Impact of naturally occurring serine/cysteine variations on the structure and function of Pseudomonas metallothioneins†

Jelena Habjanič, Serge Chesnov, Oliver Zerbe and Eva Freisinger*

Metallothioneins (MTs), small cysteine-rich metal-binding proteins, support the viability of organisms under normal physiological conditions and help them to respond to different environmental stressors. Upon metal coordination (e.g. Zn\(^{II}\), Cd\(^{II}\), Cu\(^{I}\)) they form characteristic polynuclear metal–thiolate clusters that are known for their high thermodynamic stability and kinetic lability. However, despite numerous studies, it is still not understood how MTs modulate their metal-binding properties. Pseudomonas MTs are an emerging subclass of bacterial MTs, distinct for their high number of His residues and for several unique features such as an intrinsically disordered long C-terminal tail and multiple variations in the number and nature of coordinating amino acids. These variations might provide the bacteria with a functional advantage derived from evolutionary adaptation to heterogeneous environments. Nearly 90% of the known Pseudomonas MT sequences feature a central YC\(\times\)xC motif, that is altered to YC\(\times\)xC in the rest. We demonstrate that the additional Cys residue serves as a coordinating ligand without influencing the metal-binding capacity, the overall metal-binding stability or the structure. However, the additional ligand changes intra-cluster dynamics and, as a consequence, modulates metal transfer reactions that could be functionally advantageous in vivo.

Significance to metallomics

The metal clusters of metallothioneins (MTs) are known for their unique properties such as a high thermodynamic stability that is nevertheless combined with a kinetic lability. Both are crucial for their biological functions, among others, in metal ion homeostasis and detoxification. However, insights into the modulation of metal-binding properties by amino acid sequence variations is scarce. In this work we demonstrate how altering the number of potentially coordinating ligands in form of naturally occurring Ser/Cys variations help modulating metal transfer reactions in Pseudomonas MTs. These variations do not influence the overall protein structure but significantly change intra-cluster dynamics.

Introduction

Metallothioneins (MTs) are unique, low-molecular-weight (6–20 kDa) proteins, characterised by an extraordinary high cysteine content (15–30%). The Cys residues are organized in specific Cys motifs (e.g. CxC, C\(\times\)C, CC, CC\(\times\)C, C\(\times\)CC) that allow the formation of diverse metal–thiolate clusters. Depending on the biological nature of the bound transition metal ion, MTs are involved in a wide variety of vital physiological processes, which include metal homeostasis and detoxification, as well as a defence against nitrative and oxidative stress owing to the redox properties of the Cys residues.\(^{1–3}\) The main differences between bacterial and vertebrate MTs are the involvement of His residues in metal ion coordination, the greater sequence diversity now also featuring aromatic residues, and a higher percentage of secondary structural elements.\(^{4,5}\)

Pseudomonas metallothionein (PsdMT) sequences were identified only recently\(^{5,6}\) and the encoded proteins form a highly conserved subfamily of bacterial metallothioneins (bacMTs). PsdMTs share a conserved Cys distribution pattern with cyanobacterial MTs, the most intensively investigated bacMTs, but the crucial difference is the replacement of one of the two coordinating His ligands in cyanobacterial MTs by a non-coordinating residue (e.g. Ala, Gly, Ser) in PsdMTs. Furthermore, the His content in PsdMTs is generally unusually high, some sequences contain a long C-terminal Cys-free amino acid stretch with a highly conserved sequence,\(^{7}\) and in almost 90% of the sequences identified so far an additional Cys instead of a...
Ser residue can be found in one of the central motifs, i.e. YCxxC instead of YCxxC. This additional Cys residue is also present in 30% of cyanobacterial MTs (Fig. 1).

In our previous work, we have structurally and functionally characterized some of these novel features using an MT from \textit{Pseudomonas fluorescens} Q2-87 (PflQ2 MT, sequence 8 in Fig. 1).\textsuperscript{7} Our studies revealed that the long C-terminal tail is intrinsically disordered and neither influencing the metal-binding properties nor the structure of the rest of the protein. However, it might potentially serve as a biologically relevant binding site for globular proteins or other interacting molecules. Surprisingly, PsdMTs display differences in their maximum metal-binding capacity towards Zn\textsuperscript{II} and Cd\textsuperscript{II}, metal ions that are often considered isostructurally replaceable, as reflected in the formation of a Cd\textsubscript{4} versus a Zn\textsubscript{3} cluster. The maximum binding capacity for Zn\textsuperscript{II} was restored to four by introducing a second coordinating His residue as found in cyanobacterial MTs but is accompanied by a decrease in kinetic lability. The investigation of the additional non-coordinating His residues present in the sequence revealed no significant impact on stability, structure or metal-binding properties of the protein, and the biological relevance remains unclear.

In the present study, we address another major feature of PsdMTs, that is a naturally occurring Ser/Cys variation in the central stretch of the Cys distribution pattern (YCS/CxxC). The change of Ser to Cys implies altering only a single atom, that is oxygen to sulphur. However, for MTs, this change potentially introduces an additional metal binding ligand and therefore might significantly affect the metal-binding properties, and hence function and structure. Upon metal binding, the Cys thiol group is deprotonated, reducing its apparent pK\textsubscript{a} value from approx. 8.5 in the free amino acid\textsuperscript{6,9} by 4–5 units, depending on the metal ion. In contrast, the higher electronegativity of the oxygen ($\chi = 3.44$) compared to the sulphur ($\chi = 2.58$)$\textsuperscript{10}$ prevents deprotonation of the hydroxymethyl group of Ser under physiological conditions. In fact, Ser residues have never been observed to coordinate to metal ions in MTs, despite their high abundance. For example, in vertebrate MTs they are the second most abundant residue after Cys.$\textsuperscript{11,12}$ Strikingly, Ser residues are often located in the direct vicinity of Cys residues, and mutagenesis studies showed that conserved Ser residues contribute to the strength and overall stability of metal binding.$\textsuperscript{12}$

The previously studied PflQ2 MT has a Ser residue at the specified position (Ser29, YCxxC). The S29C mutant of the protein will be used here to evaluate if the additional Cys residue is participating in metal ion coordination and whether it changes metal-binding properties, function and structure in comparison to the wild-type protein. In addition, an MT with a naturally occurring Cys residue at this position (Cys28), \textit{Pseudomonas putida} KT2440 (PpKT MT, sequence 7 in Fig. 1), is studied together with its C28S mutant.

**Materials and methods**

**Chemicals and solutions**

The enzymes and purification systems used for plasmid construction were purchased from Fermentas (Le Mont-sur-Lausanne, Switzerland) and Sigma-Aldrich (Buchs, Switzerland). All buffers and chemicals were ACS grade or comparable from Sigma-Aldrich (Buchs, Switzerland), Merck (Darmstadt, Germany), Chemie Brunschwig (Basel, Switzerland), and Roth AG (Arlesheim, Switzerland). Chelex\textsuperscript{8} 100 resin was from Bio-Rad (Reinach, Switzerland). Isotopically labelled compounds (tris(hydroxymethyl)aminomethane, Tris, (D\textsubscript{11}, 98%), D-glucose (U-13C\textsubscript{6}, 99%), ammonium chloride ($^{15}$N, 99%), and cadmium chloride ($^{111}$Cd, 95.24%)) were purchased from Cambridge Isotope Laboratories (ReseChem, Burgdorf, Switzerland), while D\textsubscript{2}O (99.8%) was purchased from Addgene (Addgene plasmid #8827). All solutions were prepared using Millipore water and were degassed under vacuum or saturated with argon. To exclude oxygen, experiments were performed in a Type B Vinyl Anaerobic Chamber (Coy Lab, USA) equipped with a palladium catalyst and operated with a 5% hydrogen/95% nitrogen gas mix wherever possible.
Protein expression and purification

The coding sequence of the PpKT MT (residues 1–73) was optimised for the expression in E. coli (Fig. S1, ESI†) and constructed from commercially obtained oligonucleotides (Microsynth AG, Balgach, Switzerland) (Table S1, ESI†). A tobacco etch virus (TEV) protease cleavage site was introduced at the 5′end of the protein-encoding sequence. Using the upstream BamHI and the downstream Xmal restriction sites, the coding sequence was cloned into a pGEX-4T-1 expression vector, which encodes an N-terminal glutathione S-transferase (GST) tag (Fig. S2, ESI†). The shortened version of the protein lacking the C-terminal tail, sh_PpKT MT (residues 1–51) was constructed in the same manner. Both newly constructed plasmids, as well as previously constructed plasmids of the full-length PFLQ2 MT and its shortened version sh_PFLQ2 MT,† were used as a DNA template for the site-directed mutagenesis to construct the C28S and S29C mutants, respectively. Oligonucleotides used for the site-directed mutagenesis are listed in Table S1 (ESI†).

All proteins were expressed and purified as previously described.† The identity of proteins was confirmed by electrospray ionization mass spectrometry (ESI-MS) (Fig. S3, ESI†).

Determination of metal ion binding capacity

To determine the maximum metal-binding capacity, an excess of the respective metal ion was added to the apo-protein and the pH increased to 7.4. To remove the metal ion excess, the samples were either purified by size exclusion chromatography (SEC; Superdex Peptide 10/300 GL column; GE Healthcare, Glattbrugg, Switzerland) or by incubation with Chelex resin, both pre-equilibrated with 50 mM Tris HCl, 50 mM NaCl, pH 7.4. Samples of the holo-protein forms obtained in this way were analysed by flame atomic absorption spectroscopy (F-AAS) on an AA240FS spectrometer (Agilent Technologies AG, Basel, Switzerland) to determine the metal ion concentration and by a capillary voltage of 2.8 V, a cone voltage of 40 V, and a source temperature of 80 °C. The apo-forms were measured in MeOH:0.15% aqueous formic acid (1:1) and the holo-forms in MeOH:30 mM aqueous ammonium acetate (1:1). The m/z data were deconvoluted using the MaxEnt1 software with a resolution of the output mass of 0.1 Da per channel and a Uniform Gaussian Damage Model at the half-height of 0.1 Da.

NMR studies

To prepare the different metallated species, an exact amount of cadmium ions in the form of a CdCl₂ solution was added dropwise to the freshly prepared apo-protein solution and the pH was increased to 7.4 with 1 M Tris-D₁₁ and/or small amounts of NaOH. Samples were lyophilized and resuspended in a buffer containing 50 mM Tris-D₁₁ HCl, 50 mM NaCl, pH 7.4 (90% H₂O/10% D₂O) just prior to the measurements. The average concentration of ¹⁵N (¹³C,¹⁵N) labelled samples and those used for ¹¹³Cd measurements was around 0.5 mM and 1–2 mM, respectively. Referencing of ¹H chemical shifts was done with the external standard 4,4-dimethyl-4-silapentane-1-sulfonic acid (DSS, 0.2% pH 7.5), while ¹⁵N and ¹³C frequencies were referenced indirectly. The ¹¹³Cd chemical shifts were referenced to an external 0.1 M Cd(ClO₄)₂ solution. Experiments were recorded on a Bruker 600 MHz spectrometer equipped with a CRYO TCI triple-resonance probe (¹H, ¹³C, ¹⁵N) if not stated differently. The ¹D ¹¹³Cd and [¹¹³Cd,¹H]-HSQC-TOCSY spectra were recorded on a Bruker 500 MHz spectrometer equipped with a CRYO 5 mm QNP probe in the temperature range from 275 to 320 K. Assignments and structure calculations were performed in the same way as described previously.†

¹⁵N relaxation measurements

The backbone amide dynamics were studied by measuring ¹⁵N longitudinal and transverse relaxation rates (Rₓ, Rₓ) as well as ¹⁵N(¹H)-NOE experiments. Delays were set to 0.01, 0.015, 0.025, 0.05, 0.1, 0.15, 0.2, 0.5, 1.0, 1.5, 2.0, and 2.5 s for the longitudinal (T₁) relaxation measurement and to 0, 5, 15, 25, 50, 75, 100, 125, 150, 250, 300, 400, 450, and 500 ms for the transverse (T₂) relaxation measurement. The peaks were integrated with SP Cornell, and the data were analysed using in-house written...
scripts. In order to extract the relaxation rate, the signal intensities were least square fitted to an exponential decay:

\[ I = I_0 e^{-\frac{t}{T_1}} \]

in which \( T_1 \) corresponds to \( T_1' \) or \( T_2' \) and \( t \) to the length of the corresponding delay for measuring that data point. \(^{15}\)N\(\{^1\)H\}\)-NOE experiments were recorded with a steady-state \(^{15}\)N\(\{^1\)H\}\)-NOE version of the \(^{15}\)N-HSQC experiment using an experiment repetition delay of 5 s. Two \(^{15}\)N\(\{^1\)H\}\)-HSQC spectra, one with and one without amide proton saturation were recorded and the \(^{15}\)N\(\{^1\)H\}\)-NOE values were computed from the ratio of peak volumes in the two experiments.

**Thermal denaturation**

NMR samples of \(^{13}\)C,\(^{15}\)N labelled Cd\(_4\)sh\_PflQ2 MT and its S29C mutant were lyophilized and resuspended in 100% D\(_2\)O. \(^{15}\)N\(\{^1\)H\}\)-HSQC spectra were measured from 290 K to 323 K in 3 K intervals. Spectra were calibrated against an internal standard (tetramethylsilane), analysed using CARA,16 and peak positions adjusted manually at each temperature. \(^1\)H and \(^{13}\)C chemical shifts differences against the standard at 290 K were computed for the different temperatures and tabulated in the chemical shifts differences against the standard at 290 K were computed for the different temperatures and tabulated in the form of their absolute values. The absolute values of the chemical shift differences were computed by subtracting corresponding chemical shifts:

\[ \Delta \delta_i = \sqrt{(\delta_i^{WT} - \delta_i^{MUT})^2} \]

**Cd\(_{11}\)** transfer reactions to EDTA

The Cd\(_4\) species of sh\_PflQ2 MT and sh\_PpKT MT as well as of their respective mutants, S29C-sh\_PflQ2 MT and C28S-sh\_PpKT MT, were prepared by addition of exactly four equivalents of cadmium ions to the respective apo-forms. Subsequently, the pH was increased to 7.4 with small amounts of aqueous ammonia solutions (5–20%). For each transfer reaction, 100 \( \mu \)L of a solution containing 10 \( \mu \)M of the respective Cd\(_4\) species and 40 \( \mu \)M ethylenediaminetetraacetic acid (EDTA) were prepared and incubated at room temperature. The stability constant for the [Cd\(_{11}\)(EDTA)]\(^{2-}\) complex is \( 10^{16.5} M^{-1} \) (25 °C, 0.1 M ionic strength).17

At different time intervals, 3 \( \mu \)L aliquots of each solution were desalted using C\(_{18}\) ZipTips (Merck Millipore, Darmstadt, Germany). The samples were eluted with 10 mM ammonium acetate in 30% MeOH and analysed with the Synapt G2 quadrupole time-of-flight spectrometer operating in the positive ion mode with settings as described above. Data were acquired for 2 min in the \( m/z \) range between 50 and 5000 Da. Relative quantification of the detected species was performed based on the sum of signal intensities of +3 and +4 ions.

**Results and discussion**

Similar to PflQ2 MT, the PpKT MT contains a 32-residue long C-terminal tail. For PflQ2 MT, it was shown that this tail is intrinsically disordered and does neither influence the metal-binding properties nor the structure of the rest of the protein.7 The latter was also confirmed for PpKT MT by comparing the \(^{15}\)N\(\{^1\)H\}\)-HSQC spectra of the full-length protein and its shortened version (residues 1–51) (data not shown). Hence, we will focus in this study only on the data obtained with the truncated constructs of the proteins to simplify assignments. Furthermore, while PsdMTs can bind both zinc and cadmium ions, we limit our investigations here to Cd\(_{11}\) because Zn\(_{11}\) ions are spectroscopically silent in most of the experiments presented.

**Impact of the Ser/Cys variation on the metal-binding properties**

To assess the effect of the Ser/Cys variation, first, its influence on the metal ion binding capacity was analysed. Based on F-AAS and the 2-PDS assay all four constructs can bind up to four Cd\(_{11}\) ions (PflQ2 MT (3.65 ± 0.16), S29C-PflQ2 MT (3.79 ± 0.14), PpKT MT (3.62 ± 0.20), C28S-PpKT MT (3.78 ± 0.16)). This was confirmed by ESI-MS (Fig. S4, ESI†) showing a major signal for the Cd\(_4\) species and a minor for the Cd\(_3\) species, the latter being an indication for a certain lability of the 4th binding site for all four constructs. The overall apparent metal binding stabilities of the Cd\(_3\) and Cd\(_4\) species were analysed using \(^{19}\)F NMR spectroscopy and the metal chelator 5,5'-difluoro-BAPTA and show only minor differences between the four constructs, the most apparent being a higher stability of Cd\(_3\)PpKT MT by...
0.75 log units compared to its C28S mutant (Fig. S5, ESI†). Hence, while obviously the Ser/Cys variation does not influence the Cd\textsuperscript{II} binding capacity and has a very minor influence on the thermodynamic stability, this does not exclude the possibility that the additional 10th Cys residue might be involved in Cd\textsuperscript{II} binding. To answer this question, the four different apo-proteins were titrated with Cd\textsuperscript{II} ions, and the absorptivity increase of the thiolate-to-cadmium charge transfer band at 250 nm was monitored by UV spectroscopy (Fig. 2). As a result, the maximum molar absorptivity between wild-type and mutant protein differs by approx. 7000 M\textsuperscript{-1} cm\textsuperscript{-1} for both MTs, that is 42 700 M\textsuperscript{-1} cm\textsuperscript{-1} versus 35 900 M\textsuperscript{-1} cm\textsuperscript{-1} for PpKT MT and its C28S mutant, and 45 000 M\textsuperscript{-1} cm\textsuperscript{-1} versus 38 200 M\textsuperscript{-1} cm\textsuperscript{-1} for PflQ2 MT and its S29C mutant. Hence the 10th Cys residue clearly increases the maximum molar absorptivity, confirming its involvement in Cd\textsuperscript{II} binding.

Impact on the structure, backbone dynamics and stability of the protein fold

In the structure of Cd\textsubscript{4}PflQ2 MT, Ser29 has an essential role as an N-cap for a short α-helix, forming two hydrogen bonds, one from its carbonyl oxygen (O) to the amide proton (H\textsuperscript{N}) of Ala33 and a second one from the side-chain oxygen (O\textprime{}) to the amide proton (H\textsuperscript{N}\textprime{}) of Ala31. Since the Cd\textsuperscript{II} binding studies showed that upon mutation of Ser29, the additional 10th Cys serves as an additional ligand, the observation of some conformational changes seems likely. Indeed, analysis of the C\textsuperscript{α} and C\textsuperscript{β} chemical shift differences of both Cd\textsubscript{4} and Cd\textsubscript{2} species of PflQ2 MT and its S29C mutant (Fig. S6 and S7, ESI†) reveals that significant changes only occur in the vicinity to the mutation. Indeed, the NMR solution structure of Cd\textsubscript{4}S29C-PflQ2 MT is highly similar to the wild-type (Fig. 3) and hence, any significant impact of the Ser-to-Cys mutation on the protein structure can be excluded.

The comparison of backbone amide dynamics between PfiQ2 MT and its S29C mutant reveals a moderate perturbation in \textsuperscript{15}N[\textsuperscript{1}H]-NOE values and longitudinal relaxation rates (R\textsubscript{1}). The most prominent differences are observed for Cys residues (Fig. S8 and S9, ESI†). More pronounced difference between wild-type and mutant protein are observed for the transverse relaxation rates (R\textsubscript{2}) (Fig. S10, ESI†). The considerably higher R\textsubscript{2} rates for the S29C mutant, again mostly for Cys residues, suggest the presence of exchange processes driven by Cys residues. Therefore, although the two protein folds are highly similar, the Ser-to-Cys mutation triggers a certain degree of change in the dynamics.

The impact on the protein fold stability was investigated with NMR melting studies, monitoring C\textsuperscript{α} and C\textsuperscript{β} chemical shift changes for the Cd\textsubscript{4} species of PfiQ2 MT and its S29C mutant upon temperature increase from 290 K to 323 K. The results demonstrate an overall destabilisation of the S29C mutant compared to the wild-type (Fig. S11, ESI†). Surprisingly, the most significant differences are not in the direct vicinity of the mutation but predominantly localized in the amino acid stretch from Pro\textsubscript{8} to Tyr27 with several other significant changes in polar or charged residues distributed along the entire protein sequence. In contrast to most other proteins, whose structures are to a large extend stabilized by hydrophobic interactions, MTs feature a rather high percentage of polar and charged amino acids that contribute to protein structure and stability via electrostatic interactions. Hence the observed changes in the S29C mutant are well in line with observed reduced protein stability. In order to further investigate if protein sequences are naturally optimised to adapt to either Ser or Cys at this specific position, we also investigated the PpKT MT, that contains a 10th Cys residue and the corresponding C28S mutant by NMR. The \textsuperscript{15}N[\textsuperscript{1}H]-HSQC spectrum of Cd\textsubscript{4}PpKT MT shows pronounced line broadening (Fig. 4A), which is significantly smaller in Cd\textsubscript{4}C28S-PpKT MT (Fig. 4B) and even more in Cd\textsubscript{4}C28S-PpKT MT (Fig. 4C) as both spectra display sharper peaks. It seems thus that the additional Cys residue introduces exchange processes, possibly indicating that interconversion between different metal-coordination modes takes place, which is more evident when the fourth Cd\textsuperscript{II} ion is bound.

This line broadening prohibits proper determination of the PpKT MT structure. However, assignment of backbone chemical shifts for Cd\textsubscript{4}PpKT MT reveals the characteristic “zinc-finger fold” pattern seen in PfiQ2 MT\textsuperscript{2} and SmtA\textsuperscript{18} (low-field shift of Cys27 H\textsuperscript{N}) (9.702 ppm due to hydrogen bonding of Cys5 S\textprime{}); high-field shift of Ala32 H\textsuperscript{α} (1.886 ppm) due to interaction with the aromatic ring of Tyr26), indicating a highly similar protein fold.

Impact on the metal ion coordination environment and the cluster structure

Considering the contribution of the additional Cys residue in PpKT and S29C-PfiQ2 MT to Cd\textsuperscript{II} binding, its influence on the
metal ion coordination environment and cluster structure was further investigated. The 1D $^{113}\text{Cd}$ spectrum of Cd$_4$PflQ2 MT displays four peaks representing four different Cd$^{II}$ coordination sites (Fig. 5A). Three of the peaks are sharp, while the fourth peak at 535 ppm with H36, C7, C32, C49 as coordinating residues is significantly broader and only visible at higher temperature. This broadening is attributed to an exchange process on the intermediate time scale for this site compared to the other three sites, which share Cys10 as a common residue. Interestingly, the integral of that resonance is also significantly reduced, beyond the error of integration. In studies of human MT-3 (also referred to as “human neuronal growth inhibitory factor” (GIF)), a similar observations was made in that some of the $^{113}\text{Cd}$ resonances were very broad at ambient temperature and became more sharp at higher temperature, while their integral was still much smaller than the integral of the other signals. This was interpreted as a coupled slow-configurational exchange process, due to Pro-Xaa cis/trans isomerization, with a fast conformational exchange that likely takes place in those Pro-Xaa isomer subpopulations that cannot coordinate to the metal ions. The investigated MTs contain a very large number of residues, and this is the case not only for residues spacing
the Cys/His residues that coordinate to the signal at 535 ppm (Cd-D). Moreover, this effect is much less visible in the Cd₃ species, indicating a non-obvious contribution to this effect from the additional Cd²⁺ ion.

Also the 1D $^{113}$Cd spectrum of the S29C mutant shows four distinct peaks, but here the fourth peak (at 549.2 ppm) is sharper and downfield shifted, while chemical shifts changes of the other three peaks are minor (Fig. 5B). Considering the extraordinary high sensitivity of the $^{113}$Cd chemical shift even to small differences in the coordination environment as well as to exchange processes, the coordination mode for the three sites (708.7, 619.3, 585.8 ppm) can be assumed to be virtually identical in wild-type and S29C mutant. In contrast, the more significant difference for the fourth site (549.2 ppm) points to a change in the coordination environment and/or an alteration in the timescale of the exchange process observed in the wild-type. For the Cd₃ species of both PflQ2 MTs and its S29C mutant (Fig. 5C and D), the 1D $^{113}$Cd spectra contain three well-defined peaks corresponding to three distinct cadmium binding sites. As the chemical shifts minimally differ between the two spectra, it is clear that changes in the chemical environment of the Cd²⁺ ions are minor.

In analogy to the $^{15}$N,¹H]-HSQC spectrum (Fig. 4A), the 1D $^{113}$Cd spectrum of Cd₄PpKT MT at 320 K shows various broad, low intensity and ill-defined peaks in the range from 550 ppm to 750 ppm (Fig. 6A). When the temperature is lowered to 280 K many low-intensity peaks, some of which are sharp, occur indicating that the exchange process is now in the slow-exchange NMR regime. In contrast and in analogy to the $^{15}$N,¹H]-HSQC spectrum (Fig. 4B), the 1D $^{113}$Cd spectrum of Cd₄C28S-PpKT MT is better defined and shows three peaks of higher intensity (704.9, 624.2, and 606 ppm), one of which is sharp (Fig. 6B). Apparently, there is a change in the timescale of the exchange processes between PpKT MT and its C28S mutant that is reflected in the number and broadness of peaks in the $^{113}$Cd spectrum. Again, Cd₄PpKT MT is even less dynamic, and its 1D $^{113}$Cd spectrum shows three well-defined peaks (683.6, 673.9, and 602.6 ppm) at a lower temperature (280 K; Fig. 6C), similar to the spectra of Cd₄PflQ2 MT and its S29C mutant. In contrast, in the spectrum of Cd₄C28S-PpKT MT only two peaks (691.8 and 676.5 ppm; Fig. 6D) are observed and only at a higher temperature (320 K). Hence, these results corroborate the view that the Ser/Cys variation influences the exchange processes by altering the timescale of intra-cluster dynamics while retaining the overall chemical environment.

Due to the well-resolved NMR spectra, we were able to further address the impact of the additional Cys ligand on the Cd₄ cluster structure in S29C-PflQ2 MT. As shown above, NMR studies of Cd₄S29C-PflQ2 MT reveal that the Cd₄ cluster has four distinct Cd²⁺ coordination sites reflected by the four defined peaks in the 1D $^{113}$Cd spectrum (Fig. 5B). As in the previously published wild-type structure, the chemical shifts of two peaks (708.7 ppm and 619.3 ppm) are in the range of Cys₄ polynuclear Cd²⁺ coordination while the other two peaks (585.5 ppm and 549.2 ppm) are indicative for either the involvement of one His residues in coordination, a deviation from ideal tetrahedral tetrathiolate geometry, or a transient coordination.⁷,²¹,²²

Like in wild-type Cd₄PflQ2 MT, His36 coordinates to one of the Cd²⁺ ions as detected by the characteristic scalar coupling of $^{113}$Cd to Nᵦ of His36 (Fig. S12, ESI†) in the long-range

**Fig. 6** $^{113}$Cd NMR of PpKT and its C28S mutant. Proton-decoupled 1D $^{113}$Cd spectra of the Cd₄ species of PpKT MT (A) and the C28S mutant (B). Cd₃ species of PpKT MT (C) and its C28S mutant (D). Peak integrals are listed in Table S2 (ESI†).
[\textsuperscript{15}N,\textsuperscript{1}H]-HSQC spectrum. In analogy to the wild-type protein, the \textsuperscript{113}Cd peak at 549.2 ppm was assigned to the Cys\textsubscript{3}His site. This was additionally confirmed during structure calculation since assigning the Cys\textsubscript{3}His site to peak at 585.5 ppm resulted in significant violations in NMR distances. Therefore, we assume that the signal at 585.5 ppm is upfield-shifted due to a deviation from ideal tetrahedral tetrathiolate coordination geometry accompanied by transient coordination of the fourth ligand as was previously discussed for the wild-type.\textsuperscript{7} Since a total of 11 ligands (10 Cys and 1 His) in Cd\textsubscript{4}S\textsubscript{29C-PflQ2} MT are involved in Cd\textsuperscript{II} coordination, a typical \(\alpha\)-type cluster as characteristic for vertebrate and echinoderm MTs\textsuperscript{23,24} and also found in cyanobacterial MTs (where two terminal Cys residues are replaced by His)\textsuperscript{24,25} would be possible in principle (Fig. 7A). However, the additional Cys ligand is located at position 29 in the sequence rather than at position 44 as expected from sequence alignments of cyanobacterial MTs. Furthermore, if Cys29 would indeed restore an \(\alpha\)-type cluster, the zinc-binding capacity should be increased from three to four as seen for A44H-PflQ2 MT.\textsuperscript{7} However, neither the zinc-binding capacity nor overall binding stability was affected by introducing the additional Cys residue (Fig. S13, ESI\textsuperscript{†}).

Due to the low number of experimentally observed metal–ligand connectivities (Fig. S14 and Table S3, ESI\textsuperscript{†}), it is not possible to directly determine the cluster structure. However, Cys–Cd chemical shifts are highly sensitive indicators to changes in metal binding, and the differences are small between the spectra of Cd\textsubscript{4}PflQ2 MT and its S\textsubscript{29C} mutant. Therefore, it can be excluded that any of the Cys residues (except for C28 that is in the direct vicinity of the mutation) changes from bridging to terminal binding mode (or \textit{vice versa}), and hence the two clusters are most likely highly similar. When these restrictions are added as restraints to the structure calculation in addition to the experimentally observed connectivities, the Cd\textsubscript{4} cluster topology as depicted in Fig. 7C is obtained for the S\textsubscript{29C} mutant while all other topologies cause a larger number of violations of experimental restraints. In the calculated structure, the additional Cys29 acts as a terminal ligand at site C, converting Cys28, that bridges sites A and C in the wild-type structure, into a terminal ligand at site A. Interestingly, Cys10 still serves as a ligand between three metal sites (A, B, C), as observed in the wild-type (Fig. 7B), forming two interchanging arrangements. The resulting cluster structure is also in agreement with the \textsuperscript{113}Cd NMR data: The peak at 549.2 ppm belongs to cadmium site D that has the same Cys\textsubscript{3}His coordination as in the wild-type. Since this peak was the most affected one (see above), we attribute the observed decreased linewidth to a change in the rate of exchange from the intermediate to the fast-regime, while the shift to higher ppm values indicates also a change in the relative populations of the interconverting states: Since the shift occurs to higher ppm, this suggests a higher population of regular tetrahedral tetrathiolate coordination. Interestingly, cadmium site C, that is the most affected by the newly coordinating ligand, does experience very little change in the \textsuperscript{113}Cd chemical shift (585.8 vs. 585.5 ppm), likely because the additional coordination is only transient and because changing coordination from Cys28 to the neighboring Cys29 requires little structural changes.

Overall, the Cd\textsubscript{4} cluster structure of the S\textsubscript{29C} mutant shows that the additional Cys ligand does not drastically change the cluster topology, but increases ligand crowding that promotes the change in coordination and increases dynamics.

The long-range [\textsuperscript{15}N,\textsuperscript{1}H]-HSQC spectrum of Cd\textsubscript{4}S\textsubscript{29C-PflQ2} MT again confirms the involvement of His36 in the coordination (Fig. S15, ESI\textsuperscript{†}), and the \textsuperscript{113}Cd peak at 569.4 ppm in the 1D \textsuperscript{113}Cd spectrum is assigned to the Cys\textsubscript{3}His site (Fig. 5D).
The $^{113}$Cd chemical shifts of the other two peaks (689.9 ppm, 674.9 ppm) are again indicative for polynuclear CdCys$_4$ coordination sites. While for Cd$_2$PflQ2 MT His36, Cys7, Cys32, and Cys49 complete the CdCys$_4$His$_3$ site, the $^{113}$Cd,$^1$H-HSQC-TOCSY spectrum of Cd$_3$S$_{29}$C-PflQ2 MT shows five Cd$^{II}$-ligand correlations, that is to His36, Cys7, Cys29, Cys32, and Cys49 (Fig. S16, ESI†). This observation is, at least at first glance, surprising: We observed in the Cd$^{III}$ titration experiments (Fig. 2) that the molar absorptivity for Cd$^{II}$ titration experiments (Fig. 2) that the molar absorptivity for the fourth Cd$^{II}$ was attributed to the transition of some of the Cys ligands from terminal to bridging coordination mode. However, the much more significant increase for S$_{29}$C-PflQ2 MT suggests that the coordination of the fourth Cd$^{II}$ involves an additional Cys residue, and obviously Cys29 would be the likely candidate for this case. An argument against this assumption is that all Cys-$^1$C$^3$ resonances in the $[^{13}$C,$^1$H]-HSQC spectrum of Cd$_3$S$_{29}$C-PflQ2 MT, including the one of Cys29, are in the chemical shift range (29–33 ppm) typical for deprotonated thiol groups, and deprotonation of Cys residues at or near physiological pH should only occur upon metal coordination (Fig. S17, ESI†). Since a pentacoordinated Cd$^{II}$ ion is highly unlikely and in addition not in line with the observed chemical shift in the 1D $^{113}$Cd spectrum, the five correlations observed in the $[^{13}$Cd,$^1$H]-HSQC-TOCSY spectrum most likely represent an equilibrium between two interchanging CdCys$_3$His$_3$ coordination modes. Hence, while Cys29 is not significantly changing the coordination environment in the Cd$_3$ species, it promotes faster ligand exchange.

Possible functional implications – metal transfer reactions

In order to test the potential functional impact of altered intracluster dynamics, the Cd$^{II}$ ion transfer from the four different MTs to EDTA was probed using ESI-MS. The transfer reaction is completed within 1–10 min for all MTs as no more changes in speciation are observed (Fig. 8 and Fig. S18–S20, ESI†). Generally, there is apparently a very fast transfer of the first Cd$^{III}$ equivalent taking place during the time period needed to collect the first ESI-MS spectra after EDTA addition (time point “1 min” in Fig. S20, ESI†). The predominant difference between the behaviour of PpKT and PflQ2 MTs is that for the entire duration of the experiment, no completely decomplexed MTs are observed for the two PtPkt constructs, while for PflQ2 as well as S$_{29}$C-PflQ2 MT already after 1 min small amounts of the respective apo-species can be detected. Comparing the amount of detected Cd$_3$ species for wild-type and mutant constructs of PflQ2 MT after equilibrium is reached, rather small but peculiar differences are observed (Fig. 8). While generally ESI-MS is considered to be a qualitative method, the similarity of wild-type and respective mutant protein, as well as the identical sample and measurement conditions, allow for some more quantitative interpretations: Mutating Ser to Cys increases the amount of the Cd$_3$ species, while the fraction of the apo and Cd$_3$ species is significantly reduced. Hence, while the fourth Cd$^{III}$ is clearly less tightly bound in the S$_{29}$C mutant (see also Fig. S4, ESI†), the other binding sites are apparently stabilized in the Cd$^{III}$MT form with the additional Cys residue. The difference between PpKT MT and its C28S mutant is also striking: Again, the amounts of Cd$_3$ species detected is higher in the form that contains 10 Cys residues, that is wild-type PpKT MT, while for the C28S mutant the Cd$_3$ species dominates. Hence, while the additional Cys residue seems to cause weaker binding of the fourth Cd$^{II}$ in Cd$_3$S$_{29}$C-PflQ2 MT, it stabilizes the Cd$_3$ species considerably both for S$_{29}$C-PflQ2 and PpKT MT.

The fact that our structural studies show the same protein fold for the Cd$_3$ and Cd$_4$ species of PflQ2 MT as well as its S$_{29}$C mutant argues against a structural role of the additional Cys residue and implies a functional relevance instead. As shown before, the additional Cys residue increases cluster dynamics by providing an additional ligand for metal ion coordination. Hence, it might modulate metal transfer reactions by transiently coordinating metal ions and enabling metal ion transfer without protein unfolding.

Conclusions

MT sequences from different Pseudomonas species exhibit features previously not observed for any other MT. These unique properties have the potential of modulating metal-binding properties and functions of MTs in vivo and presumably reflect the environmental adaptation of Pseudomonas species. One of those characteristics is the presence of a Cys residue in the YCS/CxxC motif in almost 90% of PsmDT sequences identified so far. A comparative study of four different constructs was performed including sequences with a naturally occurring Ser (PflQ2 MT) or Cys (PpKT MT) residue at the indicated position as well as the corresponding S$_{29}$C and C$_{28}$S mutants. Results show that the additional 10th Cys residue is involved in the binding of Cd$^{III}$ ions but is influencing neither the metal-binding capacity nor the overall metal-binding strength. Interestingly, the three-dimensional structure of Cd$_3$S$_{29}$C-PflQ2 MT reveals no...
significant structural influence evoked by the additional Cys residue. Furthermore, although binding of an additional Cys residue now provides enough ligands (11) to build a typical MT x-cluster, the additional ligand instead participates in transient metal-binding resulting in an even more dynamic cluster that interchanges between different coordination modes.

These alterations may be correlated with the type and amount of sub-metallated species formed during the metal transfer reaction to EDTA. For MT forms that feature such an additional Cys residue, the predominant formation of the Cd₃ species is observed, and the amount of sub-metallated species is significantly reduced. Therefore, we hypothesise that the Ser/Cys variation modulates important metal transfer reactions that might be favourable for the overall function of PsdMTs in vivo.

Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>MT</td>
<td>Metallothionein</td>
</tr>
<tr>
<td>bacMTs</td>
<td>Bacterial metallothioneins</td>
</tr>
<tr>
<td>PsdMTs</td>
<td>Metallothioneins from Pseudomonas species</td>
</tr>
<tr>
<td>PfIQ2 MT</td>
<td>Metallothionein from Pseudomonas fluorescens Q2-87</td>
</tr>
<tr>
<td>sh_PfIQ2 MT</td>
<td>C-terminal truncated form of metallothionein from Pseudomonas fluorescens Q2-87</td>
</tr>
<tr>
<td>PpKT MT</td>
<td>Metallothionein from Pseudomonas putida KT2440</td>
</tr>
<tr>
<td>sh_PpKT MT</td>
<td>C-terminal truncated version of metallothionein from Pseudomonas putida KT2440</td>
</tr>
<tr>
<td>SEC</td>
<td>Size-exclusion chromatography</td>
</tr>
<tr>
<td>ESI-MS</td>
<td>Electrospray ionization mass spectrometry</td>
</tr>
<tr>
<td>F-AAS</td>
<td>Flame atomic absorption spectroscopy</td>
</tr>
<tr>
<td>Tris</td>
<td>Tris(hydroxymethyl)aminomethane</td>
</tr>
<tr>
<td>TCEP</td>
<td>Tris(2-carboxyethyl)phosphine</td>
</tr>
<tr>
<td>5F-BAPTA</td>
<td>1,2-Bis(2-amino-5-fluorophenoxy)ethane-N,N,N',N'-tetra-acetic acid</td>
</tr>
<tr>
<td>2-PDS</td>
<td>2,2'-Dithiopyridine</td>
</tr>
<tr>
<td>GST</td>
<td>Glutathione S-transferase</td>
</tr>
<tr>
<td>TEV</td>
<td>Tobacco etch virus</td>
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<tr>
<td>EDTA</td>
<td>Ethylenediaminetetraacetic acid</td>
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</table>

Conflicts of interest

There are no conflicts of interest to declare.

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References


