Metal- and Affinity-Specific Dual Labeling of Cysteine-Rich Proteins for Identification of Metal-Binding Sites

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ABSTRACT: Here, using human metallothionein (MT2) as an example, we describe an improved strategy based on differential alkylation coupled to MS, assisted by zinc probe monitoring, for identification of cysteine-rich binding sites with nanomolar and picomolar metal affinity utilizing iodoacetamide (IAM) and N-ethylmaleimide reagents. We concluded that an S_n2 reaction provided by IAM is more suitable to label free Cys residues, avoiding nonspecific metal dissociation. Afterward, metal-bound Cys can be easily labeled in a nucleophilic addition reaction after separation by reverse-phase C18 at acidic pH. Finally, we evaluated the efficiency of the method by mapping metal-binding sites of Zn_{−}−MT species using a bottom-up MS approach with respect to metal-to-protein affinity and element(al) resolution. The methodology presented might be applied not only for MT2 but to identify metal-binding sites in other Cys-containing proteins.

Approximately one-third of human genes encode proteins that bind metal ions, and in around 10% of the proteome, Zn^{2+} ions are used as a catalytic, structural, and regulatory cofactor. Metallothioneins (MTs) play a role in homeostatic control of Zn^{2+} and Cu^{+} ions in cellular signaling and transduction networks by muffling reactions. They bind metal ions and serve as both metal donor or acceptor, controlling the cellular Zn^{2+} fluctuations in the cytosol, nucleus, or mitochondria. Mammalian MTs are low molecular mass (6−7 kDa) cysteine-rich molecules that bind multiple metal ions in a multiple tetrathiolate coordination environment within two separate M_S clusters. In humans, there exist at least a dozen MT proteins, categorized into subfamilies (MT1−4 isoforms and MT1 subisoforms) depending on sequence, tissue localization, function specificity, or metal-binding properties. Cysteine (Cys) is the most nucleophilic amino acid residue, commonly binding essential as well as toxic metal ions. Moreover, the Cys residue is a cellular target for reactive oxygen, nitrogen, and sulfur species, and it is posttranslationally modified in S-methylation and S-linked acylation, among other reactions. Thus, Cys acts in multiple proteins as a redox switch, depending on the oxidative molecules and metal ion concentration. Because of the aforementioned relevance, a range of experimental and theoretical tools has been developed aimed at identifying different Cys residue states in proteomes. Most of the chemical tools are based on the nucleophilic reaction of Cys toward thiol-specific probes, which may exhibit different reactivity, enabling differentiation of the cysteine sulfur state. Some of these common protein thiol probes are iodoacetamide (IAM), iodoacetic acid (IAA), N-ethylmaleimide (NEM), methyl methanethiosulfonate (MMTS), Cys-reactive mass tag (cyss-TMT), and p-benzoquinone (Bq). Among them, IAM and NEM are commonly used, forming covalent products with sulfhydryl groups by an S_n2 nucleophilic substitution or by nucleophilic Michael addition, respectively. NEM reacts with thiols faster than IAM and also exhibits a wider range of pH applicability. It shows an appreciable rate even at acidic pH, whereas IAM requires neutral or basic pH. However, NEM exhibits less specificity, which gives rise to side reactions with histidine and lysine residues when NEM is used at large excess or basic pH. On the other hand, IAM is preferred since it forms a very stable thioether, whereas NEM might undergo partial ring opening.

Received: April 14, 2020
Accepted: August 3, 2020
Published: August 3, 2020
step and a second alkylation.\textsuperscript{25} Other derived strategies are based on a dual parallel experiment in which a native protein and the treated protein are both labeled by the alkylation reagent.\textsuperscript{31,37} For instance, if one is interested in localizing metal-binding Cys residues, the protein is treated by removing the metal ion by chelation or reducing the pH to promote metal dissociation and then labeling.\textsuperscript{37−39} Subsequently, by comparing both experiments, the native metalloprotein labeled and the treated protein, one might inquire into a particular Cys residue participating in metal ion binding.\textsuperscript{31} This methodology coupled to state-of-art mass spectrometry (MS) techniques has been successfully used in the past in Cd\textsuperscript{2+} and Zn\textsuperscript{2+}-containing MT studies.\textsuperscript{31,35,40,41} Despite the efforts and advances made, these methods still present several limitations in metalloproteomic studies.\textsuperscript{42} In principle, a metal-bound thiolate would decrease its reactivity toward nucleophiles and thus no modification would occur under controlled conditions because it is metal-protected.\textsuperscript{24,40,43} This is a fact for a kinetically inert metal–Cys bond.\textsuperscript{44} For instance, 2 out of 10 Cys residues from the p53 core domain were found to be reactive toward NEM.\textsuperscript{45} Identification of the Cys positions revealed that they do not belong to the Zn(Cys)\textsubscript{5}((His)\textsubscript{2}) core, which remained unmodified.\textsuperscript{51} Not only the kinetic stability but also the thermodynamic stability of the metal–protein complex dictates how the reaction proceeds. In the presence of the same ligand, properties of the metal ion influence the thermodynamic stability of the protein complex. For instance, Cd\textsuperscript{2+}, which in terms of the hard–soft acid–base concept is significantly softer than Zn\textsuperscript{2+}, is bound more strongly to Cys sites and more weakly to the Cys\textsubscript{5}His\textsubscript{2} core.\textsuperscript{46} Therefore, proteins with a low or moderate Zn\textsuperscript{2+} binding affinity (log $K_{d}$ of 7−10) and kinetically labile metal–protein bonds, such as MT2, might reveal other scenarios.\textsuperscript{47} The alkylation concentration and time of reaction should be particularly optimized to avoid Zn\textsuperscript{2+} dissociation from the reactive Zn\textsuperscript{2+}-bound Cys residue. Unlike traditional differential alkylation, a similar reactivity might appear for a metal-bound Cys and for a free Cys residue with a low $pK_a$, which hinders their nucleophilic differentiation.\textsuperscript{23−25} The local electrostatic environment will affect the acidity of Cys residues present in the protein, dictating the reactivity toward the electrophile.\textsuperscript{48} This might lead to situations in which an M(Cys)\textsubscript{x} core is alkylated without metal dissociation. The situation where a reactive Cys residue participates in coordination of both a metal ion and an alkylation moiety has been previously observed.\textsuperscript{28,31,49} Three possible scenarios need to be contemplated: (i) alkylation of a free Cys residue, (ii) alkylation of a metal-bound Cys residue without metal ion dissociation, and (iii) alkylation of a metal-bound Cys residue and subsequent metal ion dissociation. Therefore, finding a reagent or reaction conditions that distinguish between free Cys and metal-protected Cys residues is a fundamental issue. Moreover, high-resolution methods such as MS are greatly needed to follow the modification extent but also to identify metal-binding sites.\textsuperscript{50,51} Using soft electrospray ionization (ESI) conditions and transmission parameters, native conditions might be preserved during ESI-MS analysis, maintaining the noncovalent metal–protein interactions, the solution-phase populations, and the conformational states for the labeled metal-bound protein.\textsuperscript{52−56} During the last 2 decades, a vast number of research studies have applied ESI-MS for MT investigations.\textsuperscript{28,31,37,40,57−59} To map the modification sites, top-down sequencing provides an easy and fast way, although it still provides a lower sequencing coverage than bottom-up approaches.\textsuperscript{60,61}

Here, we describe an improved strategy based on differential alkylation coupled to MS for identification of multi-Cys metal-binding sites with nanomolar and picomolar metal affinity in MT2 by utilizing IAM and NEM reagents. First, we studied the kinetic and thermodynamic lability of Zn\textsuperscript{2+}– and Cd\textsuperscript{2+}–thiolate bonds in Zn-MT2 and Cd-MT2 using both alkylators and analyzed them by UV–vis, matrix-assisted-laser desorption ionization mass spectrometry (MALDI-MS), and ESI-MS methods. Most biophysical research has used Cd\textsuperscript{2+} for MT study since it produces well-defined spectroscopic signals, in contrast to spectroscopically silent Zn\textsuperscript{2+}.\textsuperscript{31} Notably, MTs were originally isolated as a mixed Zn\textsuperscript{2+}/Cd\textsuperscript{2+} complex.\textsuperscript{6} For our purposes, MT2 was used as a model of a protein with various low, moderate, and high Zn\textsuperscript{2+} binding affinities. It is worth noting that the first three Zn\textsuperscript{2+} ions dissociating from MT2 demonstrate nano- and subnanomolar affinity, perfect for this investigation. Comparing the reactivity with the more thermodynamically stable Cd-MT2 protein (low picomolar affinity) illustrated the issue regarding the metal–protein stability and the phenomena of metal dissociation. Through this, we ranked and ordered the reactivity, allowing us to develop a differential alkylation strategy to map partially Zn\textsuperscript{2+}-loaded Zn\textsubscript{x−y}MT2 species. We demonstrated that IAM is more suitable than NEM to be used as the first labeling reagent in order to label free Cys residues. The lower reactivity of IAM prevents the metal ion dissociation, whereas its small size and hydrophilic character allow access to buried free thiols.\textsuperscript{40,62} This step was followed by metal ion removal with acidification and a subsequent second labeling reaction with NEM. Although we and others demonstrated that addition of NEM without acidification may dissociate all seven Zn\textsuperscript{2+} from Zn-MT2, different conclusions are found for Cd-MT2.\textsuperscript{28,30} NEM did not dissociate all seven Cd\textsuperscript{2+} due to the higher Cd\textsuperscript{2+}–to-protein affinity. Another factor that may limit the metal ion dissociation with addition of NEM without acidification is the thiol accessibility. The low pH greatly facilitates the access of NEM to all of the Cys residues, independently if they were buried or solvent-exposed Cys residues.\textsuperscript{24} For this purpose, NEM offers faster kinetics than iodoacetamide derivatives, even at low pH.\textsuperscript{26,27} Thus, in order to elaborate a general method that may be applicable for other Cys-containing proteins, with higher metal-to-protein affinity and with different thiol accessibility,\textsuperscript{63,64} we introduced an acidification step prior to NEM labeling. Altogether, this ensures metal ion removal independent of how strongly they are bound and facilitates the access of NEM to the thiols. Afterward, utilizing the differential alkylation developed, we identified the Zn\textsuperscript{2+}-binding Cys residues in Zn\textsubscript{x−y}MT2 species with a bottom-up MS approach. The methodology presented herein represents a general technique to study metal-binding Cys-containing proteins, independently of their metal-binding properties and protein structure.

**EXPERIMENTAL SECTION**

**Materials.** All reagents used were purchased from Sigma-Aldrich (Merck Group), Acros Organic, Roth, BioShop, VWR International (Avantor), and Iris-Biotech GmbH. The pH buffers and solutions were prepared with Milli-Q water (Merck Millipore), incubated with Chelex 100 resin (Bio-Rad) and degassed to eliminate trace metal ion contamination. For more
in the presence of 1 mM TCEP, followed by bu
50 mM ammonium acetate (pH 7.4) and puri
purging nitrogen, and addition of 1 mM TCEP at each round. detailed information may be found in the SI.

Avoid oxidation of the free thiols,37 using a low capillary voltage
nitrogen blanket and analyzed by ESI-MS. Care was taken to
kDa Amicon Ultra-4 Centrifugal Filter for 10 min under a
modi
previously degassed by purging with nitrogen. To measure the
oxidation of thiols, as con
ity).31 Under our experimental conditions, we did not observe
tightly than DL-dithiothreitol (DTT) (submillimolar a
We used TCEP as a reducing agent since it binds zinc less
Labeling Strategy. Zn7MT2 was puri

Overview of the hereby presented differential labeling approach. (A) Steps of the procedure for labeling free- and metal-bound Cys residues by IAM and NEM, respectively. (B) Human metallothionein-2 (MT2) sequence with indicated tryptic fragments and position numbers of Cys residues. Arrows indicate the cleavage positions of the C-terminal lysine, specific for trypsin enzyme.

Identification of Zn2+-Binding Sites by a Dual Labeling Strategy. Zn-MT2 was purified by size exclusion chromatography with 10 mM HCl, and the concentration of apoMT was estimated by 5,5′-dithiobis(2-nitrobenzoic acid) and Cd2+ titration experiments. Then, apoMT was saturated with 4, 5, 6, or 7 mol equiv of ZnSO4 under a nitrogen blanket with 4, 5, 6, or 7 mol equiv of ZnSO4 under a nitrogen blanket in the presence of 1 mM TCEP, followed by buffer exchange to 50 mM ammonium acetate (pH 7.4) and purification with a 10 min spin time (three times) using 3 kDa Amicon filters, purging nitrogen, and addition of 1 mM TCEP at each round. Partially metalated proteins were subsequently analyzed by ESI-MS. Then, a 15 μM Zn7-MT2 aliquot was incubated in 1 mM IAM (15 min, 25 °C) in darkness. From this, an aliquot was purified by ZipTip μ-C18, as done previously, and measured by MALDI-MS. Another aliquot was analyzed by ESI-MS. The rest of the sample (80 μL) followed the dual-labeling strategy. First, the pH was reduced with 0.1% formic acid (FA) and 1 mM DTT and the excess IAM removed by purification with C18 resin. Eluted protein was double-labeled with incubation in 3 mM NEM (30 min, 25 °C). After that, an aliquot was analyzed by MALDI-MS. The rest of the sample underwent a bottom-up MS approach by digestion in a solution using trypsin at a weight ratio of 1:20 (30 min, 37 °C). The trypsinization reaction was quenched by addition of 5 μL of 0.1% FA.

UV–Vis Spectroscopy. UV–vis spectra were recorded on a JASCO V-650 spectrophotometer at 25 °C with a 1 cm quartz cuvette. The Zn-MT2 and Cd-MT2 reactions with NEM and IAM were followed at 492 nm using 100 μM Zincon (ZI) or 4-(2-pyridylazo)resorcinol (PAR) in 20 mM Tris-HCl (pH 7.4, NaCl 150 mM) buffer, respectively.65,66 To avoid oxidation, 1 mM TCEP was used as a non-metal-binding reducing agent. The reactions were followed at 618 and 492 nm, respectively, for a period of 0–24 h.

ESI-MS. ESI-MS experiments were performed on a quadrupole time-of-flight (qTOF) Bruker Maxis Impact mass spectrometer (Bruker Daltonik GmbH, Bremen, Germany) calibrated with a commercial ESI-TOF tuning mix (Sigma-Aldrich). Samples were directly infused with a 1 μL/min flow rate. ESI-MS spectra were recorded in positive mode with a capillary voltage of 2 kV, end plate offset potential of 500 V, nebulizer gas (N2) pressure of 1.5 bar, drying gas (N2) flow rate of 4 L/min, and drying temperature of 180 °C. The mass range was set from 500 to 3000 m/z and recorded and averaged over 1 min. The Bruker Compass data analysis
The first part of this research attempted to study the kinetic and thermodynamic lability of Zn\(^{2+}\)-thiolate bonds in fully saturated Zn-MT2 and to compare it with its cadmium counterpart. To do so, zinc and cadmium proteins were incubated with increasing concentrations of both alkylation reagents NEM and IAM for different periods of time. The reaction was then simultaneously followed by native ESI-MS and MALDI-MS (Figure 1A). Complementarily, UV–vis spectrophotometric studies were carried out using chromophoric chelating probes ZI and PAR, respectively, to monitor the Zn\(^{2+}\) and Cd\(^{2+}\) dissociation upon protein modification. ZI forms a weak ZnZI complex with a dissociation constant of \(K_d = 2.09 \times 10^{-6}\) M at pH 7.4\(^{68}\) that ensures no Zn\(^{2+}\) competition with Zn-MT2 protein. Because Cd\(^{2+}\) binds less tightly to ZI than Zn\(^{2+}\), we could apply PAR to monitor the Cd\(^{2+}\) dissociation upon alkylation in a more quantitative way without Cd\(^{2+}\) competition with Cd-MT2.\(^{52,56}\) No doubt, ESI can retain the native-like structures, possibly because of the kinetic trapping effect.\(^{28}\) However, complementing gas-phase MS and solution experiments might require caution to be taken due to concentration-dependent rate limitations, differences in the solvent composition, or pH-induced changes.\(^{59}\) However, general trends and conclusions obtained from both gas-phase and solution experiments matched very well, as will be presented in the next section.

### RESULTS AND DISCUSSION

The oxidative alkylation and subsequent metal dissociation from Zn\(_7\)MT2 and Cd\(_7\)MT2 followed by MALDI-MS, ESI-MS, and UV–Vis Experiments at Increasing Concentrations of IAM and NEM Alkylating Reagents

| equiv protein | species (ESI-MS) | metal ions transferred (UV–vis) | Cys residues modified\(^a\) (MALDI-MS) | species (ESI-MS) | metal ions transferred (UV–vis) | Cys residues modified\(^b\) (MALDI-MS) |
|--------------|-----------------|-------------------------------|--------------------------------------|-----------------|-------------------------------|--------------------------------------|
| 15 Cd-MT     | Cd\(_6\)NEM\(_{b-1}\) | 0.4                           | 0.5                                  | Cd\(_6\)IAM\(_{b-1}\) | 0.0                           | 0.5                                  |
| Zn-MT        | Zn\(_6\)NEM\(_{b-1}\) | 2.5                           | 1.0                                  | Zn\(_6\)IAM\(_{b-1}\) | 0.0                           | 0.6                                  |
| 40 Cd-MT     | Cd\(_5\)NEM\(_{b-3}\) | 1.2                           | 2.5                                  | Cd\(_5\)IAM\(_{b-3}\) | 0.0                           | 0.5                                  |
| Zn-MT        | Zn\(_5\)NEM\(_{b-11}\) | 3.0                           | 10.0                                 | Zn\(_5\)IAM\(_{b-11}\) | 0.5                          | 1.0                                  |
| 180 Zn-MT    | Cd\(_4\)NEM\(_{b-1}\) | 1.3                           | 7.5                                  | Cd\(_4\)IAM\(_{b-1}\) | 0.3                          | 2.5                                  |
| 750 Zn-MT    | Cd\(_3\)NEM\(_{b-12}\) | 2.3                           | 11.0                                 | Cd\(_3\)IAM\(_{b-12}\) | 0.3                          | 7.0                                  |
| Zn-MT        | Zn\(_3\)NEM\(_{b-20}\) | 6.7                           | 19.0                                 | Zn\(_3\)IAM\(_{b-20}\) | 2.4                          | 14.0                                 |
| 1400 Cd-MT   | Cd\(_2\)NEM\(_{b-11}\) | 2.7                           | 12.5                                 | Zn\(_2\)IAM\(_{b-11}\) | 1.2                          | 13.0                                 |
| Zn-MT        | Zn\(_2\)NEM\(_{b-20}\) | 6.8                           | 19.5                                 | Zn\(_2\)IAM\(_{b-20}\) | 3.5                          | 14.5                                 |

*Refer to the average number of Cys residues modified. The bolded text represents the main species presented in Scheme 1.*

**MALDI-MS.** A MALDI-TOF/TOF MS Bruker UltrafleXtreme instrument (Bruker Daltonik GmbH, Bremen, Germany) was used for MALDI-MS experiments. 2,5-Dihydroxybenzoic acid (DHB) and \(\alpha\)-cyano-4-hydroxycinnamic acid (HCCA) were used as the MALDI-TOF matrix for protein and peptide analysis, respectively. The saturated matrix solution was prepared in 30% ACN and 0.1% trifluoroacetic acid. MALDI-MS analysis of proteins was performed in a linear positive mode in the 2–20 kDa range. The mass spectra were typically acquired by averaging 2000 subspectra from a total of 2000 laser shots per spot. The laser power was set at 5–10% above the threshold. The calibration was done using a standard peptide and protein calibration mixture obtained from Bruker (Bruker Daltonik GmbH, Bremen, Germany). MALDI-TOF/TOF measurements of peptides were performed in reflector positive mode in the 0–4 kDa range. Moreover, a LIFT cell was used for MS/MS analysis of detected peptides.\(^{67}\) The instrument was controlled by flexControl ver. 3.4 and flexAnalysis ver. 3.4 software. BioTools 3.2 SR3 and Sequence Editor (Bruker Daltonik GmbH, Bremen, Germany) were used to analyze the MALDI-MS data.
been dissociated (Figure 2A). The reaction intermediate \( \text{Zn}_4\text{NEM}_9 \beta \) concomitant full consistent with the 10 Cys being NEM-labeled, suggesting a agents NEM and IAM on the Cys residue modi... 3600 equiv) of alkylating agents NEM and IAM on the Cys residue modification (A and B, respectively) monitored with MALDI-MS and the metal dissociation from Zn\(_7\)MT\(_2\) and Cd\(_7\)MT\(_2\) analyzed by UV−vis spectrophotometry (C and D, respectively). MALDI-MS and UV−vis spectra were recorded after 60 and 200 min of sample incubation at 25 °C. Metal ion dissociation was analyzed using Zincon and PAR assays.59,60 Red and blue stand for Zn\(_7\)MT\(_2\) and Cd\(_7\)MT\(_2\), respectively. Cd\(_{2+}\) and Zn\(_{2+}\) refer to Zn\(_7\)MT\(_2\) and Cd\(_7\)MT\(_2\) proteins.

NEM with Zn\(_7\)MT\(_2\) were observed in the gas phase and in solution (Table 1). Meanwhile, ESI-MS captured the intermediate Zn\(_7\)NEM\(_{11}\)MT\(_2\), and approximately 2.5 Zn\(_{2+}\) dissociated rapidly in solution (Table 1 and Figure S4 (SI)). Its Cd\(_{2+}\) counterpart (Cd\(_{7}\)NEM\(_{11}\)MT\(_2\)) appeared and remained the most abundant, even at large molar excess, which clearly shows the higher Cd\(_{2+}\)-to-protein affinity. On the other hand, less reactive IAM required at least 180 equiv to provoke Zn\(_{2+}\) dissociation and to form the intermediates Zn\(_4\)IAM\(_{9,11}\)MT\(_2\). In disagreement, UV−vis experiments showed that only one Zn\(_{2+}\) dissociated from the protein. Note that the spectroscopic signal obtained is the average response from dynamic equilibria between multiple species. This hypothesis is supported by the Cys modification profile that showed a broad number of modifications (Figure 2B). Increasing to 750 equiv of alkylator leads to the formation of the intermediate Cd\(_4\)IAM\(_{10}\)MT\(_2\) and the species Zn\(_9\)IAM\(_{8,11}\)MT\(_2\) that were metal-stripped after doubling the IAM concentration (Table 1, Figure 2B, and Figure S5 (SI)). Interestingly, IAM was able to alkylate and dissociate the four Cd\(_{2+}\) from the Cd\(_4\)(Cys\(_{11}\))\(_{\alpha}\)-domain, while the intermediate Cd\(_4\)NEM\(_{9,11}\)MT\(_2\) remained stable. Thus, not only the higher metal-to-protein affinity but also the bigger size of NEM inhibits the access to the thiols buried in the Cd\(_4\)S\(_{11}\) cluster. In agreement with ESI-MS experiments, NEM reached a plateau with ca. 11 Cys modified for Cd-MT\(_2\) (Figures 2A and 3A). UV−vis confirmed that, even after 24 h of incubation with 3600 equiv of NEM, no more than 4 equiv of Cd\(_{2+}\) was detected (Figure 3B). IAM, on the other hand, did not stabilize any intermediate (Figure 3C,D).

In the light of these results, we interpret the data as follows (Scheme 1): NEM empties out the β-domain faster than IAM, forming the Cd\(_4\)MT\(_2\) α-cluster or a Zn\(_7\)MT\(_2\) where the Zn\(_{2+}\) are redistributed within the α- and β-domains (Figure 2B). Increasing to 750 equiv of alkylator NEM had been added and incubated with for 0−1450 min (B and D, respectively). Red and blue stand for Zn-MT\(_2\) and Cd-MT\(_2\), respectively. Gray and cyan stand for 15 and 60 min of alklylation, respectively.

**Scheme 1. Summary of the Structures Obtained as the Reaction Progressed Followed by either Increasing the Equivalents of NEM or IAM or Increasing the Reaction Time**

| Time (min) | Equivalents of NEM or IAM | Reaction Progressed Followed by | Termination of the reaction or Metalstripping
|------------|--------------------------|---------------------------------|---------------------------------|
| 0−1450    | 3600 equiv for UV−vis and 750 or 1400 equiv for MALDI-MS and ESI-MS | Fix: Reaction time | Fix: Reaction time
| 1450      | 0 equiv for MALDI-MS and ESI-MS | Fix: Reaction time | Fix: Reaction time

The structures are based on the combination of the methods explored in this research, through cysteine profiling with MALDI-MS and metal probe monitoring with UV−vis and native ESI-MS. The Zn\(_7\)MT\(_2\) and Cd\(_7\)MT\(_2\) proteins were incubated at different molar equivalents (e.g. 40, 180, 750, and 1400) for 60 min. In the other option, the proteins were incubated at fixed molar equivalents of NEM or IAM (3600 equiv for UV−vis and 750 or 1400 equiv for MALDI-MS and ESI-MS), and the reaction was followed over time (0−1450 min for UV−vis and 0−60 min for MALDI-MS and ESI-MS).
product, indicating lower kinetic lability (indeed, more inert to breaking metal–ligand bonds). Subsequently, IAM dissociated the four α-domain Cd$^{2+}$ and NEM still did not dissociate the remaining α-Cd$_{3}$S$_{1}$ cluster. This suggests an additional factor to Cd$^{2+}$-S bond stability preventing the disruption of Cd$_{4}$NEM$_{2}$MT2. Therefore, the lower IAM reactivity could benefit the mapping of free Cys residues without altering metal-binding sites, even in these proteins with low metal-to-protein affinity. The small size and hydrophilic character of IAM allows Cys-labeling, even for the Cys residues that are buried.

**Mapping the Zn$_{4}$MT2 Species—Pinpointing the Zn$^{2+}$-Binding Sites.** The previous results helped us with the main aim of this research: the improvement of a differential alkylation strategy to map Zn$^{2+}$-binding sites in partially Zn$^{2+}$-loaded proteins.1 We demonstrated how the use of first IAM followed by NEM to label metal-free Cys residues in the presence and absence of Zn$^{2+}$ is the best choice (Figure 4A).

For this purpose, native ESI-MS and MALDI-MS analysis for metal-free IAM-labeled samples showed modifications ranging from 17 to 20, where the latter was the maximum (Figure 4A,B). Parallel nESI-MS analysis annotated peaks corresponding to [Zn$_{4}$IAM$_{18-20}$MT2]$^{4+}$, with the fully modified species being the most abundant [Figure 4A and Table S1 (SI)]. After removing the excess IAM, NEM-labeled apoMT2 analyzed by MALDI-MS showed a peak previously fully modified by IAM (7183.3 m/z) but also double-labeled peaks with both IAM and NEM (Figure 4C). The distribution of modifications was centered at 7319 m/z, which corresponds to the IAM$_{4}$NEM$_{2}$MT2 species with a visible 7308 m/z peak that comes from overalkylation of IAM$_{2}$NEM$_{4}$MT2. With the conditions applied, we only observed one non-cysteine residue reacting with NEM. Double-labeled peaks extended up to forming IAM$_{2}$NEM$_{2}$MT2 species (7388 m/z).

Addition of 4 equiv of Zn$^{2+}$ to apoMT2 resulted in a wide IAM modification profile ranging from 0 to 14 M with a maximum centered at 8–9 M, which suggests eight or nine noncoordinating Cys residues possibly from the β-domain (Figure 4B). Comparing both the maximum for apoMT2 (20 M) and the results after addition of 4 equiv of Zn$^{2+}$ (8–9 M) reveals a difference of ~11–12 modifications (Figure 4B). This might correspond with the number of Cys residues in the α-domain. The nESI-MS analysis confirmed the presence of [Zn$_{4}$IAM$_{4-n}$MT2]$^{4+}$ ions, but also other Zn$^{2+}$-to-protein stoichiometries are visible [Figure 4A and Table S2 (SI)]. We hypothesized the existence of several conformers, where 4 Zn$^{2+}$ were located in the α-domain or there are redistributed between both β- and α-domains.

To assess the Cys residues coordinating Zn$^{2+}$, IAM-labeled Zn$_{4}$MT2 protein samples were DTT-treated with a reduced pH accompanied by Zn$^{2+}$ removal and further purification using C18 resin (Figure 1). As concluded before, NEM might not be able to displace all of the metal ions in proteins that binds Zn$^{2+}$ with high affinity. To elaborate a general method for proteins that binds Zn$^{2+}$ with low but also with high affinity, we introduced an acidification step for metal ion removal. Moreover, the acidification and metal ion removal promote destabilization of the natively folded structure, facilitating the access of NEM to all of the Cys residues.24 NEM is a suitable reagent for this purpose, since it exhibits fast kinetics even at low pH.16,25 The Zn$^{2+}$ purified protein was then incubated with an excess of NEM. The former population of singly IAM-labeled peaks now appeared double labeled with both IAM and NEM moieties (Figure 4C). For instance, the peak centered at 9 IAM now incorporated 11 NEM molecules in the metal-coordinating Cys residues, forming the IAM$_{4}$NEM$_{11}$MT2 species (7936 m/z), which suggests that before to Zn$^{2+}$ removal, the α-domain was fully occupied with 4 Zn$^{2+}$. As in the previous step, where the protein was singly labeled with IAM, multiple products are observed.

Inferring the localization of Zn$^{2+}$ binding among the Cys residues was achieved with a bottom-up approach. In order to demonstrate the approach presented, we worked with purified proteins or complexes, taking advantage of the speed, sensitivity, and mass accuracy in the analysis provided by the MALDI ion source coupled to the TOF detector. However, one needs to consider other analytical platforms in the case of more complex samples. First, we collected peptide mass fingerprints (PMF) from tryptic fragments measured by MALDI-MS [Figure 4B and Figure S8 (SI)]. A detailed analysis showed an emerging fully modified tryptic fragment, NEM$_{5}$[44–51], from the Zn$_{4}$MT sample (Figure S9, SI). This fragment was not previously found in the apoMT2 sample, suggesting that all three Cys residues had bound Zn$^{2+}$. Congruently, IAM$_{4}$[44–51] is no longer visible (Figure S10, SI). Moreover, we observed the appearance of fragment IAM$_{4}$NEM$_{3}$[31–43] sequenced with IAM-labeled Cys41 accompanied by disappearance of IAM$_{4}$[32–43] (Figures S11 and S12, SI). Fragment NEM$_{5}$[52–61] from the C-terminal region was also detected, suggesting that it had bound Zn$^{2+}$. On the other hand, tryptic peptides NEM$_{5}$[21–30] and NEM$_{5}$[26–31], corresponding to the N-terminal β-domain, emerged for Zn$_{4}$MT. De novo sequencing localized NEM-labeled Cys21 for the former tryptic peptide and NEM-labeled Cys26 and Cys29 for the latter one (Figures S13 and S14, SI). This was confirmed by observing how fragment IAM$_{2}$[23–31] with IAM-labeled Cys26 and Cys29 disappeared (Figure S15, SI). Altogether, the evidence suggests no exclusive selectivity for the first four Zn$^{2+}$ toward the α-domain but partial

![Figure 4. Native ESI-MS (A) and MALDI-MS analysis (B) of apoMT2 with 0, 4, and 7 equiv of Zn$^{2+}$ to which IAM had been added to a concentration of 1 mM and incubated in darkness for 15 min at 25 °C. After removal of excess IAM and metal ions, NEM was added to a concentration of 3 mM and incubated for 30 min at 25 °C (C). ESI-MS is shown as a deconvolved zero-charge mass spectrum.](https://dx.doi.org/10.1021/acs.analchem.0c01604)
redistribution within the β-domain region. This is opposite to its Cd2+ counterpart that demonstrated formation of a stable Cd4MT intermediate.28 In our previous research, we demonstrated an initial sequential Zn2+ binding mechanism for the α-domain forming independent ZnSx sites and Zn3S9 clusters.31 Here, by using the presented methodology, our results support the hypothesis of multiple conformations being simultaneously present, where both populations have the full α-domain saturated, forming Zn2+ clusters (α-Zn3MT), and with redistributed Zn2+ among the α- and β-domains (Zn,αZnβ/MT) coexisting in solution. Thus, it might be feasible to consider the equilibrium αZn3MT = Zn,αZnβ/MT, which indicates a similar Zn2+ stability between the strongest α-site and the weakest α-site. In a previous pH titration report, the authors reported a similar conclusion, which supports this idea.72,73 A second, alternative option is that there are multiple populations, including Zn,M2 and Zn,MT2, coexisting simultaneously. This solution is further supported by native ESI-MS, where we observed multiple metal-loaded species coexisting. The third option includes both solutions, Zn,MT with multiple conformations, where Zn2+ ions are redistributed between both domains, and coexisting Zn,M,MT. Recently, a paper reported collision-induced unfolding experiments followed through ion mobility-MS for Zn,MT2.74 The authors reported four separate conformers with regard to collision cross section (CCS) profiles that required a different collision activation for unfolding. The β-domain is not fully extended but rather folded for Cd4MT2, as opposed to Zn4MT2, which showed a CCS of around 1000, which corresponds to the β-domain being fully extended. This may support the idea of an equilibrium, αZn3MT = Zn,αZnβ/MT, that is shifted toward the formation of αZn3MT upon protein destabilization. In agreement, Stillman’s group reported that at physiological pH the four Zn2+ are bound between both domains.75 The authors suggested that the terminal Cys5 and Cys7 are involved in Zn2+ coordination. However, this idea has been supported by neither Russell’s group nor here.66,75 Russell’s group concluded that the region from Asn18 to Cys38 participates in the coordination of the first four Zn2+ ions, but the terminal Cys5 or Cys7 are at most weakly interacting. Here, with the experimental conditions employed, we suggest that the region 21–30 participates in Zn2+ binding. Despite the above differences, our results agreed and suggested that there exists a Zn2+ redistribution between both α- and β-domains.74,75,83

Subsequently, addition of 5 equiv of Zn2+ to apoMT2 shifted the distribution of IAM modifications, now centered at 6–7 with a maximum of 10 IAM, clearly right skewed, clumping up populations with few or no IAM modifications (Figure S16A, SI). The complementary, double-labeling distribution obtained was centered, forming the IAM,NEM5,MT2 species (8067.5 m/z) (Figure S16C, SI). Interestingly, this stoichiometry does not resemble that for coordination of a single ZnCys4 site binding or a completely saturated α-cluster (Zn3S11) but rather suggests a structure with redistributed Zn2+ ions between domains.84,85 Moreover, a mixture of gas-phase ions with [Zn7IAM0MT2]5+ stoichiometry was annotated by nESI, which confirms the products IAM,NEM5,MT2 (7817 m/z) and IAM,NEM5,MT2 (8339 m/z) found in MALDI-MS (Figure S16B and Table S3, SI). The former one correlated with Zn3S9 stoichiometry and the latter with Zn3S17; [Zn4(Cys)11 α-domain and Zn3(Cys)2 in β-domain].

Fragments analyzed from PMF traced increased tryptic NEM,26–31 with respect to Zn3MT, which after thorough MS/MS sequenced the NEM molecule attached to Cys29 (Figure 1B and Figure S17 (SI)). The region 1–20 appeared with two IAM modifications for Cys5 and Cys19, so the three remaining Cys residues might be involved in coordination of the fifth Zn2+ (Figure S18, SI). Additionally, tryptic NEM5,44–61, NEM5,31–43, and NEM5,31–43 were detected and sequenced (Figures S19 and S20, SI). These results provide interesting clues concerning the binding of the fifth Zn2+. As the protein is being Zn2+ saturated, the α-domain folds to an α-Zn3S11 cluster and starts to fill the β-domain. The question arises whether a Zn2+ redistribution between domains occurs, to end up with a coordinated native-like α-cluster and a single Zn3S9 site in the β-domain. It is noteworthy how the Zn3S9 coordination sphere pinpointed for the single β-site does not resemble any position from the rabbit X-ray structure but perfectly matches our previous report.51 In that study, we indicated that the fifth Zn2+ probably causes conformational changes for the 1–20 and 26–31 sequence regions (Figure 1B). We might confirm that, under the conditions employed, Cys7, Cys13, Cys15, and Cys29 coordinate the fifth Zn2+ (Figure 1B). Notwithstanding, other Zn,MT2 populations are simultaneously present, as observed by IM-MS experiments, but likely with lower probability than the one described.59

A total of 6 equiv of Zn2+ added to apoMT2 caused an abrupt change of the IAM modification profile, with none to four modifications giving 1 IAM as the most intense peak (Figure S16D). Gas-phase ions detected by the nESI-MS spectrum showed the product ions [Zn7IAM0MT2]5+, [Zn5IAM2MT2]5+, and [Zn7IAM,MT2]5+ (Figure S16E and Table S4, SI). Likewise, double-labeled proteins showed that various species formed, ranging from IAM,NEM5,MT to IAM,NEM5,MT, the latter corresponding to Zn,MT2 (Figure S16F, SI). Precisely, we found a tryptic fragment NEM5,23–31 with NEM-labeled Cys24 and Cys29 (Figure S21). These two Cys residues were not previously found to be coordinating and their participation in Zn2+ coordination appears after addition of the 6 equiv of Zn2+. Such results match perfectly with our previous research, validating the approach presented herein.51,85 Addition of the last Zn5+ equivalent did not cause an observable major change for the MALDI modification profile, with the most intensive modification corresponding to one Cys residue (Figure 4B). The nESI-MS results distinguished gas-phase ions [Zn5IAM2MT2]5+ and [Zn4IAM2MT2]5+ (Figure 4A and Table S5 (SI)). So, after removal of Zn5+, NEM labeled those Cys residues, matching with the stoichiometry obtained from MALDI-MS (Figure 4C). Interestingly, the bottom-up MS results identified the tryptic fragment NEM5,21–30 with NEM-labeled Cys21 as still existing in the spectra, confirming the gating role of Cys21 for the seventh Zn2+ previously reported (Figure 1B and Figure S22 (SI)).51

■ CONCLUSION

Cysteine is the most nucleophilic amino acid residue of all building proteins; a target for reactive oxygen, nitrogen, and sulfur species and numerous chemical reactions, including post-translational ones; and critical for binding essential and toxic metal ions. To identify the chemical or redox state of Cys residues in proteins, analytical methods are based on reactions with thiol-specific probes. Among them, IAM and NEM are commonly used following an S2+ reaction and nucleophilic addition, respectively. Differential labeling coupled to mass spectrometry is used for mapping metal-binding sites. In
principle, a single labeling approach should be sufficient to pinpoint Zn\textsubscript{2}(Cys\textsubscript{3}) protein sites, on which the Zn\textsuperscript{2+}-bound thiolate would exhibit lower reactivity toward electrophiles. However, this is not the case for low- or moderate-affinity Zn\textsuperscript{2+}-binding sites, where unwanted metal ion dissociation during protein modification might occur. Moreover, a similar reactivity for a Zn\textsuperscript{2+}-bound and free Cys residue with low pK\textsubscript{a} might occur, hampering their nuclophilic differentiation. Herein, we developed a dual-labeling methodology to overcome these obstacles. Our findings suggested proceeding with an S\textsubscript{N}2 reaction with IAM or a similar reagent, but not with a nucleophilic addition provided by maleimides, to label free Cys residues and avoid Zn\textsuperscript{2+} dissociation. The small size and hydrophilic character permits the labeling of free Cys residues, regardless if they are buried or not. Once the free Cys residues are labeled, metal-bound Cys might easily be labeled by a second nucleophile after extraction of the metal ion by acidification and followed by reverse-phase C18 separation. A priori, Zn\textsuperscript{2+} removal can be approached by increasing the alkylator concentration or reaction time. However, this approach does not work for high-stability metal–protein sites, and substantial overallkation may occur. Thus, the acidification step ensures metal ion removal independent of how strongly it is bound. Not only the metal-to-protein affinity but also the thiol accessibility may limit the labeling reaction. The acidification step greatly facilitates the access of the second alkylator to all of the Cys residues, even if they were buried in the protein structure. Although any reagent different than IAM would in principle work for the second labeling reaction, NEM reacts faster than iodoacetamide derivatives with thiols/thiolates, even at low pH.

This research demonstrated how the combination of inexpensive standard labeling reagents and mass spectrometry can directly map metal-binding sites in Cys-rich proteins. Lastly, the methodology presented is very encouraging and illustrates its future use to study other metal-binding Cys proteins.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.analchem.0c01604.

Detailed materials and protocol for expression and purification of MTs; MALDI-MS spectra of Cd-MT2 and Zn-MT2, to which 15–3600 equiv of NEM or IAM had been added, after 60 min of incubation; UV–vis spectra of the metal displacement reactions after 60 min incubation with 0–3600 equiv of IAM and NEM with Cd-MT2 and Zn-MT2; ESI-MS spectra of Cd-MT2 and Zn-MT2 incubated with increasing equivalents of IAM and NEM for 60 min; ESI-MS spectra after 60 min of incubation with 750 or 1400 equiv of IAM and NEM with Cd-MT2 and Zn-MT2; number of Cys modified determined by MALDI-MS as a function of alkylator/MT ratio after 60 min incubation; peptide mass fingerprint for IAM and NEM labeling for Zn-MT2, Zn\textsubscript{2}MT2, Zn\textsubscript{3}MT2, and Zn\textsubscript{4}MT2 samples analyzed by MALDI-MS; MALDI-MS/MS analysis for the tryptic fragment of Zn\textsubscript{2}MT2, Zn\textsubscript{3}MT2, Zn\textsubscript{4}MT2, Zn\textsubscript{5}MT2, and Zn\textsubscript{6}MT2 samples; gas-phase ions annotated for Zn\textsubscript{3}MT2, Zn\textsubscript{4}MT2, Zn\textsubscript{5}MT2, and Zn\textsubscript{6}MT2 samples labeled with IAM (Figures S1–S22 and Tables S1–S5) (PDF)

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Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

This research was supported by the National Science Centre of Poland (NCN) under the Opus grant no. 2018/31/B/NZI/00567 (to A.K.). M.D.P.-D. thanks the Erasmus+ program and the Polish National Agency for Academic Exchange under the PROM program (grant no. PPI/PRO/2018/1/00007/U/00). V.A. would like to thank to the European Research Council (ERC) for financial support from the European Union’s Horizon 2020 Research and Innovation Programme (grant agreement no. 759585).

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