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Significance to metallomics

Crosstalk of the structural and zinc buffering properties of mammalian metallothionein-2*

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Metallothioneins (MTs), small cysteine-rich proteins, present in four major isoforms, are key proteins involved in zinc and copper homeostasis in mammals. To date, only one X-ray crystal structure of a MT has been solved. It demonstrates seven bivalent metal ions bound in two structurally independent domains with $M_{4}S_{11}(\alpha)$ and $M_{\pi}S_{9}$ (β) clusters. Recent discoveries indicate that Zn(μ) ions are bound with MT2 with the range from nano- to picomolar affinity, which determines its cellular zinc buffering properties that are demonstrated by the presence of partially Zn(11)-depleted MT2 species. These forms serve as Zn(11) donors or acceptors and are formed under varying cellular free Zn(u) concentrations. Due to the lack of appropriate methods, knowledge regarding the structure of partially-depleted metallothionein is lacking. Here, we describe the Zn(II) binding mechanism in human MT2 with high resolution with respect to particular Zn(II) binding sites, and provide structural insights into Zn(m)-depleted MT species. The results were obtained by the labelling of metal-free cysteine residues with iodoacetamide and subsequent top-down electrospray ionization analysis, MALDI MS, bottom-up nanoLC-MALDI-MS/MS approaches and molecular dynamics (MD) simulations. The results show that the α -domain is formed sequentially in the first stages, followed by the formation of the β -domain, although both processes overlap, which is in contrast to the widely investigated cadmium MT. Independent ZnS₄ cores are characteristic for early stages of domain formation and are clustered in later stages. However, Zn-S network rearrangement in the β -domain upon applying the seventh Zn(11) ion explains its lower affinity. Detailed analysis showed that the weakest Zn(11) ion associates with the β -domain by coordination to Cys21, which was also found to dissociate first in the presence of the apo-form of sorbitol dehydrogenase. We found that Zn(11) binding to the isolated β-domain differs significantly from the whole protein, which explains its previously observed different Zn(μ)-binding properties. MD results obtained for Zn(μ) binding to the whole protein and isolated β -domain are highly convergent with mass spectrometry data. This study provides a comprehensive overview of the crosstalk of structural and zinc buffering related-to-thermodynamics properties of partially metal-saturated mammalian MT2 and sheds more light on other MT proteins and zinc homeostasis.

The growing interest in metallomics within metal-dependent biological systems provides an opportunity to explore the structural and buffering properties of metallothioneins (MTs). The wide relevance of MTs is obvious due to their multiple biological functions such as participation in zinc and copper homeostasis. However, the MT metalloproteome is still not fully explored and is rather complicated *in vivo* or *in vitro* because of its highly dynamic structure, which lacks secondary structural elements and metamorphic behaviour. We have undertaken comprehensive research into the Zn(n) ion binding process and localization, providing structural insights into partially Zn(n)-depleted species of MT2 through a combination of mass spectrometry strategies and molecular dynamics simulations. We attempted to decipher the Zn(n) binding process and its implications for reactivity and metal buffering.

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Introduction

Mammalian metallothionein (MT) is a small ~ 7 kDa cellular protein with a high ($\sim 30\%$) content of cysteine residues. It was discovered 60 years ago as a Cd(π)-containing protein in horse kidneys.¹ However, further studies showed that Zn(π) and Cu(π) ions are the most relevant biologically. Mammalian metallothionein is

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present in four major isoforms, MT1-MT4, which are tissuespecific proteins. MT1 (with 9 subisoforms) and MT2 are present in all kinds of tissues in various amounts and ratios, while MT3 and MT4 are mostly present in the brain and squamous epithelia.^{2,3} Structural studies of unsaturated MT have proven to be a challenge mainly due to its probably unfolded and highly dynamic structure. To date, only one X-ray structure of metallothionein has been solved: hepatic rat MT2.4 However, NMR structures of individual (isolated) domains of human, rat, rabbit and mouse MT1-MT3 were also determined.⁵⁻⁸ Mammalian metallothionein folds into two separate domains with two Zn-S clusters, one with 3 Zn(π) and 9 Cys (β -cluster) and the other with 4 Zn(π) and 11 Cys $(\alpha$ -cluster) (Fig. 1a). In both domains metal ions are present in tetrahedral geometry and tetrathiolate environments, where sulfur donors are bound to one or two (bridging donors) metal ions. It is worth noting that the X-ray structure is based on Cd(II)-induced *in vivo* protein,⁴ which has one Cd(II) ion in the β -domain at position IV and four ions in all positions (I, V, VI and VII) in the α -domain (Fig. 1b). MT reconstituted *in vitro* also binds seven Zn(II) or seven $Cd(\pi)$ ions. Most of the current knowledge regarding the structural, biophysical and even biochemical properties of MTs is based on cadmium MTs, which are more convenient to study due to their better spectroscopic properties, less dynamic nature or higher protein affinity, in comparison with a zinc counterpart. Zinc metallothioneins are much less explored mostly due to the lack of appropriate methods to study the structural or biophysical features of their complexes.



Fig. 1 Crystal structure and cluster organization in hepatic rat MT2. (a) Crystal structure of Zn_2Cd_5MT isolated from rat liver (PDB ID: 4MT2).⁴ (b) Schematic representations of β - and α -metallothionein clusters in MT2. Yellow and grey circles refer to Cd(II) and Zn(II), respectively, found in the crystal structure. Metal ion numbering corresponds to the order of ¹¹³Cd-NMR chemical shifts of Cd₇MT2.

In early studies it was shown that seven Zn(II) ions bind to the apo-form of metallothionein (thionein) with the same, undistinguishable affinity which varies from $\sim 10^{-12}$ to 10^{-13} M in terms of average dissociation constant, depending on the literature source.^{9,10} In these approaches $Zn(\pi)$ -to-protein affinity was determined by competition with either protons or chelating agents assuming either identical pK_a of all cysteine thiols or equal contribution of ligand-to-metal charge transfer (LMCT) of Zn-S bonds in observable absorbances in the UV range. Our previous study performed on highly sensitive zinc fluorescent probes, FluoZin-3 and RhodZin-3, showed that Zn(II) ions bind to human MT2 with three ranges of affinity.¹¹ Four $Zn(\pi)$ ions are bound tightly with an average dissociation constant of $\sim 10^{-12}$ M as previously determined for all metal ions. Another two Zn(II) ions bind moderately with a dissociation constant of 10^{-11} to 10^{-10} M. The last and the weakest Zn(II) is bound with nanomolar affinity ($K_{\rm d} \sim 10^{-8}$ M). The differentiation of Zn(II) stability constants has been recently shown for MT3 by ITC experiments with EDTA as a competitor, showing that differentiation of Zn(II)-to-protein affinity is common across mammalian metallothionein isoforms.¹² In consequence, Zn(II) binding or dissociation occurs sequentially and intermediates such as Zn₄MT2, Zn₅MT2 and Zn₆MT2 are formed, which has been shown in several examples in ESI-MS titrations.^{13,14} It is worth noting that these species play an important role in free Zn(II) buffering due to their unsaturated nature and various $Zn(\pi)$ affinities. They can serve as a $Zn(\pi)$ donor and acceptor at the same time and buffer Zn(II) from pico- to nanomolar concentration. Considering cellular free Zn(II) concentration and its natural fluctuations in the range from 10^{-11} to 10^{-9} M, it is apparent that partially Zn(n)-depleted metallothionein species should be present as stable MT2 forms under cellular conditions. Studies performed on HT-29 cells have shown that cells contain in the cytosol fraction a surplus ($\sim 10\%$) of unoccupied tightly Zn(II) binding proteins. Differential chemical fluorescent modification of metallothionein indicated that a major fraction of this surplus belongs to partially Zn(II)-saturated metallothionein.^{15–17} Moreover, a number of studies have already shown that various ratios of thionein (T) to total MT (T + MT) control the activity of zinc enzymes or Zn(II)-modulated enzymes, zinc finger folding and saturation of other structural zinc sites.¹⁸⁻²¹

In this study we aim to connect the current knowledge regarding the structural properties of mammalian metallothionein and its $Zn(\pi)$ buffering properties. Assuming that metallothionein is always naturally present as Zn_7MT species is not valid, and $Zn(\pi)$ -to-protein content depends on many factors such as oxidative or nitrosative state, thionein induction level and cellular or local free $Zn(\pi)$ concentration.^{22,23} The way how $Zn(\pi)$ is buffered and muffled is still not fully understood. One way to explore this issue is to understand the structural and biophysical features of highly dynamic $Zn(\pi)$ -depleted MTs species, especially those with five (Zn_5MT) and six (Zn_6MT) $Zn(\pi)$ ions, which seem to be the most relevant for the cellular zinc buffering process. Although knowledge regarding the physicochemical properties of $Zn(\pi)$ -depleted species is growing, currently little is known about the structure and exact $Zn(\pi)$

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distribution among those species of various MT isoforms.^{11,24} Moreover, the molecular bases that differentiate Zn(II)-toprotein affinity in both domains of metallothionein are also not fully understood.²⁵ In this study we combined classical chemical labelling of cysteine residues by iodoacetamide (IAA) with mass spectrometry approaches. Although metallothionein labelling with IAA has been applied in the past for $Cd(\pi)$ binding to MT, the lack of high resolution techniques made it impossible to determine the distribution of metal ions accurately and assign them to particular cysteines.^{26,27} Since spectroscopic methodologies rely on measuring an average structure of multiple species and due to the lack of secondary structural elements and aromatic amino acids, limited information about partially depleted MT could be obtained. In contrast, as demonstrated in early studies, mass spectrometry is a useful and unique technique to identify particular saturated species in MT.²⁸ Recently, the usefulness of mass spectrometry with ion mobility detection for studying conformational changes in partially Cd(II)-depleted MT2 species has been confirmed.²⁹ However, Cd(II) coordination positions across cysteine residues were not assessed. It is worth noting that besides simple ESI-MS studies, none systematic research on zinc metallothionein with bottom-up and top-down MALDI-MS has been performed. Therefore, here we aimed to explore the crosstalk of structural and zinc buffering properties of human metallothionein-2 to shed more light on the properties and importance of Zn(II)-depleted species which are critical for understanding the basis of the cellular zinc buffering process and role of metallothionein in zinc homeostasis.

Experimental

Materials

All reagents used in these studies were of high purity and purchased from Sigma-Aldrich, Acros Organic, Roth, BioShop, VWR International, Avantor, and Iris Biotech. All pH buffers prepared with Milli-Q water were incubated with Chelex 100 resin to eliminate trace metal ion contamination. More detailed information about the used materials is listed in the ESI.[†]

Expression and purification of MT2

The cDNA encoding metallothionien-2 (MT2) was synthesized (GenScript, USA) and cloned into the pTYB21 vector using SapI and EcoRV (Thermo Scientific, USA) restriction sites. The primer pair sequence used in the PCR reaction is provided in the ESI.† The pTYB21 vector encoding MT2 – deposited in the Addgene plasmid repository (https://www.addgene.org), plasmid ID 105693 (MT2a) – was transformed into BL21(DE3)pLysS *E. coli* cells. The culture medium (1.1% tryptone, 2.2% yeast extract, 0.45% glycerol, 1.3% K₂HPO₄, 0.38% KH₂PO₄) was prepared as described previously.^{30,31} Transformed bacteria (4 l) were cultured at 37 °C until OD₆₀₀ reached 0.5, then induced with 0.1 mM IPTG and incubated overnight at 20 °C with vigorous shaking. All subsequent steps of purification were conducted at 4 °C. Cells were collected by centrifugation at 4000 × *g* for 10 min, resuspended in 50 ml of cold buffer A (20 mM HEPES, pH 8.0, 500 mM

NaCl, 1 mM EDTA, 1 mM TCEP) and sonicated for 30 min (1 min "on", 1 min "off") followed by centrifugation at $20\,000 \times g$ for 15 min. The supernatant was incubated with 20 ml of chitin resin in buffer A and kept overnight with mild shaking. After the incubation resin was washed 4-5 times with 50 ml of buffer A, to induce the cleavage reaction 25 ml of buffer B (20 mM HEPES, pH 8.0, 500 mM NaCl, 1 mM EDTA, 100 mM DTT) was added to the resin and the mixture was incubated for 36-48 h at room temperature on a rocking bed. The reaction was monitored by SDS-PAGE. Briefly, the centrifuged solution containing protein was acidified to pH ~ 2.5 with 7% HCl and subsequently concentrated to a small volume using Amicon Ultra-4 Centrifugal Filter Units with a membrane NMWL of 3 kDa (Merck Millipore, USA) and purified on an SEC-70 gel filtration column (Bio-Rad, USA) equilibrated with 5 mM HCl. The identity of apo-MT2 protein (thionein, T) was confirmed by ESI-MS, using an API 2000 instrument (Applied Biosystems, USA). The average molecular weight $(M_{\rm W})$ calculated was 6042.0/6042.2 Da, respectively. Purified apo-MT2 was used for experiments immediately after purification due to its instability under aerobic conditions. The yield of pure MT2 production was 3.5 mg per liter of bacterial culture.

Synthesis of β-domain of MT2

The N-terminal fragment of MT2 (Met1-Ser32) corresponding to the β -domain (β MT2) was synthesized on TentaGel R RAM resin (loading 0.19 mmol g⁻¹, Rapp Polymere GmbH) using a Liberty 1 microwave-assisted synthesizer (CEM, USA) according to the standard Fmoc strategy described in detail before.^{32,33} The resinattached peptide was cleaved from the resin by 2 h incubation with a mixture of TFA/thioanisole/EDT/anisole (90/5/3/2, v/v/v/v), followed by precipitation in cold (-20 °C) Et₂O. The crude peptide was purified by HPLC (Dionex Ultimate 3000 system) on a Phenomenex C18 column (Gemini-NX 5 µm, 110 Å) using 0.1% TFA with a 0–45% ACN gradient in 30 min and then lyophilized. The identity of the obtained pure peptide was confirmed by ESI-MS (API 2000 Applied Biosystems, USA). The average mass calculated for the β -domain was 3252.3/3252.8 Da.

Reconstitution of MT2 and the β -domain with Zn(II)

Aliquots of apo-MT2 and apo- β MT2 in diluted HCl (pH 2.5 with 1 mM TCEP) were mixed with zinc sulfate at a molar ratio of 1:9 or 1:4 under a nitrogen blanket, and the pH adjusted to 8.6 with 1 M solution of Tris base.³⁴ The samples were concentrated with Amicon Ultra-4 Centrifugal Filter Units with a membrane NMWL of 3 kDa, and purified on an SEC-70 column (Bio-Rad, USA) equilibrated with 20 mM Tris-HCl buffer, pH 8.6. The concentration of the purified protein was obtained spectrophotometrically, *via* DTNB and PAR assays regarding thiol and Zn(II) concentration, respectively.^{35,36} Additionally, samples were analysed by ICP (ICP-AES iCAP 7400, Thermo Scientific) to confirm the spectrophotometric results.

ESI-MS-monitored titration of apo-MT2 with Zn(II)

The Zn(II) titration study was performed in degassed 10 mM $(NH_4)_2CO_3/ACN (9:1, v/v)$ buffer solution, pH 7.6. A solution of 25 µM apo-MT2 was mixed with 0–7 molar equivalents of zinc

acetate (500 μ M), and buffer solution was added to a final volume of 200 μ l. The samples were incubated for 30 s and injected by a syringe pump (40 μ l min⁻¹ flow rate) into an ESI mass spectrometer (API 2000 Applied Biosystems, USA). MS spectra were recorded in positive ion mode during 5 min in the 1000–1800 *m*/*z* range.

Alkylation of MT2 for MALDI-MS studies

All solutions were degassed by purging with nitrogen just before use. To an Eppendorf plastic tube (1.5 ml) filled with nitrogen the following were added sequentially: 1 nmol of apo-MT2 in 10 mM HCl, 20 μ l of 100 mM (NH₄)₂CO₃, pH ~ 8, and an appropriate amount of 500 µM ZnSO₄ (1, 2, 3, 4, 5, 6 and 7 molar equivalents over apo-MT2, respectively). A control experiment with no Zn(II) ions added was also performed. The final volume was adjusted to 55 µl with Milli-Q water. The pH of the final solution was 7.5. After gently mixing, 3 µl of 10 mM iodoacetamide (IAA) (30 nmol, 1.5 molar equivalents over one Cys) was added. Alkylation was carried out under a nitrogen blanket in darkness for 15 min, followed by freezing the samples on dry ice to prevent further reaction. Each sample was desalted while removing excess IAA using ZipTip µ-C18 (Merck Millipore, USA), and eluted with 10 µl of Milli-Q water/ACN solution (50:50, v/v). MALDI spectra were recorded for each sample (1 µl of sample diluted 12 times with CHCA matrix in ACN) on an ABI 4800 MALDI TOF/TOF mass spectrometer (Applied Biosystems, USA).

Trypsin digestion of alkylated MT2

4.5 µl of a desalted solution of partially alkylated MT2 was pipetted into a 250 µl PCR tube. Acetonitrile was removed before digestion with trypsin on a Maxi dry plus speed vacuum system (Heto Lab Equipment, UK) for approximately 10 minutes, followed by adding 1.5 µl of diluted trypsin (0.5 µg ml⁻¹ in 50 mM AcOH) and 3.5 µl of 100 mM (NH₄)₂CO₃ to each sample. Digestion was carried out for 10 minutes at 37 °C followed by freezing the sample on dry ice to prevent further digestion. All samples were lyophilized and kept at -20 °C before analysis.

Nano-LC separation and MS measurements of digested samples

Lyophilized samples were dissolved in 30 µl of 0.1% TFA in LC-MS quality water and transferred into low protein binding autosampler vials. Nano-LC separations were carried out on an Easy nano-LC II instrument (Bruker, Germany) using an Acclaim PepMap C18 3 μ m column (75 μ m i.d. \times 15 cm, 100 Å) with a gradient of 0-60% B in A (A: 0.1% TFA/H₂O, B: 0.1% TFA/ACN) during 24 minutes. Collected fractions were pooled every 15 s and spotted on a MALDI plate pre-spotted with HCCA matrix (PAC II 384, Bruker, Germany) as a single spot (96 spots in total for each sample). MS spectra were recorded off-line using an Ultraflextreme MALDI spectrometer (Bruker, Germany), in the 700-3500 m/zrange, calibrated before each of four adjacent spots with Peptide Calibration Standard PACII (Bruker, Germany). The laser intensity was 29% for calibration and 31% for measurement. MS/MS spectra were recorded in LIFT mode, in the 800–3000 m/z range, with the laser intensity adjusted manually.

Preparation of metal-depleted SDH

An apo-form of sorbitol dehydrogenase (SDH) was prepared according to a previously established procedure by the treatment of commercial SDH (Roche) with dipicolinic acid and subsequent purification.^{18,37} Concentration of purified apo-SDH was examined by its titration with standard Zn(II) ions followed by enzyme activity determination. The activity was determined spectrophotometrically as time-course measurements at 340 nm as a result of NADH oxidation to NAD⁺ upon conversion of D-fructose to D-sorbitol. For that purpose protein samples were incubated in 50 mM HEPES buffer, pH 7.4, with 100 μ M TCEP, 0.1 M D-fructose and 0.125 mM NADH.

Zn(II) transfer from MT2 to metal-depleted SDH (apo-SDH)

The rate and efficiency of Zn(II) transfer from MT2 to apo-SDH were monitored enzymatically using the assay described above. Metal-depleted SDH was mixed in a molar ratio of 1:1 with freshly prepared 1 µM MT2 and incubated in 50 mM HEPES buffer, pH 7.4, in the presence of 100 µM TCEP. Protein aliquots were taken after certain periods of time (0-120 min) and assayed for the recovery of enzymatic activity. Full recovery of the SDH was reached by apo-SDH saturation with Zn(II) ions under the same conditions. Samples for mass spectrometry were prepared in a different way. For that purpose 1 nmol of MT2 and apo-SDH were mixed with 20 µl of 100 mM (NH₄)₂CO₃ and incubated for 4, 30, 60 or 120 min at room temperature. After that time the sample was modified with IAA according to the procedure described above and finally measured using an ABI 4800 MALDI TOF-TOF mass spectrometer (Applied Biosystems) and subjected to nano-LC separation and a further procedure as described above.

Mass spectrometry data analysis

Obtained mass spectrometry data were analysed and processed using Data Explorer Software (Version 4.9, Applied Biosystems), flexAnalysis (Version 3.4, Bruker Daltonik) or Data Analyst (Version 1.4.2, Applied Biosystems) according to the equipment used.

Circular dichroism

Circular dichroism (CD) spectra of zinc metallothionein were recorded using a J-1500 Jasco spectropolarimeter (JASCO) at 25 °C in a 2 mm quartz cuvette, under a constant nitrogen flow over the range of 196–270 nm with a 100 nm min⁻¹ speed scan. Final spectra were averaged from three independent scans. 10 μ M thionein (apo-MT2) was incubated with 6 and 7 eq. of ZnSO₄ in 20 mM Tris–HCl buffer (100 mM NaCl, 200 μ M TCEP pH 7.4) and recorded. Independently, reconstituted 10 μ M MT2 was incubated with 0–100 μ M of EGTA or EDTA and spectra were recorded after 10 and 30 minutes.

Molecular dynamics

Initial coordinates were taken from the crystal structure of rat Zn_2Cd_5MT2 (PDB ID: 4MT2). All molecular dynamics simulations were performed using an AMBER99SD force field for

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protein atoms and a semi-bonded approach for Zn_xS_y clusters.38,39 Hydrogen bonds were added to the metalloprotein using the LEAP module of the AMBER program. The structure was solvated in a box of TIP3P water molecules with a density of 0.813 g cc^{-1} and counter-ions were added to neutralize the system. After solvation and neutralization, periodic bound conditions were set up using the particle mesh Ewald method with a cut-off of 12 Å. The system was firstly energy minimized for 10000 steps by means of a conjugate gradient algorithm followed by a steep descent algorithm, setting up the switching cut-off from 10 to 12 Å for computing electrostatics and Lennard-Jones energies. The LINCS constraint was applied to all bonds involving hydrogens. MD simulations were performed in two steps. The system was equilibrated using the NVT ensemble with leapfrog Verlet integrator (leap) and Hoover thermostat (303.15 K and 1 atm) for 1 ns with a timestep of 0.002 (2 fs) and NVT production for 4 ns with a 2 fs timestep. Molecular dynamics simulations were carried out with the module LEAP from AMBER and with the GROMACS software.⁴⁰ The PROPKA 2.0 method was used for the prediction of the pK_a values of ionizable residues.41

Results

In order to gain detailed structural insights into the $Zn(\pi)$ complexation pathway in mammalian MT2, we combined differential alkylation of free cysteine thiols by iodoacetamide (IAA) in metal-free and partially Zn(II)-depleted protein with mass spectrometry techniques. To do so, samples of human MT2 were subjected to alkylation with IAA to "fix" the coordination pattern in the desired structural state, and subsequently analysed by (i) MALDI-MS, (ii) nano-LC-MALDI-MS for the partially alkylated tryptic digest and finally by (iii) MS/MS for the nano-LC separated tryptic peptides (Fig. 2). The same approach was used to study the complexation process of the isolated β -domain of MT2 (β MT2) and to map free cysteine residues formed upon Zn(II) transfer from fully Zn(II)-loaded MT2 to metal-depleted sorbitol dehydrogenase (apo-SDH). We supported our study with direct titration of apo-MT2 with consecutive Zn(II) molar equivalents, ESI-MS measurements and molecular dynamics, which were used to examine structure conformational changes upon Zn(II) binding. In the first step of the study, MALDI spectrum of metal-free or partially Zn(II)-loaded MT2 modified by IAA was analysed and compared with the signal profile of unmodified protein obtained by ESI-MS. Then, detailed analysis of tryptic digest MS and MS/MS spectra were carried out and taken into account to propose the stoichiometry of the formed MT2 species and coordination mode. Finally, experimental results were compared with molecular dynamics simulations.

Alkylation of metal-free metallothionein-2

The MALDI spectrum recorded for unmodified MT2 demonstrates one major signal of 6043.7 m/z (Table S1 and Fig. S1a, ESI[†]), which is expected for a pure apo-form. However, alkylated apo-MT2



Fig. 2 Schematic workflow and three types of MS experiments used in this study. (a) Human apo-MT2 was incubated with 0–7 molar equivalents of Zn(II), followed by free cysteine alkylation with iodoacetamide (IAA). Samples were either directly analysed by MALDI-MS or trypsinized and separated on a nano-LC column prior to MALDI-MS. Chosen tryptic peptides were subjected to MS/MS analysis. (b) Sequences of tryptic peptides with the notation of cysteine residue numbers. The arrows indicate cleavage sites of C-termini of lysine residues. Squared brackets correspond to tryptic fragments.

indicates signal distribution with pairwise intensities with clearly visible signals with 12, 14, 16, 18 and 20 modifications (Fig. 3a, Fig. S1b and Table S1, ESI⁺), which is contradictory to the expected single signal of an entirely alkylated protein (20 acetamide moieties). The observed pattern did not change when either extending the reaction time or applying higher IAA concentrations. This effect was largely suppressed when the protein was treated with 6 M urea (Fig. S1c, ESI[†]). In this case the most abundant signal has 20 modifications, but peaks with 17, 18 and 19 modifications are still observed (Fig. S1c, ESI⁺). The tryptic digest revealed significantly incomplete alkylation in regions [1-20] and [32-43], but incomplete modification was also observed in other peptides (Table 1, Tables S2 and S3, ESI^{\dagger}). The pK_a values of cysteine thiols predicted by PROPKA method revealed an acidity shift of Cys residues, which ranges from 6.9 to 11.9.41 The protonation state of Cys affects solvent exposure and accessibility for Cys to be modified by IAA. Calculations of absolute surface area (ASA) of each residue in apo-MT2 obtained by MD after 4 ns showed cysteine residues from Cys6 up to Cys20 and from Cys34 to Cys42 to be less accessible for solvent (Table S4, ESI[†]). These results are in





agreement with those obtained from tryptic digestion and match fragments [1–20] and [32–43] in which alkylation was not completed, possibly due to the lower solvent accessibility.

Alkylation of partially Zn(II)-depleted metallothionein-2

Alkylation of apo-MT2 was performed in such a way as to determine the positions of modified and protected Cys residues upon addition of each molar equivalent of Zn(II) from 1 up to 7 in order to map the $Zn(\pi)$ -bound position with the highest possible resolution. The MALDI spectrum of apo-MT2 with one $Zn(\pi)$ equivalent shows a maximum of 18 modifications, which is a sign of two cysteine residues protected from alkylation (Fig. 3b). The most abundant signal corresponds to 12 modified residues, whereas in the spectra for metal free MT it corresponded to 16 modifications. Therefore, these four cysteine residues not observed in MS spectra should be involved in metal binding when one Zn(II) equivalent is applied. The ESI spectrum of the same Zn(II)-to-protein ratio reveals the co-existence of three species with 0-2 bound Zn(II) ions, Zn₀₋₂MT2 (Fig. S2b and Table S5, ESI[†]). Additionally, MS/MS spectra show that the [32-43] region is more resistant to alkylation since it showed a tryptic fragment with fewer modifications (2 M) compared to the metal-free protein sample (3 M) (Table 1). MS/MS spectra of selected tryptic fragments revealed Cys36, Cys48 and Cys60 residues as highly protected, so more likely involved in Zn(II) ion coordination (Fig. S3, ESI^{\dagger}). Because Cys41 is modified, and Zn(II) coordinates tetrahedrally, one of Cys33 or Cys34 and Cys37 may be involved in coordination of the first Zn(II) ion, although these occur in both modified and unmodified forms (Fig. 4a). Molecular dynamics results indicated that Cys37 is coordinated to Zn(II), whereas Cys33 and Cys34 remain far from the metal ion coordination sphere (Fig. S4a, ESI⁺). A stabilized structure by interaction between Lys44 and Cys37 was found, which would increase $Zn(\pi)$ -bound thiolate reactivity towards the $Zn(\pi)$ ion. Positively charged lysine will lower the pK_a value of cysteine thiol and thus promote more nucleophilic thiolate formation.^{42,43} The observed peaks indicate labile coordination of the first $Zn(\pi)$ ion under the applied conditions. Surprisingly, the residues responsible for $Zn(\pi)$ binding do not correlate with any positions observed in the crystal structure of fully saturated MT2 (Fig. 1).4,5,7

The addition of two Zn(II) equivalents to apo-MT2 and subsequent alkylation results in 4 up to 16 cysteines modified observable in the MALDI spectrum, indicating a minimum of four highly protected cysteine residues, which is two modifications less in comparison with results obtained for a sample with one Zn(II) equivalent (Fig. 3b and c). Characteristic pairwise peak distribution is no longer visible. Interestingly, a singly modified cysteine residue (1 M) in the tryptic fragment [32–43] emerged; hence if only one cysteine could be modified, the other four Cys residues might be protected in this region (Table 1). In the tryptic fragment [44–51], the signal intensity in MALDI-MS spectra suggests the prevalence of double modification, which would lead to one protected cysteine (Fig. S5, ESI†). MS/MS data revealed that both Cys48 and Cys50 are protected

Table 1 Selected tryptic fragments of human MT2 with various equivalents of Zn(n) obtained after nano-LC off-line MALDI-MS analysis. The number and range of cysteine residues (<u>C</u>) modified by IAA are indicated by *n* M (where *n* indicates the number). n.d. – not detected. All identified tryptic fragments can be found in Table S3 (ESI). The full list of calculated and found *m*/*z* for all tryptic fragments can be found in Table S4 (ESI)

Zn(II) eq.	MDPNCSCAAGDSCTCAGSCK [1-20] MDPNCSCAAGDSCTCAGSCKCK [1-22]	<u>C</u> TS <u>C</u> KK [26-31]	SCCSCCPVGCAK [32–43]	CAQGCICK [44-51] CAQGCICK GASDKCSCCA [44-61]	GASDK <u>C</u> S <u>C</u> A [52–61]
0	3-5 M	1 M	3-5 M	2 M	0 M, 2 M
1	3–5 M	2 M	2–5 M	2 M	2 M
2	2–5 M	1 M	1–5 M	1–2 M	1-2 M
3	5 M	1 M	1–5 M	1–2 M	0-2 M
4	3 M	n.d.	0–5 M	0-2 M	0 M
5	0-2 M	1 M	0–2 M	0-1 M	0 M
6	0 M	0 M	0 M	0 M	n.d.
7	0 M	n.d.	0 M	0 M	n.d.
7	1 M	n.d.	0 M	0 M	n.d.



Fig. 4 Scheme of the most prevalent sequential Zn(II) binding pathway in MT2 based on MS/MS analysis of tryptic peptides differently modified by IAA and MD calculations.

from alkylation, wherein Cys50 is less resistant to those two (Fig. S6a, ESI†). Tryptic fragment [52–61] occurs in singly and doubly modified form, which corresponds to two and/or one protected cysteine (Table 1 and Table S2, ESI†). Because Cys60 was alkylation protected for the sample with one Zn(π) equivalent, similarity with the two Zn(π) equivalent sample is highly possible. The tetrahedral ZnS₄ coordination sphere is probably filled up by either Cys57 or Cys59 (Fig. 4b). Therefore, from the

results obtained we may conclude that Cys33, 34, 50 and 59 or 57 residues were available to coordinate Zn(n), besides those reported previously (Cys36, 48 and 60). Regarding molecular dynamics simulations, Cys33, 34, 36 and 48 coordinate one Zn(n), whereas Cys36, 50, 57 and 60 coordinate a second Zn(n) (Fig. S4b, ESI†).

The MALDI spectrum of undigested apo-MT2 with three Zn(n) equivalents shows a maximum of 12 modifications, strongly suggesting at least eight protected Cys residues, which is four less

than in the previous step (Fig. 3d). Strikingly, a small fraction of unmodified protein (indicated as WT, which corresponds to Zn₇MT2) is observed. ESI titration results are consistent with this finding, as the observed peaks correspond to co-existing forms from Zn₃MT2 up to Zn₇MT2 (Fig. S2d, ESI⁺). The presence of the unmodified tryptic fragment [52-61] showed that Cys57, 59 and 60 are protected against alkylation to some extent (Table 1), but singly and doubly modified [52-61] forms are still visible (Fig. S7, ESI[†]). Fragment [32-43] exhibits an almost unchanged modification mode as compared to the protein with two $Zn(\pi)$ equivalents, so Cys33, 34, 36 and 37 are visible as modified and unmodified (Table 1). MS/MS data for singly and doubly modified tryptic fragments [44-51] show that one or two cysteines are involved in Zn(II) binding, i.e. Cys48 and/or Cys50 may possibly be involved in cluster formation (Fig. S8, ESI⁺). Based on the above results, the existence of a Zn₃S₈ or Zn₃S₉ cluster is inferred (Fig. 4c). Molecular dynamics results indicated that both Cys48 and 50 are involved in Zn₃S₉ cluster formation (Fig. S4c, ESI[†]).

Iodoacetamide modification of apo-MT2 incubated with four Zn(II) equivalents results in 0 and up to 11 modifications of cysteines observed in the MALDI spectrum, which can be attributed to maximally 9 alkylated cysteines in the β -domain and two alkylated residues in the α -domain (Fig. 3e). The most abundant signals correspond to five and four modified cysteines, which translates to 15 or 16 cysteine residues involved in $Zn(\pi)$ binding. This can be attributed to the most present Zn₄MT2 and Zn₅MT2 forms. The ESI-MS of Zn(II) titration study indicates species diversity and the presence of Zn₄MT2, Zn₅MT2, Zn₆MT2 and Zn₇MT2 (Fig. S2e, ESI⁺), but one should note that the presence of particular Zn(II)-depleted species must be taken qualitatively not quantitatively. Tryptic fragment [1-20] is triply modified, which means two cysteine residues resistant to alkylation reside in the β-domain and probably coordinate to the same Zn(II) ion. Tryptic fragments [32-43], [44-51] and [52-61] exist in non-alkylated forms (Table 1), which proves full saturation of the $Zn_4\alpha MT$ cluster (Fig. 4d). Molecular dynamics simulation showed Zn(II) distribution along the Zn₄αMT domain (Fig. S4d, ESI[†]). Whilst fragment [52-61] is observed in unmodified form only, both [32-43] and [44-51] exist in partially modified forms. For fragment [44-51] a fully modified form was not detected (Fig. S9, ESI⁺).

The most abundant signal in the MALDI profile for apo-MT2 with five Zn(π) equivalents conforms to three modified cysteine residues (Fig. 3f); hence the dominant form should have 17 cysteine residues involved in the coordination, which is shifted to two modifications less than in apo-MT2 with four Zn(π) equivalents. This relates to the Zn₄ α Zn₂ β MT2 form (Zn₄S₁₁ in α - and Zn₂S₆ in β -domain). Moreover, the maximum number of modifications observed is 10, so the rest would be involved in Zn(π) coordination forming α -cluster Zn₃S₉₋₁₀ with one labile cysteine (Fig. 3f). Furthermore, increased abundance of unmodified protein is observed (*e.g.* Zn₇MT2 WT form) in MALDI-MS spectra. ESI-MS titration shows that the forms Zn₆MT2 and Zn₇MT2 predominate, although minor intensities from Zn₃MT2, Zn₄MT2 and Zn₅MT2 are still observed (Fig. S2f, ESI†). Tryptic fragments [32–43] and [44–51] show decreased heterogeneity regarding the number of cysteine modifications in the α -cluster (with a range of 0 to 2 and 0 to 1 modified cysteines, respectively) (Table 1). Interestingly, a new pattern is observed when adding 5 Zn(n) equivalents compared to previous titrations regarding tryptic fragment [1–20] since 0 to 2 cysteine residues are modified (0–2 M) (Table 1). So, a minimum of 3 and a maximum of 5 cysteine residues would be involved in Zn(n) coordination. Tryptic fragment [26–31] with two cysteines is observed as singly modified, so the other cysteine might be involved in metal binding to fill up Zn(n) coordination in the β -domain. MD simulations present a structure with Zn₄S₁₁ and ZnS₄ clusters in α - and β -domains, respectively (Fig. 5a and Fig. 4e). As observed in Fig. S12a (ESI†), the structure obtained by molecular dynamics showed a comparable α -domain to the X-ray MT2 structure (PDB ID: 4MT2).⁴

Apo-MT2 with six Zn(II) equivalents shows in the MALDI spectrum only two peaks, with one and without any modification (Fig. 3g). This is consistent with ESI-MS titration (Fig. S2g, ESI⁺), where in the case of six Zn(II) equivalents only Zn₇MT2 and Zn₆MT2 species are observed. All tryptic fragments are found to be unmodified, except for the region 21-25, which was not detected (Table 1). MS/MS spectra for tryptic peptides confirm the lack of modification in region 31–61 (α -domain) (Fig. S10, ESI⁺). Because there is one single modification, and it is proved not to be found in this region (31-61), there is strong evidence that either Cys21 or Cys24 should be modified by IAA. MD results showed a possible structure where $Zn(\pi)$ binds tetrahedrally to Cys24, whereas Cys21 is not coordinated (Fig. 5b). Regarding whether Cys21 or Cys24 is modified, our MD results indicate H-bonding interaction from Ser6 to Cys21 stabilizing the latter, and thus Cys21 is modified (Fig. 4f). On the other hand, Cys5 is found to not be involved in coordination. Fig. S12b (ESI⁺) showed overlapping structures between the X-ray MT2 structure (PDB ID: 4MT2), validating the MD structures obtained.⁴

Similarly, in the case of apo-MT2 with seven $Zn(\pi)$ equivalents only two peaks are well observed in the MALDI spectrum with one and no modification (Fig. 3h). This is also consistent with ESI-MS titration where Zn7MT2 and Zn6MT2 species are observed (Fig. S2h, ESI[†]). All found tryptic fragments show no modifications, except for fragment [1-22], which is singly modified (Table 1). Fig. 6 shows the comparison of two tryptic fragments [1-20] with no modification and [1-22] with one modification, which indicates that Cys21 might be modified. Moreover, the MS/MS spectrum of the singly modified [1-22] fragment strongly confirms localization of the acetamide modification on the Cys21 residue (Fig. S11, ESI†). Addition of the last $Zn(\pi)$ ion caused overall structure arrangement that involved coordination of those previously non-coordinated cysteines, namely Cys21 and Cys5, and Zn(II) coordination filled it with Cys7 and Cys24 (Fig. 4g). Our molecular dynamic simulation results for Zn7MT2 species are in good agreement with the X-ray structure, thus assessing and validating the quality of the results previously reported (Fig. S12c, ESI⁺).

Alkylation of metal-free and Zn(π)-depleted βMT2

In order to investigate whether or not the Zn(n) binding process in the isolated β -domain of MT2 is comparable with the whole



Fig. 5 Simulated metallothionein-2 structures obtained from molecular dynamics. (a) Molecular structure of Zn_5MT2 ; (b) molecular structure for Zn_6MT2 . Zn(II) ions are represented by grey colour.

protein, we performed a MALDI-MS/MS study on non-digested and trypsinized protein. The MALDI spectrum of undigested β MT2 in the absence of Zn(II) presents a single peak corresponding to the β -domain fragment with all nine Cys modified by IAA (Fig. S13 and Table S6, ESI†). No trace of threedimensional structure is identified, in contrast to the full apo-MT2 protein (see above Fig. 3a). An isolated β -domain is shown in Fig. S14a (ESI†) and has been obtained from molecular dynamics simulation.

Addition of one molar equivalent of Zn(II) results in a maximum of seven modifications, while the most abundant signal corresponds to the species with three modified cysteines (Fig. S13 and Table S6, ESI⁺). The MALDI spectrum does not show the presence of a fully modified β -domain. This together suggests that two species can be formed with one (ZnS_4) and two Zn(II) ions (Zn_2S_{6-8}) bound to the β -domain peptide. ESI-MS titration confirms that the system is highly dynamic due to the presence of multiple species with 0-3 bound $Zn(\pi)$ ions; however, species with one and two Zn(II) predominate, which is consistent with MALDI data (Fig. S13, S15 and Table S7, ESI⁺). Upon addition of one Zn(II) equivalent tryptic fragments [1-20] and [1-25] are less extensively modified, with no significant changes for others; hence the first Zn(II) is probably coordinated in the [1-25] region (Fig. 7 and Table S8, ESI⁺). To elucidate those Cys residues involved in Zn(II) coordination, MS/MS analysis over undigested 5 M β MT2 with 1 Zn(π) eq. was performed (peak 5 M in Fig. S13b, ESI[†]). The produced y fragmentation series revealed that Cys19, 21, 24, 26 and 29



Fig. 6 Comparison of MALDI spectra of two spots indicating acetamide modification of Cys21 residue. (a) Peak [1-22] with 1 M (2209.853 *m/z*); (b) peak [1-20] with 0 M (1923.633 *m/z*). M indicates the presence or lack of modification in the selected fragment.

are modified (Table S9, ESI[†]), so Cys5, 7, 13 and 15 should form a coordination site for the first $Zn(\pi)$ ion, which is contrary to annotations in the X-ray structure (Fig. 1b, β -cluster). The most abundant fourfold modified tryptic fragment [21–31] of β MT2 from a 1 eq. $Zn(\pi)$ sample indicates modified Cys21, 24, 26 and 29 residues, and the triply modified tryptic fragment [23–31] from the same sample confirms Cys24, 26 and 29 modifications (Fig. 7). Therefore, all data consistently prove that coordination



Fig. 7 Comparison of MALDI spectra of trypsinized and differentially modified β MT2 samples incubated with 0–3 Zn(II) equivalents. All found tryptic fragments are listed in Table S7 (ESI†). Asterisk and M indicate the occurrence of two disulfide bridges and modifications in the selected fragment, respectively.

of the first $Zn(\pi)$ is formed by Cys5, 7, 13 and 15, contradictory to all coordination sites for available crystal and NMR structures.^{4,5,7} This result is supported by the simulated structure obtained from MD calculations (Fig. S14b, ESI†).

Addition of two molar equivalents of Zn(n) ions causes the maximum modification of four cysteines and the singly modified form is the most abundant (Fig. S13c, ESI[†]). The results are consistent with ESI titration results, as for two Zn(n) molar equivalents two species exist, $Zn_2\beta$ MT2 and $Zn_3\beta$ MT2 (Fig. S15c, ESI[†]). Moreover, the results are consistent with observed tryptic peptides, which exist mostly in unmodified forms (Fig. 7). Specifically, the [1–25] fragment exists in unmodified form only, while [1–20] and [21–31] both demonstrate an accompanying singly modified form. The obtained structure based on MD simulations displayed Cys5 and Cys29 free of Zn(n) coordination (Fig. S14c, ESI[†]), supporting the MS results.

Application of the third molar equivalent of Zn(n) shows all Cys residues protected from alkylation (Fig. S13d, ESI[†]); hence all are involved in Zn(n) coordination. These results are in agreement with those obtained by ESI titration results in which only the $Zn_3\beta$ MT2 form is observed (Fig. S15d, ESI[†]) and with MD simulations (Fig. S14d, ESI[†]). Furthermore, all visible tryptic peptides are unmodified (Fig. 7).

Zn(II) transfer from MT2 to apo-SDH

Apo-SDH activity is restored relatively quickly during incubation with fully Zn(II)-loaded MT2 at room temperature and pH 7.4, in such a way that 50% of the enzyme activity is recovered after ~ 17 min of incubation. Almost full recovery (more than 80%) was observed after 2 h of protein incubation (Fig. 8a). Since the SDH molecule binds one $Zn(\pi)$ ion in the active centre, and both proteins (MT2 and apo-SDH) were mixed equimolarly, we assumed that one Zn(II) ion from MT2 was transferred to the metal-depleted metalloenzyme. The MALDI spectrum recorded for the sample alkylated after 120 min of incubation shows up to three possible modified cysteine residues (Fig. 8b). Tryptic peptides obtained in the subsequent experiment prove single modification in region [26-31] (Fig. S16 and Table S10, ESI[†]). Because the undigested protein bears three modifications, the other two should be located on Cys21 and Cys24. Similarly, the sample incubated with apo-SDH for 5 min showed no modifications in regions 1-20 and 31-61 (Table S10, ESI⁺), but the undigested protein bears a single modification (data not shown). Because region 21-30 was not detected, it is clear that Cys21, 24, 26 or 29 was modified. These data show that the modification pattern of Zn₆MT2 species obtained by one Zn(II) ion transfer from Zn₇MT2 to apo-SDH seems to be identical to Zn₆MT2 obtained by addition of 6 Zn(II) equivalents to apo-MT2.

CD spectra of Zn_6MT2 obtained by $Zn(\pi)$ association or dissociation

Independently from mass spectrometry studies we examined whether or not CD spectra of Zn_6MT2 species differ depending on the preparation pathway. For this purpose, the CD spectrum of 10 μ M thionein (apo-MT2) incubated with 6 equivalents of



Fig. 8 Transfer of Zn(II) ions from MT2 to apo-SDH. (a) Time-dependent apo-SDH activity recovery with fully loaded MT2 mixed in molar ratio 1:1, 50 mM HEPES (l = 0.1 M from NaCl), pH 7.4, 25 °C. (b) MALDI spectrum of undigested Zn₇MT2 incubated with apo-SDH for 120 minutes and alky-lated with IAA. M and asterisks indicate modifications in the selected fragment and impurities, respectively.

 $Zn(\pi)$ (60 μ M) was recorded in the UV-range. Addition of the seventh Zn(II) ion almost does not change the spectrum in the applied UV range (Fig. S17a, ESI†). Because apo- and holo-SDH forms demonstrate intensive CD signals in the UV range, the use of apo-SDH enzyme as an acceptor of $Zn(\pi)$ is not possible in this case due to very weak signals of zinc metallothionein species and possible effect alteration.⁴⁴ To avoid this effect we used EGTA as an acceptor of Zn(II). This chelator has been shown in the past to compete with the weakly bound Zn(II) in human MT2.^{11,18} For that purpose fully loaded metallothionein was incubated with various concentrations of EGTA for 10 or 30 min and spectra were recorded. Fig. S17b (ESI⁺) shows that the CD spectrum of Zn7MT2 incubated with 20 µM EGTA does not differ from that of Zn₆MT2 obtained by the addition of 6 eq. to apo-MT2. Only adding EDTA (\sim 3.5 order of magnitude tighter chelator than EGTA) causes virtual changes in the weak CD spectrum (Fig. S17b, ESI[†]).

Discussion

Intramolecular interactions in metal-free MT2

Early structural studies on mammalian apo-MTs originally concluded that the lack of UV and CD signals specific for α -helices and β -sheets is proof of a disordered conformation with no secondary structural features,^{45,46} although obtained NMR resonance shifts suggested some residual, untypical hydrogen bonding network.⁴⁷ Later on, molecular simulation studies confirmed that thionein retains a significant amount of structural features upon metal dissociation,⁴⁸ which are probably held together *via* the hydrogen bonding network.⁴⁹ Those observations were supported by the shift of charge state distribution observed in the ESI spectra of sequentially Zn(π)-depleted metallothionein,⁵⁰ IM-MS results and molecular dynamics simulation studies.²⁹

The encountered problems with obtaining entirely alkylated apo-MT2 in our study confirm the existence of a residual, threedimensional-like molecular scaffold. Furthermore, pairwise peak distribution observed in the MALDI spectra suggests the presence of intermolecular interactions involving thiol groups, lowering their solvent exposure, rotation or reactivity. Literature data suggest that an untypical hydrogen bonding network may be present, either involving the OH group from Ser residues which may contribute to steric hindrance of SH groups due to their close proximity or involving the SH group directly.^{29,47,51} Despite the lowered susceptibility to alkylation of thionein, literature data demonstrate that it is readily oxidized by a series of compounds, including DTNB used for the quantitative determination of thiol/ thiolate group concentration.^{11,52} Molecular dynamics simulation studies revealed that thionein exists in multiple conformation states in solution, fluctuating between extended and globularshaped final forms, with only a transient network of hydrogen bonds.²⁹ Literature data as well as our results support the conclusion that apo-MT2 demonstrates residual structural features, which exist in a constant equilibrium with an extended, random coil form, although under the applied conditions the structurally ordered form seems to prevail.

$Zn(\pi)$ binding to the $\alpha\text{-domain}$ of MT2 is sequential under cellular pH

Knowledge of the Zn(II) binding mechanism to metallothionein is a critical issue for the understanding of the zinc sites' features and functions in Cys-rich zinc proteins. This is especially important for cellular zinc homeostasis and the zinc buffering mechanism, in which the presence of partially saturated species of metallothionein seems to function as a Zn(II)donor and acceptor buffering Zn(II) in the appropriate free concentration. In this study, with the combination of Zn(II)to-protein titration, alkylation, enzymatic digestion and several mass spectrometry approaches, we were able to look deeper into the Zn(II) coordination mechanism of MT2 with a resolution that was not experimentally obtained before. However, due to high internal dynamics and heterogeneity of differentially $Zn(\pi)$ -loaded forms, a multitude of information was not deciphered. In the following text one has to take into account several issues: (i) co-existence of several diversely saturated forms in each sample, (ii) high flexibility and internal dynamics of the system, and (iii) the observed peaks result from all present forms and provide overall information about molecular entities in each sample. Nonetheless, we attempted to elucidate the coordination site for each $Zn_{0-7}MT2$ stoichiometry.

The first binding event occurs by Zn(II) coordination to Cys36, 48 and 60 residues, with Cys33/Cys34/Cys37 completing the coordination sphere (Fig. 4a). Primary sequence motifs $C^{57}SCC^{60}$ and $C^{33}CSCC^{37}$, present in the α -domain, possess a well-established tendency for preferential metal ion binding and thermodynamic advantage of proximal cysteine ligands.⁴ Inspection of the MT2 sequence shows that Cys60 and Cys36 residues are a part of both motifs. It is worth noting that CXXC sequence motifs used for Zn(II) binding are the most wide-spread binding motifs in zinc proteins and form the most

stable metal-protein complexes.53-55 Since experimental data revealed that Cys36 was involved in Zn(II) coordination, Cys33 could complete the Zn(II) coordination sphere ligand because it forms a CXXC binding motif with Cys36. Interestingly, this binding site does not resemble any of the sites denoted in crystal and NMR structures. It is also different from the putative nucleation centre of rabbit MT2 (49-61 region) for Cd(II)induced folding.⁵⁶ Also, molecular dynamics results indicated the prevalence of Cys37 over Cys33 and Cys34 to coordinate the first $Zn(\pi)$ ion (Fig. S4a, ESI⁺). In this model, Cys33 and Cys34 remain far away from the coordination sphere (12.2 and 6.7 Å, respectively); moreover, the former interacts weakly, H-bonding with Lys20 (3 Å). This is not a surprising result for Zn₁MT in which probably a residual network of hydrogen bonds maintains structural order of the α - and β -cluster, although flexibility and dynamic behaviour still occur. This indicates the need of Zn-S bond rearrangement during complexation of subsequent Zn(II) ions and confirms the high coordination dynamics of MT2 metal clusters.57-59

Binding of the second Zn(II) ion is associated with four more cysteine residues resistant to modification, which is clear evidence for the formation of two independent ZnS₄ sites. The obtained results suggest the presence of two distinct cysteine groups that are alkylation protected to different extents, i.e. Cys48, 50 and 60 are highly protected, while Cys33, 34, 36 and 37 are less protected. Such a pattern suggests separate binding of two Zn(II) ions to CSCC and CCSCC motifs, respectively, accompanied by one Zn-S bond rearrangement upon incorporation of the second Zn(II) ion (Fig. 4b). The proposed bond re-formation is caused by steric hindrance in the C³³CSCC³⁷ motif, which could involve only three cysteine residues for coordination of the same $Zn(\pi)$ ion.^{56,60} The second metal ion is coordinated by Cys50, Cys60 and possibly Cys36/37. Since Zn(II) shows preference towards CXXC, it is highly possible that the fourth ligand is thiolate of the Cys57 residue. Another possible model of the second Zn(II) ion coordination is its binding to Cys50, 57, 59 and 60, as has been shown by Munoz and co-workers.56 In such a scenario Cys37 remains free yet resistant to alkylation due to steric hindrance resulting from adjacent binding residues. Independent formation of two ZnS4 sites was also predicted by Rigby and co-workers in their molecular dynamics study on cadmium MT.61 Our experimental and simulation results are highly consistent with their simulation study, but differ in some Zn(II)-Cys coordination. While they proposed Cys33, 34, 44, and 48 for one out of two Zn(II) ions, slightly different results where Cys36 replaces Cys44 were obtained. As well as for the second Zn(II), in our model Cys36 acts as a bridge and Cys59 remains free. Furthermore, studies performed on cobalt MT showed that first 3-4 Co(II) ions are bound in independent sites, although their exact localization has not been studied in detail to date.^{62,63} However, the results obtained for $Zn(\pi)$, $Cd(\pi)$, $Co(\pi)$ and other divalent metal ions should be treated with caution, as discussed below.

Analysis of the binding process of the third $Zn(\pi)$ ion uncovers even higher sample heterogeneity, compared to the previous two ions. ESI spectra indicate the formation of

 $Zn_{3-7}MT2$, which shows that α -cluster formation is not strictly cooperative since multiple metalated species are observed simultaneously (Fig. S2, ESI⁺). It is worth noting that signal intensities in the ESI spectrum should not be treated quantitatively because of the electrospray ionization operating principle and do not necessarily reflect the real distribution of species in the solution phase. This fact has been underlined recently by two recent reports proving that investigations of Zn(II)-peptide/ protein equilibria by ESI in the gas phase are not quantitative due to zinc deposition or protein supermetalation during analysis.^{64,65} Nevertheless, such a dynamic picture shows that two possible Zn(II) binding pathways may occur "clustered" via cluster formation and "beaded" via formation of individual coordination sites prior to coalescence into a cluster, as was postulated in a recent study.¹⁴ On the other hand, our results strongly support the model in which initial metal ion binding takes place only in the α -domain. Obtained tryptic digest patterns for two and three Zn(II) equivalents do not show any major differences; hence it is reasonable to assume two independent ZnS₄ sites in the Zn₂MT2 and Zn₃S₉ cluster in Zn₃MT2 form. Moreover, measured molecular masses observed in the ESI titration study confirm this hypothesis (Table S5, ESI[†]). Fig. 4c presents the possible location of three metal ions in the α -domain. However, it should be mentioned that other configurations of bridging cysteines may occur or several similar conformers may be present in the solution. In this conformation, Cys41 and Cys44 are not involved in coordination, although Cys41 interacts with Lys31 by weak H-bonding (3 Å) and remains far away from the coordination sphere (Fig. S4c, ESI⁺).

Recent semi-structural studies on Cd(n) formation intermediates of MT2 by ESI-IM-MS and MD simulations suggest the presence of several conformers of Cd_3MT2 that differ in overall shape and total protein area, which is in good agreement with our experimental results.²⁹ Molecular dynamics also allowed prediction of two independent CdS_4 sites for Cd_2MT2 and formation of a cluster for Cd_3MT2 , although in contrast to our stoichiometry, Zn_3S_9 , they proposed a slightly different stoichiometry, Cd_3S_{10} .⁴⁹

Eleven modifications in MT2 observed in the MALDI spectrum after addition of four $Zn(\pi)$ equivalents may be attributed to nine alkylated cysteines in the β-domain and a fully formed α -cluster (Zn₄ α MT), proven by the obtained tryptic peptides (Table 1). Peaks with lower modification number (Fig. 3e), signals observed in the titration study (Fig. S2e, ESI[†]) and the obtained tryptic peptides (Table 1) all point to the existence of higher metalated species, *i.e.* Zn₅₋₇MT2. The different modification numbers of cysteines in the [32-43] fragment (Table 1) and co-existence of several variously Zn(II)-depleted forms probably indicate high internal dynamics of Zn-S bonds, with their constant breakage and re-formation, as has been proposed in earlier studies.^{57-59,66} Another plausible explanation relates to the stability of particular metal binding sites. The fourth Zn(II) equivalent may be partially distributed amongst α - and β -domains, as was proposed in the literature,⁶³ which could explain the observed protection of certain β-domain regions from alkylation (Table 1).

Overall, our results regarding α -domain folding allowed us to conclude that Zn(II) binding to the α -domain is moderate sequential rather cooperative. The first two Zn(II) ions bind to independent ZnS_4 sites, the third forms the Zn_3S_9 cluster, and in the case of four Zn(II) ions the Zn₄S₁₁ cluster is formed with saturation of the α -domain. Association of the $Zn(\pi)$ ion with the β -domain before complete filling of the α -domain shows that there is no significant difference in the affinity of the weakest site in the α -domain and the strongest in the β -domain. This is in good agreement with a pH titration study performed on Zn7MT2, where one clear absorbance increase isotherm around pH 4-5 is observed.9,30 Analogous pH-titration of Cd₇MT2 shows a significant difference between α - and β-domains, which is observed as two clearly separated absorbance increase steps.9 Very early studies showed that the α -domain saturated with Cd(π) is much more stable than the β -domain,²⁶ which was more carefully analysed in recent years. Formation and dissociation of Cd7MT2 performed by bottomup and top-down ion-mobility mass spectrometry showed that Cd_4MT2 species are highly stable and all four $Cd(\pi)$ ions are located in the α -domain.^{67,68} A recent pH-variable study on MT1a supported the observation of Cd₄MT1a high stability even at low pH.¹⁴ The same study suggested that the Zn(II) binding process is significantly less cooperative and the Zn₄MT1a form is not as stable as the Cd(II) counterpart. Overall, our and the above-mentioned studies confirm the differences in $Cd(\pi)$ and $Zn(\pi)$ affinities towards the two respective domains in whole metallothionein.

$Zn(\pi)$ -Dependent folding of the isolated β -domain differs from that of whole MT2

Metal-induced β -domain folding should be expected to resemble the α -domain folding in terms of the formation of independent and clustered sites. However, their sequence motifs differ significantly from each other. Instead of the CXCC motifs of the α -domain, there are four CXC and two CXXC motifs in the β -domain sequence. The crystal structure of hepatic rat MT2 indicates a type-I reverse turn-like conformation with an internal NH···S^{γ} hydrogen bond for all CXC motifs. According to Stout and co-workers, this type of hydrogen bonding is expected to stabilize the local conformation in apo-MT2 and thereby facilitate metal ion binding.⁴ Indeed, as discussed above, three cysteine residues from region 1–20 are recalcitrant to modification after addition of four Zn(π) equivalents. This fragment contains two CXC motifs and indicates quite similar affinity to the weakest binding site in the α -domain.

Addition of a fifth molar equivalent to apo-MT2 sheds more light on the first $Zn(\pi)$ ion bound in the β -domain due to the lack of resolution and high similarities with the previous titration step in all experimental pathways (Table 1, Fig. 3e, f and Fig. S2e, f, ESI†). Most likely this ion is bound by three thiolate ligands from the 1–20 region with one from 26–31, the exact localization of which can be assumed based on the third ion in the whole cluster (Fig. 1b and 4e). Molecular dynamics simulations performed on partially saturated cadmium metallothionein showed that the first metal ion in the β -domain occupies

the site formed by Cys7, 13, 15 and 26, which agrees with our experimental and MD findings (Fig. 4e and 5a) and coordination mode in the 3D structure (Fig. 1).⁶¹ Comparison of the α -domain between the X-ray MT2 structure and MD results shows high structural similarity, which is a sign of a stable α -domain. The defined $Zn(\pi)$ binding site in the β -domain observed after addition of five Zn(II) equivalents causes a decrease in coordination dynamics in 32–43 and 44–51 regions in the α -domain. It may be partially explained by an equilibrium shift of a metal ion from $Zn_3\alpha Zn_1\beta MT2$ to $Zn_4\alpha Zn_0\beta MT2$ species, which again indicates similar Zn(II) affinities of both sites and high coordination dynamics of the weakest $Zn(\pi)$ ion in the β -domain. It should be noted that pronounced β-domain tryptic fragments also demonstrate a fully protected form (Table 1), and the signal intensity of the unmodified full protein in the MALDI spectrum is much higher in comparison with the previous titration step, which is a sign of co-existence of Zn₆MT2 and Zn₇MT2 species (Fig. 3e, f and Fig. S2f, ESI[†]). The high internal dynamics of Zn-S bonds and/or the presence of several partially Zn(II)-depleted MT species is maintained.

Although clustering is one possibility in handling the second $Zn(\pi)$ ion in the β -domain, our results obtained for MT2 with six Zn(II) equivalents show that the formation of another separate binding ZnS₄ site is much more preferred (Fig. 5b). Full protection from modification of all identified tryptic fragments (Table 1) indicates that at least seven cysteines from the β -domain are involved in Zn(II) binding. Therefore, the last cysteine that fills the coordination sphere should be either Cys21 or Cys24. Single modification of Cys21 in tryptic fragment [1-22] detected in the case of seven Zn(II)equivalents (Fig. 6) is strong evidence that the second metal ion in the β -domain is bound to the Cys24 residue. Moreover, molecular dynamics simulations strongly support this experimental result, as the $Zn(\pi)$ is coordinated by Cys24 while Cys21 remains free (Fig. 5b). It should be emphasized that the employed molecular dynamics methodology makes use of a semi-bonded approach, which in contrast to the bonded approach, is able to simulate coordination changes and the ligand exchange phenomenon. Moreover, H-bonding stabilization of the non-coordinated Cys21 residue by Ser6 is observed. Localization of Zn(II) ions in the β -domain presented in Fig. 4f is based on the structure and coordination bond networks in the full cluster. Our results therefore show that the Cys21 residue more likely is the only one free cysteine in Zn₆MT species. Utilization of the maximum possible number of cysteine residues in two separate binding sites in the β -domain is in agreement with previous studies indicating that there are virtually no free cysteines in Zn₆MT2 species.¹¹ This indicates that clustering does not occur until the last step of domain formation. Coordination of the last Zn(II) ion takes place by metal ion binding to Cys21 and subsequently to Cys5, 7 and 24. This binding mode causes rearrangement in the existing Zn-S bond network. Sulfur donors of Cys7, 15 and 24 become new bridging ligands. A similar conclusion came from differential modification of cadmium MT by radioactive IAA in a very early study.^{26,27} The authors postulated that the seventh

metal ion is bound to the 20-30 region, but they did not obtain single Cys resolution. Our results demonstrate one more important feature of the last binding event in the β -domain, namely its thermodynamic difference from the other two binding sites. Full protection from alkylation should be observed for Zn₇MT2 if binding sites were comparable, which is not the case (Fig. 3g and h). The singly modified peak observed for Zn7MT2 (Fig. 3h and 6) indicates partial dissociation of the last Zn(u) ion under alkylating conditions. Interestingly, Bernhard's report states that protection from modification of the N-terminal region of the β-domain causes distinctly lower affinity of the central region 20-30 for the metal ion.^{26,27} By identification of particular metal sites and lastly modified cysteine we may extend that conclusion forward. Binding of the first two $Zn(\pi)$ ions to independent sites located on the opposite tails of the β-domain lowers the protein affinity towards the third metal ion, probably due to the significant energy-requiring conformational changes needed for Zn-S rearrangement and Zn₃S₉ cluster formation. Interestingly, results on a $Zn(\pi)$ transfer process from Zn7MT2 to apo-SDH showed that dissociation of the first Zn(II) ion from the fully Zn(II)-loaded protein is structurally reversible to association of the last (seventh) Zn(II) ion to Zn₆MT2 species. Such a mechanism is not obvious because various energetic minima can be reached during association or dissociation of a metal ion. The same modification scheme observed here for the reaction forward and back indicates that the same Zn₆MT2 species is preferentially formed, which indicates its thermodynamic stability. In this case, the energetic cost related to the seventh Zn(II) ion binding is comparable with the energetic cost of its dissociation from Zn_7MT2 . Studies performed on Zn(II) transfer from Zn_7MT2 to PTP1B or fluorescent probes FluoZin-3 and Rhod-Zin-3, which bind Zn(n) with $K_d \sim 10^{-8}$ M show that $-\log K_{d1}$ of MT2 is the same as $\log K_{b7}$ strongly suggesting that this process is reversible.^{11,18} Moreover, the CD spectra recorded for Zn₆MT2 species obtained by association of 6 $Zn(\pi)$ ions to apo-MT2 or dissociation of the seventh Zn(II) from Zn7MT2 are identical and are not very different from the spectrum recorded for fully $Zn(\pi)$ -loaded protein, which indicates that dissociation and association of weakly bound $Zn(\pi)$ does not influence significantly the secondary structure of metallothionein, while being highly disordered.

The isolated β -domain demonstrates a more distinct Zn(II) coordination pathway than this domain in the whole MT2 protein. The first Zn(II) ion is bound rather weakly and probably delocalized throughout the whole β -domain peptide, as the MALDI spectrum shows 6 and 7 modifications (Fig. S13b, ESI†). Based on all the obtained results, it is reasonable to assume two, not mutually exclusive, possibilities: (i) all four Zn₀₋₃ β MT2 species are present in dynamic equilibrium, with prevalence of Zn₁ β MT2, and (ii) distribution of the first Zn(II) throughout all binding sites, with higher appearance frequency in the N-terminal part of the peptide. The obtained MS/MS results confirmed the presence and localization of the Zn₁ β MT2 form and localized its binding to Cys5, 7, 13 and 15. The MD results were in agreement with the experimental results, and showed

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that Cys5 was H-bonding stabilized by Lys22, in contrast to Zn_5MT2 , where $Zn(\pi)$ bound in the β -domain resulted in Cys5 free coordination (Fig. S14b, ESI†). Therefore, the data obtained for the isolated domain are contradictory to those obtained for the whole protein and do not correspond to any site in the 3D structure (Fig. 1b).

Addition of the second Zn(II) equivalent to the isolated B-domain caused a significant equilibrium shift towards $Zn_2\beta MT2$ and $Zn_3\beta MT2$ species, but the results did not allow the $Zn(\pi)$ position to be sharply elucidated in the $Zn_2\beta MT2$ form. However, there is indirect evidence for the formation of a Zn₂S₇₋₈ form, with one modified Cys from the 1-20 region and possibly another one from the 25-31 region (Table S8, ESI⁺). Unique behaviour of the Cys29 residue was postulated by Babu and co-workers in a study on Zn(II) release from the isolated β-domain of rat MT2.⁶⁹ They proposed that Cys19 and Cys29 release Zn(II) faster than any other cysteine residues with concurrent formation of a Cys19-Cys29 disulfide bridge. Interestingly, those residues are exposed to the solvent in the isolated β -domain, but are buried in the whole protein MT2.67 Our molecular dynamics indicated that Cys5 and Cys29 are free of coordination (Fig. S14c, ESI[†]) in which the first one interacts with Gln23 via H-bonding (2.6 Å).

Lack of any modifications after addition of the third Zn(II)equivalent to apo-BMT2 demonstrates a more stable cluster in the isolated β -domain than in the whole protein and points to the conclusion that the isolated domain is not a good model of the whole protein. The single peak in the MALDI spectrum corresponding to the unmodified β -domain (Fig. S13d, ESI[†]) points to the absence of weakly coordinating cysteine and supports our statement. Lower stability of the seventh Zn(n)site in metallothioneins and the difference between isolated domains and the whole protein molecule have been shown several times in the past.^{11,12,18,70,71} For example, the higher affinity of $Zn(\pi)$ ions in the isolated β -domain was confirmed in its competition with PAR (4-(2-pyridylazo)resorcinol).⁷⁰ The most likely reason for such behaviour is higher main chain flexibility due to lack of stabilizing interdomain contacts, and lack of possible conformational restraints due to the presence of the α -domain.

The mechanism of $\text{Zn}(\pi)$ and $\text{Cd}(\pi)$ binding to metallothionein is entirely different

To date, our knowledge regarding the structure, chemical and biophysical properties of metallothioneins has been mostly based on studies performed with cadmium metallothionein. Historically, it is associated with the fact that metallothionein was discovered in equine kidney cortex as a cadmium protein.¹ Another reason why the vast majority of metallothionein studies use this ion is that Cd(π) produces well-defined spectroscopic signals, in contrast to spectroscopically silent Zn(π) ions. Usage of Cd(π) ions allows one to study association, thermodynamic and kinetic properties, as well as the complex structure of metallothionein. However, it is now well known that metallothioneins are Zn(π) and Cu(π) metalloproteins in healthy living organisms.^{72–76} Comparison of NMR data obtained for

zinc and cadmium metallothioneins in the early 1990s shows that both molecules are structurally similar in terms of cluster geometry, metal–sulfur bonds and overall shape, although some differences in intraprotein connectivities and protein volume were detected.^{77,78} Differences between these metal ions include (i) their ionic radii (109 pm for Cd(π) and 88 pm for Zn(π)), (ii) hard and soft acid–base properties – Cd(π) is a soft Lewis acid while Zn(π) is considered borderline,⁷⁹ and (iii) formed M–S coordination bond lengths (averages between terminal and bridged metal centres: Cd–S ~ 2.5 Å, Zn–S ~ 2.3 Å).^{80,81}

To date, there are a limited number of reports focusing on the biophysical properties of zinc metallothioneins, although it is the most abundant MT form in mammals.⁷⁶ Various physicochemical studies including CD, MCD, UV-Vis, and NMR spectroscopy on mammalian metallothioneins show that $Cd(\pi)$ binds more tightly than $Zn(\pi)$ ions, which is well understood when considering the enthalpy of M-S bond formation and highly similar structure.^{9,12} A number of spectroscopic studies have demonstrated that the Cd(II) binding affinity in the α -domain ($K_{\rm d} \sim 10^{-15}$ M) is almost three orders of magnitude higher than in the β -domain ($K_d \sim 10^{-12}$ -10⁻¹³ M).^{9,82-84} Contrastingly, many reports state that all seven Zn(II) ions have identical or almost identical protein affinity and both clusters are basically indistinguishable in terms of their stability, with an approximate dissociation constant of $\sim 10^{-12}$ M (for MT1 and MT2).9,10,24,83,85 Several recent studies on MT2 and MT3, however, demonstrated otherwise, namely that one Zn(II) ion is bound relatively weakly with $K_{\rm d} \sim 10^{-8}$ –10⁻⁹ M, two other ions are bound with medium affinity with $K_{\rm d} \sim 10^{-10}$ – 10^{-11} M, and the remaining four ions are bound with indistinguishable high affinity of $K_{\rm d} \sim 10^{-12}$ M.^{11,12} These discrepancies in the measured affinities of Zn(II) ions to metallothioneins are a current subject of debate.12,76,86

The results presented in this article support a model with varied Zn(II) affinities towards MT2. Binding of Zn(II) to the α -domain is sequential, whilst Cd(II) seems to be cooperative, as it is known from the literature. Observed ¹¹³Cd or ¹¹¹Cd NMR signals corresponding to α -cluster formation in NMR titration studies indicate identical affinities.^{25,87,88} On the other hand, corresponding ESI-MS titration studies showed that intermediate forms (Cd₁₋₃MT2) are also formed; however, Cd₄MT2 is highly favourable.^{29,67} It is worth noting that exact identification of successive Cd(II) binding events and sites in the α -domain has not been performed to date and such studies could possibly reveal differences or the lack of differences in Cd(II) affinities toward the α -domain. Data collected to date show that both metal ions bind to the α -domain specifically with high affinity, but Zn(II) binding is more sequential.

While the binding of $Zn(\pi)$ and $Cd(\pi)$ to the α -domain is only slightly different, binding of these two ions to the β -domain is much more divergent. The most visible difference besides the fact that $Cd(\pi)$ ions are bound more tightly than $Zn(\pi)$ ions is the clear differentiation of $Zn(\pi)$ affinities in the β -domain compared to $Cd(\pi)$.⁸⁹ This phenomenon is observed in pH-dependent binding of both metal ions to metallothioneins regardless of protein isoforms.^{9,25,82–84} ¹¹³Cd-NMR studies performed on whole

protein show that $Cd(\pi)$ binding to the β -domain is cooperative, although the coordination dynamics of the cluster is much more pronounced compared to the α -cluster.²⁵ However, pH titrations clearly show that the slope of Cd(n) binding to the β -domain is significantly lower compared to the α -domain. This is due to a lower number of protons released during coordination of three ions. It also reflects more pronounced differentiation of Cd(II) affinities towards the β -domain than the α -domain. Such differentiation was observed for MT4⁸⁴ and is less pronounced for MT3⁸¹ and least visible for MT1 and MT2 isoforms.9,82,83 This shows that the relative affinities of metal ions in the β -domain vary depending on the metallothionein isoform. It should be noted that several reports in the past also reported the presence of weaker bound Cd(II) ions in MTs. These include differential modification of cadmium MT using IAA,^{26,27} NMR studies on Cd(II) release from MT by EDTA^{85,87} and ICP-MS studies on the displacement of zinc MT with Cd(II) ions.⁹⁰

Our results on the mechanism of Zn(II) association with MT2 also shed new light on $Zn(\pi)/Cd(\pi)$ exchange in this protein. The addition of Cd(II) or Cd7MT to Zn7MT causes preferential displacement of $Zn({\rm II})$ ions in the $\alpha\text{-domain}$ and $^{113}\text{Cd-NMR}$ studies show that this process is highly cooperative.^{91,92} If the affinity of the remaining three $Zn(\pi)$ ions in the β -domain were close to each other, then Zn(II)/Cd(II) exchange should indicate cooperative character. Despite this, mixed Zn/CdMT complexes are preferentially formed in vivo and in vitro, although their compositions differ. Such formation is possible when Zn(II) ion affinities towards their sites in the β -domain vary significantly, which is the conclusion from our study. The crystal structure (Fig. 1) shows localization of $Cd(\pi)$ ions in the β -domain in position IV, which indicates that the exchange of the fifth $Zn(\pi)$ ion occurs exactly in this position, where the metal is bound by bridging Cys7 and Cys24 residues and terminal Cys21 and Cys5. It is reasonable that the fifth equivalent of $Cd(\pi)$ displaces the weakest bound $Zn(\pi)$ ion in MT. The pathway of $Zn(\pi)$ metalation (Fig. 4) identified in the course of this study shows that Cys21 is the residue that more likely terminates cluster formation and is bound to the weakest $Zn(\pi)$ in the β -domain. A similar conclusion comes from the competition between Zn₇MT2 and apo-SDH, for which Cys21 and Cys24 residues were identified to be the weakest bound to the metal ion (Fig. 8). It should be mentioned that the composition of metal clusters in MT is believed to be a consequence of metal ion selectivity in the protein, according to which the major factor governing the cluster type is the protein structure perturbation due to the cluster volume variations. Vašák and co-workers state that metal-thiolate affinity is important in the folding process, while size-match selectivity is the dominant factor in the metal-loaded protein.93 In fact, the cluster composition strictly depends on the mechanism of the formation and conditions under which it is formed. The pathway of its formation is associated with the substrates' stoichiometry, reaction kinetics and thermodynamics. Overall, our results give a new, higher resolution picture of Zn(II)/Cd(II) exchange and metal binding processes.

Buffering properties of metallothionein are related to sequential binding of Zn(n) and the presence of Zn(n)-depleted species

Data presented here show that although huge efforts have been made to understand the $Zn(\pi)$ binding pathway in MT2, it is not fully defined, leaving the possibility for the existence of several species with the same number of metal ions per molecule. Moreover, the metal binding process may depend on several factors such as protein concentration or slightly different pH.14,94 However, our results are the first ones obtained with such a resolution and focused on biologically relevant zinc metallothionein-2. Sequential binding of Zn(u) in the β -domain supports the idea of $Zn(\pi)$ affinity differentiation, pointing out the different biological function of the β -domain compared to the α-domain, which has been indicated based on MT evolution.⁹⁵ According to this, the α -domain seems to function as a storage domain, and the metal binding process is highly efficient and occurs with similar affinity of Zn(II) ions. However, the data obtained so far show that Cd(II) binding to this domain is even more efficient, which is reasonable when we consider the major role of this domain in cellular detoxification. The β-domain plays another, regulatory function according to which it distributes $Zn(\pi)$ among other $Zn(\pi)$ -dependent or regulated proteins. Variously lowered affinities of Zn(II) ions bound in the β -domain place it either as a Zn(II) donor for other metal-depleted proteins or as an acceptor which allows binding of excess $Zn(\pi)$ when it exists in partially saturated species. Cellular free Zn(II) changes and dissociation constants of Zn(II)ions bound in the β -domain (from 10^{-8} to 10^{-11} M) match each other very well, which allows metallothionein to be considered as a major component of the cellular zinc buffer.¹¹ It is even more understandable when one takes into account the fact that this protein is easily inducible under several stimuli such as metal ions, glucocorticoids, and oxidative stress.⁹⁶⁻⁹⁸ Therefore, induction of metallothionein guarantees the binding of excess metal (biogenic and toxic) ions in such a way that the β -domain is still able to function as a Zn(II) regulatory module.

The Zn(II)-binding pathway in metallothionein-2 proposed here is a relation between the structural properties of the Zn(II)depleted species and the buffering properties of this protein. So far, our knowledge regarding partially depleted forms of MT is very limited, besides some information that the β-domain binds Zn(II) less efficiently than the α -domain. On the other hand, some reports stated that although $Zn(\pi)$ binding is sequential, the affinities of particular Zn(II) ions almost do not differ.²⁴ It is, however, in contrast with other experiments, where MT functions as a Zn(II) donor towards metal-depleted enzymes, and inhibitory and structural sites in proteins.^{11,12,18,20,52,70,99} By the combination of the high-resolution Zn(II)-binding process determined here together with molecular dynamics simulations and thermodynamic studies performed so far, we are able to describe structures of partially saturated species of zinc metallothionein for the first time. One needs to note that normal concentrations of freely available Zn(II) support the existence of Zn5MT2 and Zn6MT2 species.¹¹ Therefore the understanding of this process is not only important for exploration of the structural nature of this enigmatic protein, but also critical for understanding the principles of other metallothionein functions, which we are still trying to understand 60 years after its discovery.⁷⁶

Conclusions

In this study, by the combination of mass spectrometry approaches to partially saturated and alkylated human metallothionein-2 together with molecular dynamics study, we analysed the Zn(II) binding process with high resolution, which allowed us to propose the structure of Zn1-6MT species with respect to metal ions' localization and their coordination spheres. The results show that all metallothionein species are highly dynamic even if they are clustered and the proposed structures could be a representation of one possibility, however the most prevalent under the used conditions. Our data show that the four Zn(n) ions bind preferentially to the α -domain when they are added to the apo-MT2 form and then fill the β -domain. However, this process is much less distinct than in the case of cadmium MT, which shows some thermodynamic similarities in coordination with the fourth and fifth Zn(II) ion. The binding of all $Zn(\pi)$ ions is sequential, with high preference for the formation of independent ZnS4 sites in early stages of domain formation. However, the $Zn(\pi)$ binding process in both domains differs in later stages of cluster formation, indicating more distinct Zn–S network rearrangement in the β -domain upon binding of the seventh $Zn(\pi)$ ion. This process is interpreted as an energetic reason for significantly lower affinity of the last Zn(II) ion in the whole domain, which is in contrast to the Zn(II) binding pathway in the isolated β -domain. Our results have shown that Cys21 is more likely the last coordinating cysteine residue, which demonstrates that weakly bound $Zn(\pi)$ in the β -domain corresponds to position IV (Fig. 1b). The same conclusion is reached when MT2 serves as a $Zn(\pi)$ donor and supports a catalytic cofactor to $Zn(\pi)$ -depleted sorbitol dehydrogenase. Then the same Cys21 residue is modified first, which shows that the Zn(II) binding and dissociation process of the weakest Zn(II) ion is fully reversible, which has also been demonstrated by fluorescent zinc probes in a previous investigation.¹¹ The presented study sheds more light on the mechanism of zinc metallothionein-2 cluster formation with the highest resolution to date and should be useful for the investigation of metal binding and dissociation processes in other isoforms and metallothioneins from other organisms. By the comparison of various MS approaches with MD calculations, and $Zn(\pi)$ release and binding equilibria in water media with other structural and thermodynamic data we provided a new opportunity to look into the function of various metallothionein proteins with high indication of the role and importance of partially or differently metal-loaded species, in which interest is constantly growing.

Conflicts of interest

There are no conflicts to declare.

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