdMT1* encode for a Cys-rich protein of 72 amino acids (theoretical molecular weight 7645 Da) comparable with the small size MTs of other organisms (human MT1 theoretical molecular weight 6120 Da), while *OdMT2* encode for a Cys-rich protein of 399 amino acids (theoretical molecular weight 42 716 Da) representing the longest MT reported to date for any living being. Interestingly, sequence analysis reveals that *OdMT2* gene arose from a duplication of an ancestral *OdMT1* gene followed by up to six tandem duplications of an ancestral repeat unit (RU) in the current *OdMT2* gene.

OdMT promoters appear rich in stress-responsive elements, including binding sites for MREF, YAP1/CREB, RXR/PPAR and HSF transcription factors, envisaging a role of MT genes in adverse environmental conditions. Finally, heterologously expressed *OdMT1* and *OdMT2* proteins showed that they are capable to form metal complexes, with preference for cadmium ions. Overall, these results pave the way for a better understanding of the homeostasis of the physiological metals in *O. dioica*, as well as the genetic mechanisms that Appendicularians might use against the harmful effects of heavy metal exposure.

2. Material and methods

2.1. Genome survey of *O. dioica* MT genes

To identify *O. dioica* MT homologs we made low-restrictive tblastn searches³⁹ in the genome database of *O. dioica* (Oikobase, <http://oikobase.biology.uiowa.edu/Oiko>) with a high value of expect threshold (1000) and no filtering for low complexity sequences, using the two only available MTs of Urochordate ascidians *Ciona robusta* and *Herdmania curvata* MTs (accession numbers ACN32211 and AY314949, respectively), as well as Cephalochordate and Vertebrate MTs as queries.^{4,5} Gene annotations were manually reviewed and corrected based on ESTs availability: EST FP794470.1 for *OdMT1* and a consensus sequence derived from the assembly of 18 SRA sequences for *OdMT2* (Fig. S1 and Table S1, ESI[†]).

2.2. Promoter analysis and stress-responsive elements

The analysis of putative transcription factor binding sites for *OdMT1* and *OdMT2* genes was performed by means of the MatInspector 8.4.1 program with default parameters, using the MatBase 11.0 from the Genomatix software suite.⁴⁰ The MatBase 11.0 includes 2029 weight matrices of 482 families, representing binding sites descriptions of more than 11 000 TF. We restricted our analysis to a subset of the 33 families that included TF involved in metal- or stress-response in animals, plants or fungi (Table S3, ESI[†]). A 1000-bp region upstream of predicted CDS of both *OdMT* genes was selected for the promoter analysis.

2.3. Cloning of *OdMT1* and *OdMT2* for recombinant expression

A full-length synthetic cDNA for predicted *OdMT1* gene was synthesized by Integrated DNA Technologies Company (Coralville, IA, USA), while for technical limitations, the cDNA of *OdMT2* was split in two fragments of 610 nt (5'-fragment 1) and 660 nt (3'-fragment 2), overlapping 52 nt among them. *Bam*HI and *Xho*I restriction sites and 6–7 additional 5'-nucleotides were added to both *OdMT* cDNA ends to facilitate the cloning processes. The synthetic cDNA for *OdMT1* was PCR amplified –94 °C 5 minutes (min); 25 cycles of 94 °C 30 seconds (s), 55 °C 30 s and 72 °C 30 s; and 72 °C 7 min – with specific primers (5'GGGGATCCATGGATCCGGTTTGCTCTTCCGCTG3' and 5'GGGCTCGAGTTATCCGCTGTGCTGGTCCGGCAG3') using Expand High Fidelity PCR system[®] (Roche). The full-length



synthetic cDNA for *OdMT2* was reconstructed in two steps by adapting previous methods for assembling overlapping DNA fragments.^{41–44} In brief, in the first step, the two *OdMT2* fragments were denaturalized at 98 °C for 3 min, annealed at 80 °C for 1 min, and extended at 94 °C for 30 s. Ten cycles of annealing and extension were repeated to generate a single full-length cDNA molecule from the assembly of the two overlapping fragments. In the second step, the resulting cDNA for *OdMT2* was PCR-amplified – 15 cycles of 76 °C 30 s and 94 °C 30 s; and 76 °C 5 min – with specific end-terminal primers (5'GGGGGATCCATGGAAGTAAAACGACC3' and 5'GGGCTCGAGTTAACAGCATTTTTTGG3') using Phusion High Fidelity DNA polymerase (New England Biolabs, Thermo Scientific). *OdMT1* and *OdMT2* PCR products were *Bam*HI/*Xho*I-digested and cloned into the *E. coli* pGEX-4T-1 expression vector (GE Healthcare) with the DNA Ligation kit 2.1 (Takara Bio Inc.), and transformed into *E. coli* Dh5 α strain. Plasmid DNA was purified from bacteria using the GeneElute™ Plasmid Miniprep Kit (Sigma-Aldrich), screened for insert presence by digestion with *Sca*I enzyme, 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). DNA from each recombinant *OdMT*-pGEX plasmid was used to transform *E. coli* BL21 strain, a protease deficient strain used for heterologous protein expression.

2.4 Production of recombinant metal-*OdMT1* and -*OdMT2* complexes

For heterologous protein production 500 mL of LB medium with 100 $\mu\text{g mL}^{-1}$ ampicillin were inoculated with *E. coli* BL21 cells transformed with the *OdMT1* or *OdMT2* recombinant plasmids. After overnight growth at 37 °C/250 rpm, the cultures were used to inoculate 5 L of fresh LB-100 $\mu\text{g mL}^{-1}$ 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 ZnCl_2 (300 μM), CdCl_2 (300 μM) or CuSO_4 (500 μM) in order to generate metal-*OdMT* complexes. Cells were harvested by centrifugation for 5 min at 9100g (7700 rpm), and bacterial pellets were suspended in 125 mL of ice-cold PBS (1.4 M NaCl, 27 mM KCl, 101 mM Na_2HPO_4 , 18 mM KH_2PO_4 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 s, and then centrifuged for 40 min at 17 200g (12 000 rpm) and 4 °C.

2.5 Purification of recombinant metal-*OdMT* complexes

Protein extracts containing GST-*OdMT1* or GST-*OdMT2* fusion proteins were incubated with glutathione sepharose beads (GE Healthcare) for 1 h at room temperature with gentle rotation. GST-*OdMT* fusion proteins bound to the sepharose beads were washed with 30 mL of cold 1 \times PBS bubbled with argon to prevent oxidation. After three washes, GST-MT fusion proteins were digested with thrombin (GE Healthcare, 25 U L^{-1} of culture) overnight at 17 °C, thus enabling separation of the metal-*OdMT* complexes from the GST that remained bound to the

spharose matrix. The eluted metal-*OdMT* complexes were concentrated with a 3 kDa Centriprep Low Concentrator (Amicon, Merck), and fractionated on a Superdex-75 FPLC column (GE Healthcare) equilibrated with 20 mM Tris-HCl, pH 7.0 for *OdMT1*, and with fresh 50 mM ammonium acetate, pH 7.0 for *OdMT2*, and run at 0.8 mL min^{-1} . The protein-containing fractions, identified by their absorbance at 254 nm, were pooled and stored at -80 °C until use.

2.6 Analysis of metal-*OdMT* complexes

Metal-*OdMT* complexes were analyzed by different techniques: inductively coupled plasma atomic emission spectroscopy (ICP-AES) for protein quantification and element composition (S, Zn, Cd and Cu) on a Polyscan 61E spectrometer (Thermo Jarrell Ash Corporation, Franklin, MA, USA) at appropriate wavelengths (S, 182.040 nm; Zn, 213.856 nm; Cd, 228.802 nm; Cu, 324.803 nm) under conventional (dilution with 2% HNO_3 (v/v)) conditions.⁴⁵ *OdMT* concentrations in the recombinant preparations were calculated from sulfur measurements, assuming the only contribution to their S content was that made by the *OdMT* peptides.

Electrospray ionization mass spectrometry with a time-of-flight analyzer (ESI-TOF MS) was used to determine the molecular mass of the species formed, using a Micro Tof-Q Instrument (Bruker Daltonics GmbH, Bremen, Germany) calibrated with ESI-L Low Concentration Tuning Mix (Agilent Technologies, Santa Clara, CA, USA), interfaced with a Series 1100 HPLC pump (Agilent Technologies) equipped with an autosampler, both controlled by the Compass Software. The experimental conditions for analyzing the proteins were as follows: 10–20 μL of the sample were injected at 30–50 $\mu\text{L min}^{-1}$ at 3.5–5.0 kV capillary-counter electrode voltage, 90–110 °C desolvation temperature, and dry gas at 6 L min^{-1} . Spectra were collected throughout an *m/z* range from 800 to 2000. The liquid carrier was a 90:10 mixture of 15 mM ammonium acetate and acetonitrile, pH 7.0. For the analysis at acidic pH the conditions used were the same as those used in the analysis of the divalent metals, except in the composition of the carrier liquid, which in this case was a 95:5 mixture of formic acid and acetonitrile at pH 2.4. All samples were injected in duplicates to ensure reproducibility. In all cases, molecular masses were calculated according to the reported method.⁴⁶

3. Results and discussion

3.1 Identification, structural features and evolutionary origin of *O. dioica* MTs

We conducted an exhaustive survey of *MT* genes in the *O. dioica* genome.^{19,21} Because standard blast strategies³⁹ were unsuccessful for identifying putative *MT* genes in *O. dioica* sequence databases, we used low-restrictive tblastn searches using *C. robusta* and *H. curvata* MTs as well as Cephalochordate and Vertebrate MTs as query sequences. We manually inspected all the retrieved sequences for Cys-rich ORFs with MT-distinctive Cys arrangements (*i.e.* CXC, CXXC, CC and CCC) as structural criteria for identification of new putative *MT*-coding genes.



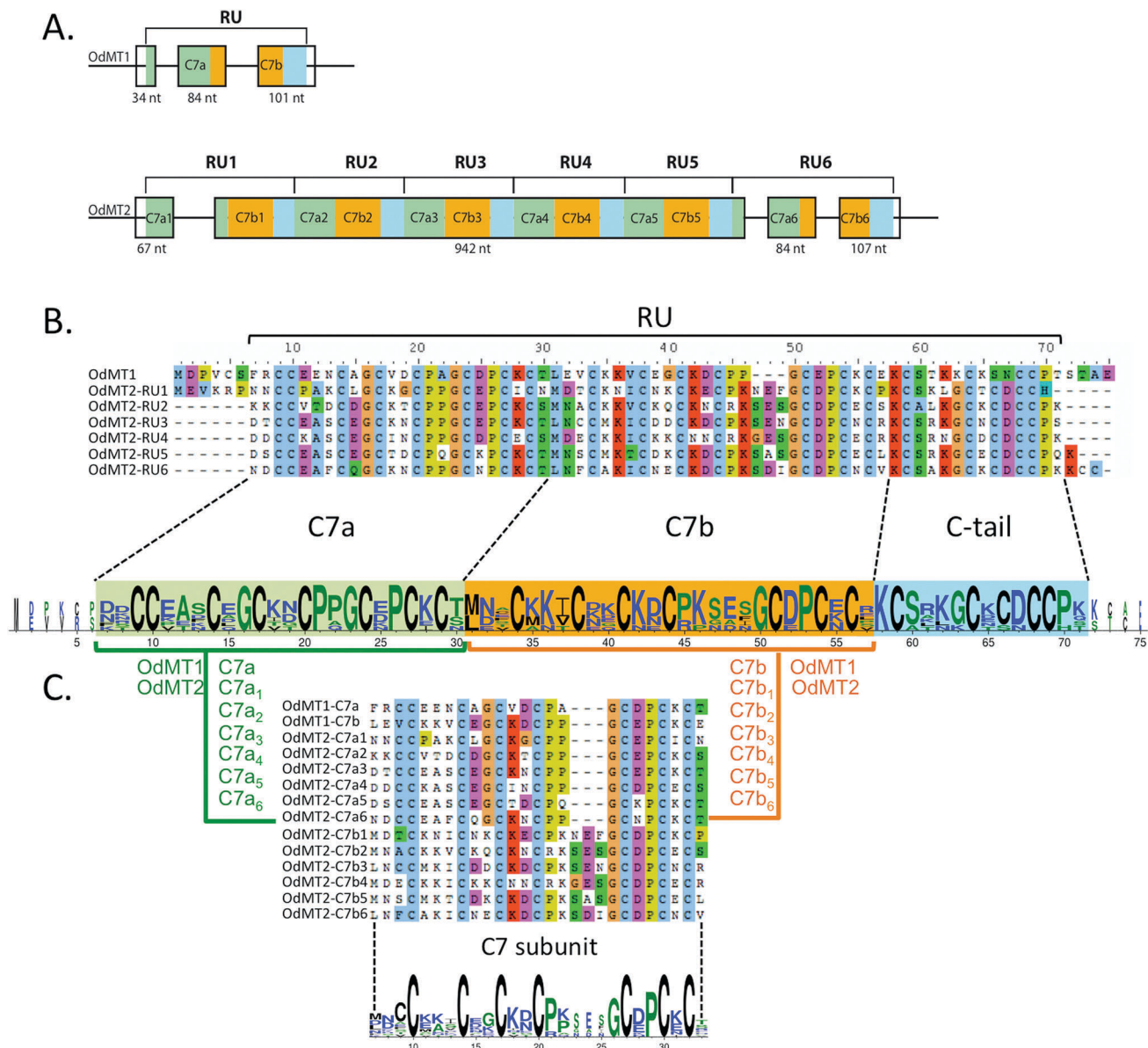


Fig. 1 *O. dioica* metallothioneins. (A) Schematic representation of the exon/intron structure of *OdMT1* and *OdMT2* genes. *OdMT1* is split in three exons and encodes for a protein made of a single repeat unit (RU) with an internal repeat of a C7 subunit – C7a (green box) and C7b (orange box) – followed by a C-terminal tail (blue box). *OdMT2* is split in four exons and encodes for a protein made of 6 RU (RU1 to RU6) with the same C7a/C7b/C-tail structure. (B) Amino acid alignment of *OdMT1* and RU1–RU6 of *OdMT2*, and graphical representation using WebLogo ³⁷ highlighting the 20 conserved Cys in each RU. Each RU contains a repetition of a C7 subunit (C7a in green background, and C7b in orange background) followed by a C-terminal tail (in blue background). (C) Comparison of C7a and C7b defines the C7 subunit as a 24–27 amino acid sequence with 7 Cys ($X_3CX_3CX_2CX_2CX_{3-6}CX_2CX_3CX$). In the WebLogo, the overall height of the stack indicates the conservation degree at a given position, while the height of symbols within the stack indicates the relative frequency of each amino acid at that position. Color code represents amino acid hydrophobicity scale: hydrophobic amino acids (YVMCLFIW) in black, neutral (SGHTAP) in green, and hydrophilic (RKDENQ) in blue.

Two *O. dioica* genomic sequences satisfied the criteria, one in the scaffold 50 and another in the scaffold 16, which we named *OdMT1* and *OdMT2*, respectively. *OdMT1* gene spanned 317 nt and its coding region (CDS) was organized in 3 exons of 34 nt, 84 nt and 101 nt (Fig. 1A and Fig. S2, ESI[†]). *OdMT2* gene extended over 1359 nt, and its CDS was organized in three small exons (exons 1, 3 and 4) of 67 nt, 84 nt and 107 nt, respectively, and one large exon (exon 2) of 942 nt (Fig. 1A and Fig. S2, ESI[†]).

OdMT1 encoded a 72 amino acid protein with 20 Cys (28%), and *OdMT2*, in contrast, encoded a much larger protein, with 399 amino acids containing 123 Cys (31%), being to our knowledge the longest MT so far described in any living being. A detail comparison of *OdMT1* and *OdMT2* sequences revealed that *OdMT2* was made of 6 direct tandem repeat units (RU) of approximately 65 amino acids, each RU resembling to *OdMT1* (Fig. 1A and B and Fig. S2, ESI[†]). Moreover, close inspection revealed that each RU was made of an internal repeat of 24–27



amino acids with 7 conserved Cys (C7a and C7b subunits: X₂CX₃CX₂CX₂CX₃₋₆CX₂CXCX; Fig. 1B and C) followed by a C-terminal tail of 14 amino acids with 5 additional Cys residues. The fact that OdMT1 corresponded to a single RU whereas OdMT2 had 6 RUs suggested that the latter might have been originated by duplication of an ancestral *OdMT1* gene with one RU followed by successive tandem duplications of the primeval RU. These results indicated a modular and step-wise evolution of OdMTs, in which the unit of expansion was a RU made of two C7 subunits, likely related with functional and structural constraints that would link the number of RUs and C7 subunits to the metal-binding capacity of OdMTs. Interestingly, a modular structure based on different C7 subunits had been found in several long fungal MTs, that is, in *Cryptococcus neoformans* CnMT1 and CnMT2⁴⁷ and in *Tremella mesenterica* TmMT,⁴⁸ although it had never been reported in multicellular eukaryotes, and never to the level reached by *O. dioica* MTs. Tandem amplifications of C7 subunits yielding MTs with high metal-binding capacity had been associated with biological adaptations, as virulence factors in the pathogenic fungus *C. neoformans*,⁴⁷ or for Cu-providing to lignin-metabolizing enzymes in saprophytic fungus living in decaying woods as *T. mesenterica*.⁴⁸ The selective advantage that MTs with high metal-binding capacity might provide to a marine animal as *O. dioica* is still unknown, but it has been suggested that the evolution of multi-domain MTs in some animal species (e.g. *Littorina littorea* and *Pomatias elegans* with 3-domain MTs⁴⁹⁻⁵¹ *Crassostrea virginica* with 4-domain MTs,⁵² and *Alinda biplicata* with 10-domain MTs⁵³) might have improved their response to metal stress and adverse environmental conditions, offering a selective advantage to organisms that experience a greater exposure to metals due to their ecological niche.^{50,52}

3.2. Promoter prediction analysis

Metals, stress and hormones regulate transcription of *MT* genes (reviewed in ref. 10). In animals, *MT* expression can be activated in response to metals by MRE-binding transcription factor 1 (MTF-1) that recognizes Metal Responsive Elements (MRE) present in *MT* promoters.⁵⁴ MRE have been experimentally and bioinformatically identified in proximal *MT* promoters of most metazoans, from insects and mollusks to fish and mammals⁵⁵⁻⁶¹ (reviewed in ref. 62). We surveyed for MRE in 1 kb upstream sequence from the predicted CDS of both *OdMT1* and *OdMT2* genes using MatInspector 8.4.1 program but, unexpectedly, we did not find any putative MRE sequence in their promoter regions. In agreement with the absence of MRE sequences, a genome survey for a *O. dioica* homolog of the *MTF-1* gene was also unsuccessful, making thereby unlikely that these elements – MRE sequences and MTF-1 proteins – contributed to the transcriptional regulation of *MT* genes in this species. In contrast, bioinformatic analysis predicted the presence MREF-binding sequences (ACE1/AMT1) in the promoters of *OdMT1* and *OdMT2* genes (Fig. 2A and Fig. S2, ESI[†]), which are known to confer metal response to fungal metallothioneins *CUP1* and *CRS5* genes. Binding of metal regulatory elements factors (MREF) such as CUP2 (a.k.a ACE1 or AMT1

transcription factor) to ACE1/AMT1 sequences regulates *CUP1* and *CRS5* in response to copper levels,^{63,64} as well as they regulate fungal *copper-zinc superoxide dismutase* (*Cu/ZnSOD1*) genes.⁶⁵ Although CUP2 orthologous have not been identified outside fungal species, ACE1/AMT1 sequences have been related to transcriptional regulation of *Cu/ZnSOD1* genes in maize⁶⁶ and *Arabidopsis*.⁶⁷ Therefore, we also analyzed the promoters (1 kb upstream sequence from the CDS) of the five *SOD1* genes predicted in *O. dioica* genome, and interestingly, we identified MREF (ACE1/AMT1) sequences in four of them (Fig. 2A), suggesting that as described in fungi, *MT* and *SOD* genes shared a common transcriptional response in *O. dioica*.

Because MTs are activated in many stress circumstances, we extended our promoter analyses to 33 families of transcription factors with members involved in stress response in animals, plants and fungi (Table S2, ESI[†]). Interestingly, in addition to MRFE sequences, we discovered that *OdMT1* and *OdMT2* promoters shared binding sites for many stress response elements (Table S3, ESI[†]). First, *OdMT* promoters included binding sites for the family of fungal YAP1, most of them overlapping with sites for vertebrate (CREB) bZIP transcription factors (Fig. 2B, Fig. S2 and Table S3, ESI[†]). Fungal YAP1 transcription factor belongs to the bZIP (basic leucine zipper factors) family, and it is required for cadmium and oxidative stress tolerance in yeast.⁶⁸ Vertebrate bZIP transcription factors of the CREB family (e.g. CREB, ATF-1, XBP1, E4BP4) might act also as activators of vertebrate *MT* genes,⁶⁹ and CRE binding sites have been identified in the promoter of earthworm *MT* genes.^{70,71} Second, it has also been reported that transcription factors of the nuclear receptor family such as the steroid receptors, retinoid receptors and orphan receptors might be involved either directly (e.g. GR;⁷² PPAR^{56,57}) or indirectly (e.g. RXR⁷³) in the transcriptional regulation of *MT* expression. Our analysis of *OdMT1* and *OdMT2* genes revealed several binding sites for nuclear receptors, mainly for RXR/PPAR, in the promoter region (Fig. 2B, Fig. S2 and Table S3, ESI[†]). Third, it has been demonstrated that heat shock stress also induces *MT* expression through the binding of HSF to the heat shock elements (HSE),^{74,75} which appear remarkably abundant in some *MT* promoters.⁵⁷ Our computational analysis revealed a preponderant abundance of putative HSF binding sites in the *OdMT1* and *OdMT2* promoters (Fig. 2B, Fig. S2 and Table S3, ESI[†]).

Overall, the promoter region of *OdMT1* and *OdMT2* genes did not appear to contain MRE sequences, which are a common feature of metazoans *MT* genes, but they had a MREF conserved motif, which is typical of fungal *MT* genes.^{63,64} The presence of a MREF in the *OdMT* promoters was shared by the promoters of *O. dioica* *Cu/ZnSOD1* genes, and although the functionality of these sites needs to be experimentally demonstrated, they suggested a coordinated transcriptional activation of both (*MT* and *SOD*) stress-activated genes. *OdMT* promoters appeared, indeed, rich in stress-responsive elements, including binding sites for YAP1/CREB, RXR/PPAR and HSF transcription factors, envisaging a role of *MT* genes in adverse environmental conditions.



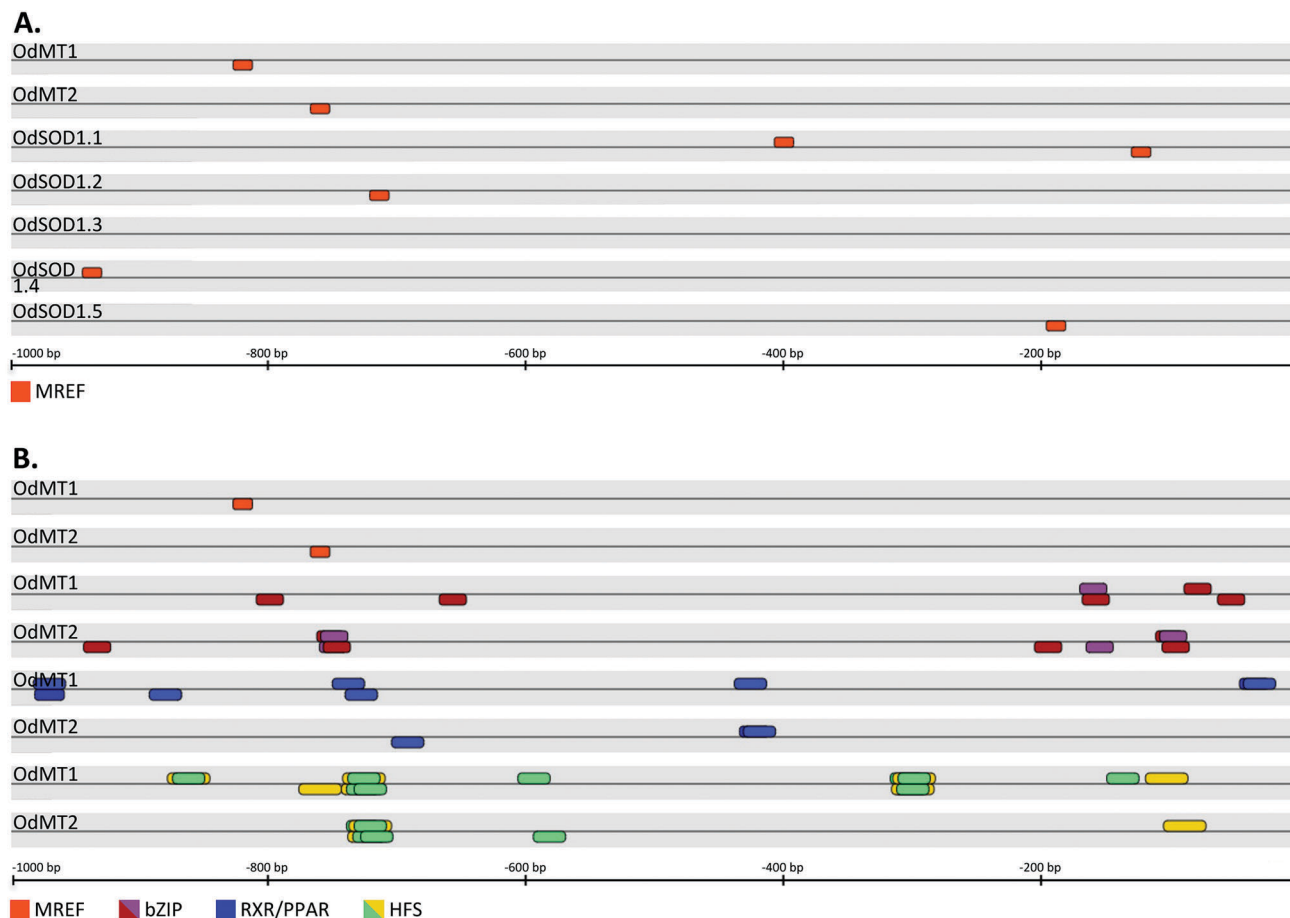


Fig. 2 Schematic representation of putative binding sites for transcription factors in the regulatory regions (1.0 kb upstream of CDS) of *O. dioica* genes. (A) MREF binding sites (orange boxes) were predicted in the *OdMT1* and *OdMT2* promoters as well as in the promoters of four out of five *OdSOD1* genes. (B) In addition of the MREF elements, several stress-response elements were predicted in *OdMT1* and *OdMT2* promoters, including those for bZIP transcription factors such as YAP1 (violet boxes) and CREB (purple boxes), for RXR/PPAR (blue boxes) and for animals or yeast HSF (green and yellow boxes, respectively). A box above the lines indicates that the binding site was in the plus DNA strand, whereas a box below indicates that it was in the minus strand (see Fig. S2 and Table S3 for additional details, ESI[†]).

3.3. Metal-binding capacity of *OdMT1* and *OdMT2*

In order to verify the MT nature of the two *O. dioica* MTs, and to evaluate their metal-binding affinity and capacity, we studied the formation of metal–MT complexes of *OdMT1* and *OdMT2* proteins heterologously expressed in *E. coli* and grown in medium supplemented with copper, cadmium or zinc salts. Metal–*OdMT1* and –*OdMT2* complexes were purified and analyzed by ICP-AES and ESI-MS. Acidification of the Zn–*OdMT* complexes yielded the corresponding apo-forms, with molecular masses of 7789 Da for *OdMT1* and 42 861 Da for *OdMT2* (Fig. 3A and B), fully concordant with the calculated average theoretical values for the synthesized products (7789.10 Da and 42860.48 Da, respectively; notice that recombinant proteins had two additional amino acids at N-terminus). This confirmed both the identity and purity of the recombinant proteins.

The ICP-AES analyses showed that both recombinant *OdMT1* and *OdMT2* proteins could form metal complexes, principally with cadmium ions (9.7 and 55.0 Cd/protein ratios for *OdMT1* and *OdMT2*, respectively; Table 1). ESI-MS analysis

of the metal–*OdMT1* species formed allowed us to observe the formation of a variety of species in the productions in Zn- and Cu-enriched media: from Zn₄- to Zn₇-*OdMT1* (Fig. 3C), and from Cu₈- to Cu₁₄-*OdMT1* (Fig. 3E), respectively. In contrast, the production in Cd-enriched media rendered a single Cd₇-*OdMT1* species, indicative of the formation of a highly favored species (Fig. 3D). Both ICP-AES and ESI-MS results indicated, therefore, that *OdMT1* exhibits a clear preference for coordinating Cd²⁺. In agreement with the Cd preference, the yield of recombinant *OdMT1* produced in Cd-media was higher than in Zn- or Cu-enriched media (Table 1) because metal binding might be contributing to stabilize recombinantly expressed protein. Regarding *OdMT2*, the high molecular weight of *OdMT2* and its complex structure made of repeated modules may be responsible for the low concentration of the samples recovered from the metal–*OdMT2* productions, even in the presence of Cd, impairing to obtain valid ESI-MS data of the metallated species. However, ICP data (Table 1), together with the fact that the yield of *OdMT2* production was higher in Cd-enriched



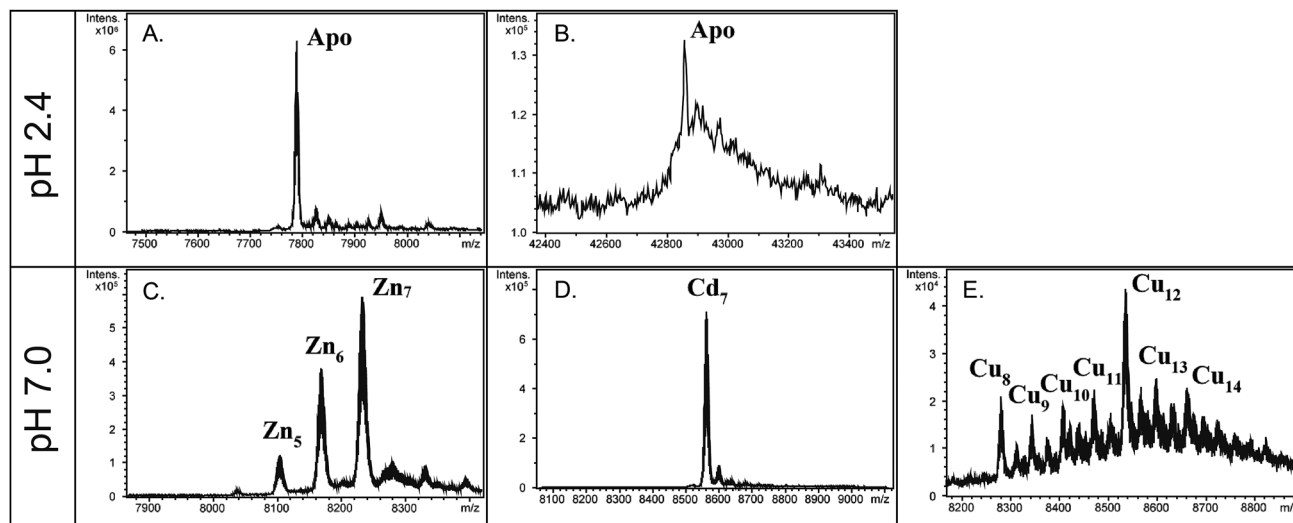


Fig. 3 Deconvoluted ESI-MS spectra of recombinant OdMT productions with different metal-enriched media. The apo-OdMT1 (A) and apo-OdMT2 (B) were recorded from the Zn- or Cd-productions, respectively, analyzed at acidic pH. Metal-OdMT1 species were recorded from the Zn-, Cd- or Cu- productions (C, D and E, respectively) at neutral pH.

Table 1 Protein concentration and metal content in metal-OdMT complexes

Metal-OdMT complex	Protein concentration (μM)	Metal/protein ratio
Zn-OdMT1	12	6.4 (Zn)
Cd-OdMT1	34	9.7 (Cd)
Cu-OdMT1	12	15.3 (Cu)
Zn-OdMT2	< 1	LDL ^a
Cd-OdMT2	15	55.0 (Cd)
Cu-OdMT2	< 1	LDL

^a LDL: lower than detection limit.

media than in Zn- and Cu-media (Table 1 and Fig. S3, ESI[†]) indicated a preference for Cd²⁺, likewise it has been demonstrated for OdMT1. In fact, OdMT2 seems to bind up to 6 more times higher amounts of Cd than OdMT1 (Table 1) accordingly with its high Cys content.

4. Conclusions

In this work, we have identified two *MT* genes in the chordate *O. dioica* revealing a particular modular structure of the encoded proteins based on repeat units of two C7 subunits. The structure of these chordate MTs together with data from fungal^{47,48} and protozoan MTs^{76,77} suggest that during metallothionein evolution, diverse living beings with different *MT* sequences have used a similar modular and step-wise mechanism for generating long proteins with high metal-binding capacities. In addition, sequence analyses have revealed the presence of metal- and stress-response elements in the regulatory regions of both genes, while ICP-AES analyses have shown the capacity of both *O. dioica* proteins to form metal-MT complexes, with preference for cadmium ions. These results support the MT nature of the *O. dioica* genes and proteins, and suggest that they might play a role in the physiological response of this

animal against metal toxicity and other stress situations. Overall our data pave the way for better understanding the evolution and function of the detoxification mechanisms of zooplankton species, which might be crucial to counteract the increasing amounts of heavy metals in marine ecosystems.

Conflicts of interest

There are no conflicts to declare.

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