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Zn^{II} and Hg^{II} binding to a designed peptide that accommodates different coordination geometries†

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Designed metal ion binding peptides offer a variety of applications in both basic science as model systems of more complex metalloproteins, and in biotechnology, e.g. in bioremediation of toxic metal ions, biomining or as artificial enzymes. In this work a peptide (HS: Ac-SCHGDQGSDCSI-NH₂) has been specifically designed for binding of both Zn^{II} and Hq^{II}, i.e. metal ions with different preferences in terms of coordination number, coordination geometry, and to some extent ligand composition. It is demonstrated that HS accommodates both metal ions, and the first coordination sphere, metal ion exchange between peptides, and speciation are characterized as a function of pH using UV-absorption-, synchrotron radiation CD-, ¹H-NMR-, and PAC-spectroscopy as well as potentiometry. Hq^{II} binds to the peptide with very high affinity in a {HgS₂} coordination geometry, bringing together the two cysteinates close to each end of the peptide in a loop structure. Despite the high affinity, Hq^{II} is kinetically labile, exchanging between peptides on the subsecond timescale, as indicated by line broadening in ¹H-NMR. The Zn^{II}-HS system displays more complex speciation, involving monomeric species with coordinating cysteinates, histidine, and a solvent water molecule, as well as HS-Zn^{II}-HS complexes. In summary, the HS peptide displays conformational flexibility, contains many typical metal ion binding groups, and is able to accommodate metal ions with different structural and ligand preferences with high affinity. As such, the HS peptide may be a scaffold offering binding of a variety of metal ions, and potentially serve for metal ion sequestration in biotechnological applications

Introduction

Metal sensor proteins^{1–5} display high selectivity for both essential and toxic metal ions, as demonstrated by representative members of the MerR family,^{6,7} such as the Cu^I-sensing CueR, Zn^{II}-sensing ZntR, and Hg^{II}-sensing MerR.⁸ In this work we have attempted to design a peptide with a broader metal ion binding profile. In a biotechnological perspective, overexpression of such a peptide in suitable bacteria could endow the cells with the capacity to sequester metal ions, including toxic elements, from the environment.⁹ Additionally, elevated levels of such a peptide could ensure metal ion buffering of the

Zn^{II} is rather promiscuous in terms of coordination characteristics as compared to the clearly soft, often two-coordinated

cytosol, allowing the bacterium to survive in harsh conditions of both deprivation and over-exposure to metal ions in the surrounding medium, and serve as an engineered organism with improved properties for biomining and bioremediation.¹⁰⁻¹² The template for the design was the Cu^I binding loop of CueR from V. cholerae, SCPGDQGSDCP. In the related sequence from E. coli, Cu(1) ion is coordinated by two cysteines in a linear coordination geometry.8 The peptide is also expected to possess the capacity to bind the soft HgII ion, due to the thiophilicity of this ion. In order to broaden the metal ion binding profile, and increase the peptide solubility, proline to histidine and proline to serine substitutions were introduced at positions 3 and 11, respectively. The positions of substitutions were chosen to increase ligand-flexibility, and to mimic the presence of His and Ser at these positions in some of the metalloregulatory MerR family members.8 The modifications were expected to promote the coordination of the borderline soft/hard Zn¹¹ ion. In a recent study we demonstrated that this designed 12-mer HS peptide (see Scheme 1) forms various species with Cd^{II}, including loop structures and metal ion bridged bis-ligand complexes, depending on pH and metal to ligand ratio. 13

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Scheme 1 Schematic structure of Ac-SCHGDQGSDCSI-NH₂ (HS).

Hg^{II}. In general, Zn^{II} can easily adopt four-, five- or six-coordinate environments. 14 Nevertheless, in zinc-containing enzymes and proteins the most typical coordination number is four. 14,15 The preference of ZnII for a tetrahedral coordination geometry in proteins is supported by detailed statistical analyses of crystal structures of zinc-containing proteins deposited in the Protein Data Bank (PDB). 16,17 Five- and six-coordinated ZnII centers are typically present due to the complementary coordination of solvent or inhibitor molecules in zinc-containing enzymes.¹⁷ Depending on the type of zinc-centers the abundance of Cys and His side chains significantly varies in the donor set patterns (number and type of bound donor groups). At catalytic zinc-centers any three N, O or S donors of Cys, His, Asp and Glu residues bind Zn^{II} in a 4-5 coordinate distorted-tetrahedral or trigonal-bipyramidal geometry, with His being the predominant ligand. 18 A water molecule is always found in such centres. His and Asp donors are dominant at the co-catalytic zinc-sites consisting of two or three metal ions in close proximity, two of which are bridged by one of the amino acid side chains or a water molecule. 18 Cysteines, however, are not utilized at these motifs. Four protein side chain ligands are bound to ZnII in a tetrahedral or distorted tetrahedral geometry at structural zinc-sites. 18 Such a binding mode is characteristic for e.g. the nucleic acid binding zinc finger proteins15 and for the zinc-clusters in metallothioneins.19 In all classes of the structurally diverse zinc fingers^{20,21} Zn^{II} ions are ligated by a combination of four Cys/ His side chain donors, at least two of which are Cys thiolates. 15 Thiolate donors, complemented with side chain carboxylates and His-imidazoles, are also typical at the metalloregulatory Zn^{II} binding sites in various zinc sensor proteins, however, coordination number and geometry appears to be more decisive in metal ion selectivity than donor ligand types. 15

HgII can tolerate various coordination numbers and geometries, although, six-coordination is much less common than for the other two group 12 metal ions CdII and ZnII. 22 Linear two-coordinate, trigonal planar or T-shaped three-coordinate or tetrahedral four-coordinate structures are representative for complexes with monodentate ligands and higher coordination numbers might be accessible mostly with multidentate compounds.^{22,23} Hg^{II} forms complexes with coordination number 2 more commonly than any other metal ion, 22 which can be explained by relativistic effects.²⁴ Low coordination numbers are characteristic for complexes formed with thiolates, a class of ligands displaying an outstanding affinity towards the large and soft HgII ion,25 and in biological systems HgII is usually

complexed by low molecular weight thiolates or by the Cys side chains of proteins.²⁶ Amongst others, some representative examples are provided by the bacterial mercury resistance systems, e.g. MerP where Hg^{II} is bound to a CXXC (X = amino acid other than cysteine) fragment in a typical linear twocoordinate fashion,27 or the metalloregulatory protein MerR where Cys residues from the two protein monomers form a tricoordinate metal binding site for HgII. 28,29 Additionally, distorted tetrahedral Hg^{II} coordination environment was reported in a few Hg^{II}-substituted proteins. 30-32

The substantially different preferences of ZnII and HgII for four- and two-coordinated structures and the negligible role of His side chains in HgII biocoordination prompted us to investigate whether the His residue incorporated in the flexible ligand sequence of HS might have an influence on the binding of either of the two metal ions. In this work we characterize the binding of ZnII and HgII to the peptide in terms of the metal site coordination geometry and exchange dynamics.

Results and discussion

UV absorption and SRCD studies monitoring the formation of thiolate-metal ion bonds and ligand structure

Comparison of pH-dependent series of UV-spectra in the presence of 0, 0.5 and 1.0 equivalent of HgII or ZnII as compared to the ligand provides information on the interaction of the metal ions with donor groups of **HS**. The occurrence of $S^- \rightarrow$ Hg^{II} ligand to metal charge transfer (LMCT) transitions^{25,33–35} upon the addition HgII to the peptide imply that the cysteine side chain thiolate groups of the ligand are coordinated to Hg^{II} already at low pH (see the full spectra in ESI, Fig. S1A-B†). The Hg^{II}: HS 1:1 system shows almost a constant absorbance at $\lambda = 230$ nm at pH = 4–11 that is significantly higher than the absorption observed for the ligand in the absence of metal ion between pH \sim 4.0-9.0 (Fig. 1). This suggests that the thiolate groups of HS are bound to HgII from acidic to alkaline pH. It is important to note that the deprotonation of the cysteine residues of the free peptide between pH \sim 8-10 is accompanied by the appearance of an n $\rightarrow \sigma^*$ transition around 230-240 nm characteristic for deprotonated thiols36,37 (Fig. 1 and S2†), and as this is absent for the HgII-HS species, we infer that the {HgS₂} coordination geometry is formed at pH lower than 4.

The pH-dependent absorbances detected for the sample containing 0.5 equivalent of HgII compared to HS are in between the values observed for the ligand alone and the

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Fig. 1 Change of the measured absorbances at 230 nm as a function of pH in the Hg^{II}: HS 1:1, 0.5:1 and 0:1 systems ($c_{HS} = 1.0 \times 10^{-4}$ M, I = 0.1 M NaClO₄, T = 298 K).

7.0

8.0

9.0

10.0

11.0

5.0

6.0

HgII-HS 1:1 system at any pH (Fig. 1). This suggests that ~50% of the cysteine residues are bound to Hg^{II} even under acidic conditions and the remaining thiol groups deprotonate in parallel with the free ligand. The spectra recorded in the presence and absence of Hg^{II} reflect that the $S^- \to Hg^{II}$ charge transfer transitions are located below $\lambda = 220$ nm (ε_{215} nm \sim 15 900 M⁻¹ cm⁻¹) independently of the pH and metal ion to ligand ratio (see the difference spectra of Hg^{II}-HS 1:1 and the free ligand in Fig. S3†). Such high energy LMCT transitions and the observed molar absorbances imply that two thiolates are coordinated to the metal ion, as proposed in previous reports on HgII - oligopeptide model systems. 38-42 Three or four HgII-bound thiolates in a trigonal/tetrahedral coordination geometry would result in LMCT peaks or shoulders at lower energies^{25,31,35,40,42-44} which is not observed here even in the excess of HS over HgII indicating that metal ion bridged species are not formed.

In contrast to Hg^{II} , the LMCT band characteristic for S^--Zn^{II} interactions in zinc(II)-bound proteins^{32–34} and peptides^{45–49} emerges only above pH ~ 5.0 in the solutions of Zn^{II} and HS (Fig. 2 and S4A–B†), reflecting the expected, substantially weaker affinity of Zn^{II} towards the ligand. A remarkable spectral change, *i.e.* a further absorbance increase occurs above pH ~ 7.5 in the presence of one equivalent Zn^{II} per HS. A similar, but less pronounced spectral change, attributed to the formation of hydroxo mixed ligand species, was also observed in the Zn^{II} -complex of a related 12-mer peptide,⁵⁰ however, at a higher pH. Thus, the metal bound water appears to display a lower pK_a of 8.65 in the Zn^{II} -HS complex, *vide infra* (potentiometric data).

The $A_{230~\rm nm}$ νs . pH curve obtained for the $\rm Zn^{II}$ -HS 0.5:1 sample runs in between those of the free peptide and the equimolar system in the whole studied pH-range (Fig. 2). The observed profile is closer to that seen in the presence of 1 eq. $\rm Zn^{II}$ between pH 5–9, contrary to the data recorded for $\rm Hg^{II}$. Thus, a more complex speciation must occur for $\rm Zn^{II}$, with more than half of the thiolates bound to the metal ion at a

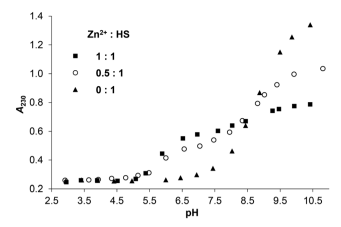


Fig. 2 Change of the measured absorbances at 230 nm as a function of pH in the Zn^{II}: HS 1:1, 0.5:1 and 0:1 systems ($c_{HS} = 1.0 \times 10^{-4}$ M, I = 0.1 M NaClO₄, T = 298 K).

stoichiometry of 0.5:1 Zn^{II}: HS, indicating the formation of metal bridged species. At high pH, however, the absorbances detected for twofold ligand excess seem to be *ca.* the averages of those of the free ligand and the equimolar sample (see Fig. 2, S2 and S4†), suggesting similar speciation at any metal ion to ligand ratios.

In order to gain information on the metal ion induced conformational change of the peptide SRCD (synchrotron radiation circular dichroism) spectra were recorded both for Hg^{II} and Zn^{II} complexes. Previously we have demonstrated that HS displays a disordered structure with varying levels of transient helicities, 13 represented by an intense negative CD-extremum slightly below 200 nm and a less intensive shoulder around 220 nm. 34,47,51,52 Addition of Hg^{II} to the acidic solution of HS results in a notable decrease of the negative peak at $\lambda \sim$ 198 nm while the shoulder is less affected (Fig. 3). A similar type of change was reported to accompany the Hg^{II} -coordi-

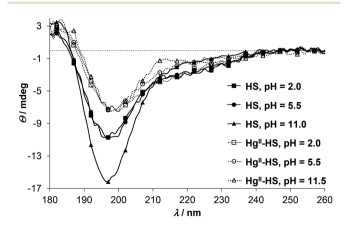


Fig. 3 SRCD spectra of HS in the absence (continuous lines with filled markers) and presence of 1.0 eq. of Hg^{II} (dotted lines with open markers) at selected pH values ($c_{\rm HS} = 1.0 \times 10^{-3}$ M, I = 0.1 M NaClO₄, T = 298 K, I = 0.1 mm).

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nation of a 18-mer peptide, comprising the metal binding loop of MerP possessing a CAAC motif. 53,54 The spectral change was assigned to the folding of the peptide to a thermodynamically (but not necessarily kinetically) stable conformation,⁵³ although the reduction of the negative ellipticity around 200 nm was also observed with other metal ions and two other peptide derivatives with alterations in the metal binding sequence (CCAA and CACA).⁵⁴ By all accounts, Hg^{II}-binding to HS clearly induces a conformational change of the ligand towards a loop structure, presumably similar to the metalloaded forms of CueR.8 One, however, has to bear in mind that due to the high energy ligand to metal charge transfer bands of the Hg^{II}-bound species, CD features of these bands may overlap with the backbone-related CD-effects. This is a known problem in the interpretation of the secondary structures of metalloproteins and metal ion-peptide complexes, 34,45,55,56 particularly when relatively small molecules, like the present 12-mer HS peptide, are studied. Distinction of the different contributions may be easier when thiolate to metal ion transitions appear separately at lower energies compared to the peptide backbone bands, like in the tetrahedral {CysS₄} type Hg^{II}-rubredoxin complex³¹ or in metallothioneins, where metal induced bands dominate the wavelength region above 220–230 nm. ⁵⁷ Comparison of the SRCD spectra of **HS** at pH \sim 2.0 in the presence and absence of HgII (Fig. 3) suggests that any effect of the HgII-binding of the thiolate donors dominate below $\lambda \sim 210$ nm. The increase of pH has practically no further effect on the ellipticity around 198 nm for the Hg^{II}-HS complex, however, it slightly influences the lower energy shoulders. This can be assigned to the deprotonation of the Asp and His residues of the peptide inducing modest changes in the backbone of the loop-forming ligand.

Zn^{II} has no impact on the SRCD spectra of **HS** up to pH 5.5 (Fig. 4), which correlates well with the UV-spectra where the LMCT bands emerge above pH \sim 5. At higher pH, however, the

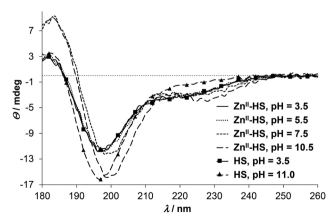


Fig. 4 pH-Dependent SRCD spectra recorded in the Zn^{II}-HS 1:1 system. pH = 3.5: continuous line; pH = 5.5: dotted line; pH = 7.5: short dashes; pH = 10.5: long dashes. For comparison, spectra of the free ligand is also shown at pH = 3.5 (continuous line with squares) and 11.0 (long dashes with triangles) ($c_{HS} = 1.0 \times 10^{-3}$ M, I = 0.1 M NaClO₄, T =

position of the main negative CD-minimum is slightly redshifted (see spectra at pH = 7.5 and 10.5 on Fig. 4), while the ellipticities around 180 and 230 nm are remarkably increased, as compared to the spectra of the free ligand. As hinted already for Hg^{II}, influences of the S⁻-Zn²⁺ chromophore and the peptide secondary structure may be superposed in the observed CD-pattern of ZnII-protein/peptide structures. 34,55,56 Nevertheless, the direction of the observed changes is rather similar to the ZnII-induced effects on the conformation of a phytochelatin analogue⁴⁷ and other relatively short oligopeptides^{52,58} and may suggest an increasing helical content^{47,52} in the Zn^{II}-bound **HS**. It was proposed that different coordination properties of metal ions may develop selectivity in the stabilization of the α-helical conformation of 20-mer peptides.⁵⁸ The fundamentally distinct CD-features of **HS** in the presence of Hg^{II} and Zn^{II} may imply that the different coordination geometry preference of the two metal ions promote large dissimilarity between the HgII- and Zn^{II}-bound structures of the ligand. The characteristic shoulder seen in the spectra of ZnII-HS (Fig. 4 and S5†) starts to develop from ca. pH 6 (data not shown) but increases up to pH 9.5-10. The ZnII: HS ratio dependence of the discussed CD-peak at pH 10.5 reflects a simple equilibrium between the free and ZnII-bound HS (Fig. S5†).

^{199m}Hg PAC spectroscopy for the elucidation of the coordination environment of HgII

The local environment and coordination geometry of Hg^{II} was also monitored by 199mHg PAC (perturbed angular correlation of γ-rays) spectroscopy in the presence of one equivalent metal ion at pH = 2.0 and pH = 8.0. The fundamentals of PAC spectroscopy and the interpretation of the parameters obtained by the technique are described in detail in the literature.⁵⁹ The PAC data may be analyzed with one nuclear quadrupole interaction (NQI) at each pH, and the PAC parameters (ν_0 , the nuclear quadrupole coupling constant, and η , the so called asymmetry parameter, which is zero for an axially symmetric coordination geometry) for the observed NQIs are collected in Table 1. The fitted $\nu_{\rm O}$ and η values are similar at pH 2.0 and 8.0 and comparable to literature data obtained for compounds with two-coordinate {HgS2} structures^{60,61} (Table 1). The spectrum recorded at low pH is slightly more complex than that obtained at pH \sim 8.0 as reflected in the lower signal amplitude and the broader and less visible second and third peaks, respectively (Fig. 5). This may suggest the co-existence of a small amount of species with a different structure, nevertheless, the main spectral features, with a support of UV-data, clearly indicate that the major component has a two-coordinate {HgS₂} coordination mode.

Potentiometric investigation of distribution and stabilities of the species formed in the ZnII: HS system

The formation constants ($\log \beta$) determined for the proton and Zn^{II} complexes of **HS** are summarized in Table 2.

The ligand undergoes five (de)protonation processes in the studied pH-range that were attributed to the carboxylate

Table 1 PAC parameters fitted for Hg^{II}: HS and for different Hg^{II}-thio-

late complexes of known structures

System/pH	$ u_{ m Q}/{ m GHz}$	η	Coordination geometry	Ref.
Hg ^{II} -HS 1:1 $(pH = 2.0)$	1.43(5)	0.07(6)	Two-coordinate, 2 thiolates	This work
Hg^{II} -HS 1:1 (pH = 8.0)	1.43(1)	0.13(3)	Two-coordinate, 2 thiolates	This work
[Hg(Cysteine) ₂]	1.41	0.15	Two-coordinate, 2 thiolates	60
Ac-Cys-dPro- Pro-Cys-NH ₂	1.42	0.19	Two-coordinate, 2 thiolates	41
MerA (77 K)	1.42	0.15	Two-coordinate, 2 thiolates	61
MerR (77 K)	1.18	0.25	Three-coordinate, 3 thiolates	61
Hg-rubredoxin	0.10	0 (fixed)	Four-coordinate, 4 thiolates	31

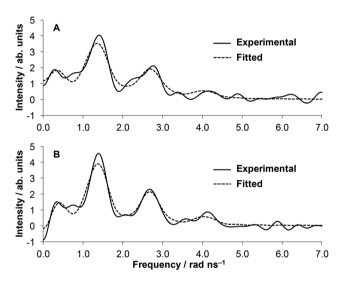


Fig. 5 Fourier transformed experimental (solid lines) and fitted (dashed lines) 199m Hg PAC data of the Hg II : HS 1:1 system at pH = 2.0 (A) and pH = 8.0 (B) $(c_{Hg^{||}} = c_{HS} = 8.03 \times 10^{-5} \text{ M}).$

groups of two Asp residues (pH \sim 3-5), the imidazole side chain of His (pH \sim 6-7.2) and the thiol moieties of the two Cys units (pH ~ 7.8-9.7). The deprotonation constants (p K_a) of the ligand have been re-determined for the present study and are in a good agreement with those published earlier. 13

In Table 2 the species model obtained by best fit of the Zn^{II}: HS system titration curves is presented. Introducing bisligand complexes (ZnHxL2) in the model was necessary for the correct description of titration data when HS was used in excess over Zn^{II} (see Experimental). Contrary to this, considering the presence of dinuclear species (Zn₂H_xL), did not improve the fit of the experimental data neither for the Zn^{II}: HS 0.5:1 and 1:1 samples nor for those containing a two-fold Zn^{II}-excess over the ligand (the latter was evaluated only up to pH \sim 7).

Table 2 Formation constants (log β) of the Zn^{II} complexes of HS (estimated errors in parentheses (last digit)) and derived equilibrium data $(I = 0.1 \text{ M NaClO}_4, T = 298 \text{ K})$

Species ^a	pqr^b	$\log \beta_{pqr}$		pK_{pqr}^{c} , $\log K_2^{d}$
[ZnHL]	111	16.58(4)	p <i>K</i> ₁₁₁	5.95
$[ZnL]^{2^{\frac{1}{n}}}$	101	10.63(4)	pK_{101}	8.65
$[ZnH_{-1}L]^{3-e}$	1-11	1.98(5)		
$[ZnH_2L_2]^{4-}$	122	31.3(2)	pK_{122}	7.6
$[ZnHL_2]^{5-}$	112	23.7(1)	pK_{112}	8.7
$[ZnL_2]^{6-1}$	102	15.0(2)		
NP^f		544	$\log {K_2}^{ m L}$	4.37
$FP^g(cm^3)$		0.005		

 $a \log \beta$ values for the protonation processes of **HS**, re-determined in the present study are: $\log \beta_{051} = 31.68$, $\log \beta_{041} = 28.33$, $\log \beta_{031} = 24.10$, $\log \beta_{021} = 17.50$, $\log \beta_{011} = 9.06$. $^b p, q$, and r reflect stoichiometric numbers of the fundamental components the complex species are composed of, as defined in the experimental section. ${}^c p K_{pqr} = \log \beta_{pqr} - \log \beta_{p(q-1)r}$, ${}^d \log K_2 = \log \beta_{102} - \log \beta_{101}$, ${}^e H_{-1}$ represents an extra deprotonation, beyond the proton releases of the peptide, *i.e.* deprotonation of a Zn^{II}-bound water molecule. ${}^f NP =$ number of points. g FP = fitting parameter representing an average deviation in cm3 between the experimental and fitted data for the full data set (including all evaluated titrations).

Complex formation processes start from pH ~ 4.5 by the appearance of a protonated mono-complex ZnHL as reflected by the calculated species distributions (Fig. 6). A consecutive deprotonation process ZnHL → ZnL + H⁺ leads to the formation of the parent complex ZnL where all of the dissociable protons of the peptide are already released. The pK_a value for this process (= 5.95, see Table 2) is significantly lower than those attributed to the deprotonation processes of the HL and H_2L forms of the free ligand (p K_{HL} = 9.06, p K_{H_2L} = 8.44) and somewhat below the third pK_a of HS ($pK_{H_3L} = 6.60$). This strongly suggests that at least two, but potentially all the three neutral/basic donor groups of the ligand (histidine imidazole and two cysteine thiolates) are bound to ZnII in the ZnL species. Coordination of both cysteines to ZnII in ZnL is also supported by the observed absorbance increase in parallel with the formation of ZnHL/ZnL (A230 traces are overlaid with species distributions calculated for the concentration of UV data, see Fig. S6A-B†).

The determined stability of ZnL (log K = 10.63) reflects a remarkable affinity of HS to ZnII. This stability constant is, indeed, several orders of magnitude higher than those of the parent Zn^{II} complexes of shorter peptides containing a CXH motif,62 but also surpasses the stabilities of terminally protected tripeptides composed of a CXC sequence, 62 in spite of the substantially longer peptide chain and the larger distance between the two Cys residues in HS. Besides, HS has a notably higher affinity to ZnII compared to a similar 12-mer oligopeptide possessing no histidine residues (studied by us, $\log K$ = 9.93).⁵⁰ Although higher stabilities were found for the Zn^{II} complexes of some 10-mer peptides, all of these contained 2-3 histidines in addition to the two cysteine units. 63 Thus, the affinity of HS for ZnII falls in range that indicates the coordination of both cysteine and the histidine residues to the

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Dalton Transactions

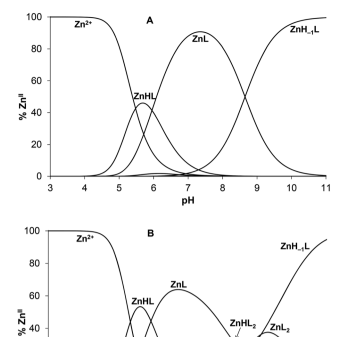


Fig. 6 Species distribution diagram for the Zn^{II}: HS 1:1 (A) and 0.5:1 (B) systems ($c_{HS} = 1.0 \times 10^{-3}$ M). The speciation curves for the concentrations applied in the UV experiments are depicted in ESI (Fig. S6A-B†).

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metal ion. The ZnII-binding affinity of HS can also be demonstrated by the conditional stability calculated at pH 7.4 and 1:1 metal ion to ligand ratio based on the equations below,

$$H_qL + Zn \rightleftarrows ZnH_qL$$

$$\mathit{K}_{\mathrm{a}} = \frac{[\mathrm{ZnH}_{q}\mathrm{L}]}{[\mathrm{Zn}][\mathrm{H}_{q}\mathrm{L}]}$$

where Zn denotes the free Zn^{II} concentration while $H_{\alpha}L$ and ZnHaL represent the overall concentration of the free and complexed ligands in any protonation states, respectively. The apparent stability constant for the above conditions is $K_a = 7.5$ \times 10⁷ (log $K_a = 7.9$) which is in the lower range of affinities reported for various wild-type bacterial ZnII-regulators64 or variants.65

The deprotonation of ZnL above pH ~ 8 (Fig. 6A) leads to the species ZnH₋₁L being strongly dominant under alkaline conditions. The observed extra deprotonation is most likely not a ligand-related proton release since the formation of a Zn^{II}-amide bond is a very scarce event in the complexes of Zn^{II} formed with terminally protected peptides. 62,66-69 Accordingly, the ZnH₋₁L composition may represent a species with a deprotonated water ligand, described as Zn(OH)L. The pK_a value of the deprotonation process is 8.65 that is ca. 1.7 log units

lower than the pK_a determined for the same type of proton release of the CdII complex of HS,13 as expected, due to the smaller ionic radius of ZnII as compared to CdII. The deprotonation of the bound H₂O occurs also at a somewhat lower pH than in the ZnL complex of a similar ligand containing no His residue in position 3 of the peptide chain ($pK_a = 9.11^{50}$). The {Zn(Cys)₂HisH₂O/OH⁻} coordination sphere is also found in horse liver alcohol dehydrogenase (LADH), where the pK_a of the metal ion bound water molecule is 9.2 for the native ZnII containing enzyme, 70 and 11.0 for the CdII substituted species.⁷¹ Interestingly, the pK_a of the Zn^{II} -bound water is lower in HS than in LADH. It seems that above neutral pH the histidine of HS significantly influences speciation, the coordination sphere of ZnII and even the peptide structure, as indicated by UV and SRCD data.

Monomeric ZnHL and ZnL complexes dominate in the acidic/neutral pH-range when HS is present in a twofold excess over ZnII (Fig. 6B). As indicated by UV data, metal-bridged bis-ligand species with different protonation states are also formed above pH ~ 6. Although the determined stabilities of the various bis-complexes do not provide direct information on the binding mode of the ligands, the relatively high pK_a value for the $ZnHL_2 \rightarrow ZnL_2 + H^+$ process (= 8.7, Table 2) suggests that there are protonated thiol groups in the ZnH₂L₂ and ZnHL2 species. The stability constant calculated for the binding of the second ligand in ZnL_2 (log $K_2 = 4.37$) and the relative stability of the parent mono- and bis-complexes $(\log(K_1/K_2) = 6.26)$ shows a notably weaker binding of the second ligand as compared to the same process in the CdII: **HS** system ($\log(K_1/K_2) = 5.33^{13}$) or to the Zn^{II}-binding of the above cited His-free peptide ($log(K_1/K_2) = 5.14^{50}$). This finding provides a further support for the important role of histidine in controlling the interaction of ZnII with HS.

¹H NMR experiments

Assignment of the 1H NMR resonances of HS and the pHdependence of the recorded spectra in the absence of metal ions were published previously. 13 Hg II coordination to the peptide has a strong effect on the resonances of the Cys C₆H₂ protons (Fig. 7). These signals shift from 2.93 ppm to relatively broad peaks at ~3.3-3.4 ppm (in an accidental overlap with one of the His C_6H_2 resonances at pH 4.0-6.0) in the presence of one equivalent of HgII. The significant, ca. 0.4 ppm downfield shift of the Cys C_βH₂ resonances of the bound ligand, as compared to the same signals of the free HS indicates the binding of both thiolates to the metal ion, as also suggested by UV titrations.

Two separate signal sets of the Cys C6H2 protons are observed at pH 4.0-6.0 when HS is in a twofold excess over Hg^{II} (Fig. 7) One set is reminiscent of the resonances of the free ligand, whereas the other coincides with those observed in the HgII: HS 1:1 system. Increasing pH to 8.0 results in coalescence of the two signal sets to a very broad bulge-like feature in the range of $\delta \sim 2.8-3.4$ ppm overlapping with the His C_βH₂ resonances (Fig. 7). This coalesced signal, with a chemical shift found in between those observed for Hg^{II}: HS

Fig. 7 Selected regions of the 1H NMR spectra of HS recorded in the absence and presence of 0.5 and 1.0 eq. of Hg II (H $_2O/D_2O=90:10\%$ v/v, $c_{HS}=1.3\times10^{-3}$ M, T=298 K). The resonances marked by symbols are: His $C_{\beta}H_2$: \blacktriangle ; Cys $C_{\beta}H_2$: O; Asp $C_{\beta}H_2$: \blacksquare . The region of $\delta=2.7-3.6$ ppm from the Hg II : HS 0.5:1 spectrum at pH = 8.0 is magnified in the frame. Note that the sharp signals at $\delta\sim1.9$ ppm are those of the acetate anion of the added mercury(II) salt which coincide with the acetyl protecting group resonances of HS at pH ~4.0 .

1.6

2.4

4.0

0.8 ppm

1:1 and the free ligand, becomes sharper on increasing pH but is still broad at pH = 10.0. These findings indicate that the ligand exchange rate between the free and bound forms gradually increases from the slow/intermediate to the intermediate/ fast time regime in parallel with the deprotonation of the unbound thiol groups of the presumably free ligand being present in the Hg^{II} : HS 0.5:1 system.

The exchange rate, $k_{\rm ex}$, between the bound and non-bound ligand forms may be roughly estimated from the observed line-broadening⁷² at pH 4.0–6.0 which is dominated by slow exchange. The line-broadening, $w_{\rm e}$ – $w_{\rm o}$, occurring for the Cys $C_{\rm \beta}H_2$ resonances of the free peptide due to the addition of 0.5 eq. Hg^{II} is *ca.* 12 Hz which leads to $k_{\rm ex} \sim \pi \times (w_{\rm e} - w_{\rm o}) \sim 38~{\rm s}^{-1}$ at pH = 6.0 ($w_{\rm e}$ and $w_{\rm o}$ represent the line width of signals at half height with and without exchange, respectively). The calculation is based on the assumption of a two-site exchange of HS between a specific Hg^{II}-peptide bound form and the non-bound form under the applied experimental conditions. $k_{\rm ex}$ may also be expressed by a formulae involving the rates of the association and dissociation processes, as follows⁷³

$$k_{\rm ex} = k_{
m on}[{
m M}] + k_{
m off} = rac{k_{
m off}}{1-f}$$

where $k_{\rm on}$ and $k_{\rm off}$ stands for the second order rate constant of the complex formation and the first order rate constant of the complex dissociation, respectively, [M] is the concentration of the metal ion and f represents the bound fraction of the ligand (0.5 in the present case). If one assumes that the association is diffusion controlled and thus $k_{\rm on} \sim 7.4 \times 10^9~{\rm M}^{-1}~{\rm s}^{-1}$ that is the rate constant for diffusion controlled reactions in water at 298 K,⁷⁴ solving the above equation for [M] would result in a concentration of [Hg^{II}] that corresponds to a $K_{\rm d} \sim 2.5 \times 10^{-9}~{\rm M}$ dissociation constant of the complex at pH = 6.0. This $K_{\rm d}$ value suggests a many orders of magnitude weaker binding than expected for a typical {HgS}₂} complex,⁴¹ indicating that the picture is too simplistic, and presumably a more complex speciation occurs.

The increase of pH also induces the upfield shift of the resonances of the Asp ($C_\beta H_2$ protons – Fig. 7) and His ($C_\beta H_2$ – Fig. 7 and the $C_{\epsilon 1}H$ and $C_{\delta 2}H$ protons of the imidazole ring – Fig. 8) reflecting the deprotonation of the side chains of these residues. The chemical shift values are practically independent of the metal ion to ligand ratio at all selected pH values. These findings indicate that the proton releases from the Asp carboxyl groups and the His imidazole moiety are practically unaffected by the presence of Hg^{II} and therefore that these groups do not participate in Hg^{II} -binding. Nevertheless, coordination of the cysteine residues to Hg^{II} has a slight line

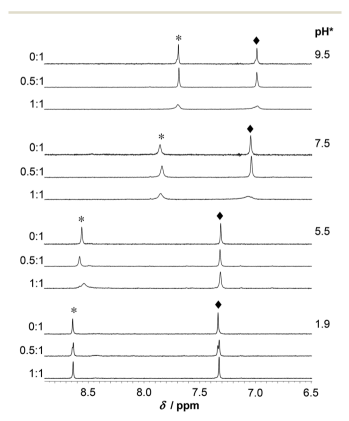


Fig. 8 Part of the 1H NMR spectra of the peptide HS recorded at various pH* values in D₂O as a function of the Hg^{II} to peptide ratio (c_{HS} = 1.3 × 10⁻³ M, T = 298 K). The symbols denote the C_{e1}H (*) and C₆₂H (\spadesuit) resonances of His.

width increasing effect on the neighbouring Asp side chain resonances under acidic conditions (Fig. 7) and a rather pronounced impact on the His $C_{e1}H$ and $C_{\delta 2}H$ signals in neutral/alkaline solutions (Fig. 8). This shows that although the chemical shifts, apart from those of the cysteines, do not change significantly, the dynamics of the peptide is affected by the binding of Hg^{II} .

The spectra of **HS** obtained at pH ~ 4.4 in the presence and absence of ZnII reflect no differences either in terms of the chemical shifts or the shape of the various ¹H-resonances (Fig. S7†). This suggests that, as opposed to Hg^{II}, Zn^{II} is not bound to HS under such conditions, which is in agreement with the potentiometric and UV absorption studies, vide supra. At pH ~ 5.5, however, the presence of Zn^{II} gives rise to pronounced broadening of most resonances. In the presence of 0.5 eq. Zn^{II} this may indicate exchange between the bound and free states of the peptide, but in the fully loaded ZnII: HS system it implies equilibria between conformers falling into the intermediate exchange time regime (ms-s) (Fig. 9-10). At a 1:1 ratio of Zn^{II} and HS the $C_{\epsilon 1}H$ and $C_{\delta 2}H$ signals of the His imidazole are shifted slightly upfield as compared to the resonances of the free ligand (Fig. 10). At 0.5 eq. of ZnII the chemical shifts of the imidazole ring protons appear in between those of the free HS and the 1:1 system reflecting an equilibrium between the non-bound and metal-bound peptide forms, and fast exchange dynamics for these resonances. The C_{s1}H and C₈₂H resonances are significantly shifted upfield by a further pH increase (pH $5.5 \rightarrow 7.0$), similarly to the metal ion free solution, which indicates that His-coordination is not completed at pH 5.5. A combined interpretation of the ¹H NMR, UV absorption, and potentiometric data at pH \sim 5.5, (see Fig. 6) leads us to propose co-existing binding isomers of the ZnHL species, with the participation of two Cys-thiolates or one of the Cys-thiolates and the His side chain in metal ion binding.

The increase of pH to pH \sim 7.0 gives rise to a substantial change of the spectral pattern. According to our data, all the metal ions are complexed under such conditions (Fig. 6). Most of the resonances, in addition to those of the Cys residues, display line broadening, in contrast to the signals observed for Hg^{II}: HS, where resonances from non-coordinating groups are not affected to the same extent. Thus, ZnII-coordination affects the internal dynamics of the entire peptide on the NMR time scale. Additionally, the ligand exchange dynamics is slowed down to the moderately slow exchange time regime causing the splitting of several ¹H resonances (C_βH₂, C_{ε1}H and C_{δ2}H of His and all the resonances of Ile) into clearly distinguishable separate signals at a 0.5:1 Zn^{II}: HS ratio (see spectra at pH ≥ 7.0 on Fig. 9-10). The decrease of exchange rate by pH-increase coincides with a remarkable change of the CD-signals (Fig. 4), and occurs in parallel with the formation of the ZnL parent complex. It implies that the participation of several donor groups in metal ion binding leads to a reduced lability of species. At a 1:1 Zn^{II} : HS ratio, the $C_{\beta}H_2$ protons of the Cys and His residues experience a significant chemical shift change relative to the free ligand, as do the $C_{\epsilon 1}H$ and $C_{\delta 2}H$

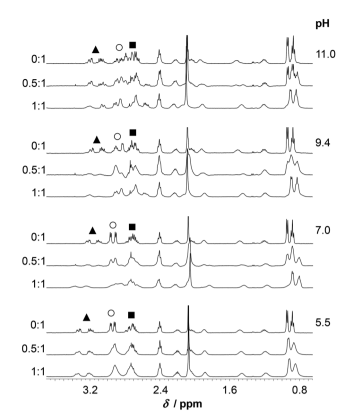


Fig. 9 Aliphatic region of the 1 H NMR spectra of HS recorded in the absence of Zn II and in the Zn II : HS 0.5:1 and 1:1 systems (H₂O/D₂O = 90:10% v/v, $c_{HS}=1.3\times10^{-3}$ M, T=298 K). $C_{\beta}H_{2}$ resonances of the residues with potential donor groups are indicated by the following symbols: His $C_{\theta}H_{2}$: \blacktriangle ; Cys $C_{\theta}H_{2}$: \bigcirc ; Asp $C_{\theta}H_{2}$: \blacksquare .

signals of the imidazole ring (Fig. 10). This supports the coordination of the two Cys-thiolates and the His-imidazole groups to ZnII in ZnL, but the poorly resolved spectrum at pH = 7.0 does not provide information on the binding of Aspcarboxylates. As pointed out above, various resonances of the C-terminal Ile residue in the spectral region 0.8-1.0 ppm $(C_8H_3, C_{y2}H_3)$ are also strongly affected by metal ion coordination as those of the bound ligand are clearly shifted upfield compared to the ones of the free peptide-like resonances (Zn^{II}: HS 0.5:1, Fig. 9). Analogous spectral features were not observed in the systems of either CdII and HS13 or ZnII and a closely related peptide⁵⁰ differing only in the His-residue from the presently studied ligand. Thus, while the exact origin of the impact of ZnII-binding on the Ile resonances is not clear, metal ion coordination of the histidine unit very likely plays a key role here.

Based on the observed line broadening of the Cys $C_{\beta}H_2$ protons at Zn^{II} : HS 0.5:1 (Fig. 9) a similar or slightly lower exchange rate between the bound and non-bound ligands, as compared to Hg^{II} : HS, may be predicted. However, the overlap of the various resonances and the complexity of the system (see the distribution curves at $pH \sim 7.0$, Fig. 6B) do not allow a deeper discussion. It is, however, an interesting contrast to Hg^{II} : HS, that the exchange rate in the presence of Zn^{II}

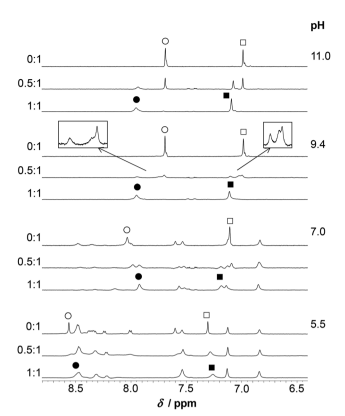


Fig. 10 Aromatic/H $_{N}$ region of the 1 H NMR spectra of HS recorded in the absence of Zn II and in the Zn II : HS 0.5:1 and 1:1 systems (H $_{2}$ O/D $_{2}$ O = 90:10% v/v, c_{HS} = 1.3 × 10 $^{-3}$ M, T = 298 K). The open circle and square symbols show resonances of the non-bound ligand (C $_{e1}$ H and C $_{62}$ H, respectively) while the filled symbols mark the same resonances of the bound His residues in the mononuclear species. The regions of δ = 6.9–7.15 and 7.4–8.05 ppm from the Zn II : HS 0.5:1 spectrum at pH \sim 9.4 are magnified in the frames.

remains relatively slow even at higher pH (see below) approaching the deprotonation-range of the thiol groups of the free ligand. The increased metal ion exchange rate, as observed by the resonances of the Cys C_6H_2 protons (see Fig. 7) with 0.5 eq. Hg^{II} for pH above the p K_a of the thiols, imply that the free thiolates take part in the exchange process, and thus that it occurs via an associative mechanism. The low coordination number may be important for this process, as it may allow for coordination of additional thiolates in the equatorial plane. This is analogous to a proposed mechanism of transfer of Cu^I between proteins, 42,75,76 where the metal ion is also found in a structure with two thiolates coordinating. Contrary to this, the metal exchange rate does not change into the fast exchange regime with 0.5 eq. Zn^{II} for pH above the p K_a for the thiols, see Fig. 9. This may reflect that the exchange occurs via a dissociative mechanism, although not necessarily via free ZnII, in analogy to the common interpretation of ligand binding reactions for the ZnII aqua ion involving dissociation of coordinated water as the rate determining step.⁷⁷

As a conclusion, and in line with SRCD data, the simultaneous binding of (at least) three side chain donors induces a

more defined ligand structure in the Zn^{II} -bound **HS**, unlike the loop-like conformation proposed for Hg^{II} : **HS**.

At pH ~ 9.4 the spectra of the Zn^{II}-containing solutions are still very poorly resolved as the resonances are strongly broadened. Coordination of His to the metal ion in the $\mathrm{Zn^{II}}: HS \ 1:1$ system is unambiguously demonstrated by the significant downfield shift of the $C_{\epsilon 1}H$ and $C_{\delta 2}H$ resonances, as compared to the metal ion free sample (Fig. 10). A similar shift was observed and attributed to His-coordination in the CdIIcomplex of the peptide. 13 At least three distinguishable, broad $C_{\delta 2}H$ peaks and three $C_{\epsilon 1}H$ peaks, albeit less clearly, are observed at 0.5:1 Zn^{II}: peptide ratio (see the enlarged spectrum segments on Fig. 10). One of the $C_{\delta 2}H$ and $C_{\epsilon 1}H$ peaks appear very close to the free ligand-like signals while the third observed signals have chemical shifts resembling those measured for the 1:1 system. In order to elucidate the processes occurring in the presence of ligand excess above neutral pH, a more detailed series of spectra were recorded at pH ~ 8. One may follow the evolution of the $C_{\epsilon 1}H$ and $C_{\delta 2}H$ signals from 0:1 to 1:1 ZnII: peptide ratio on Fig. 11. The series describe the complete transformation of the non-bound ligand to the 1:1 species (mostly ZnL at this pH, see Fig. 6A). The spectra recorded at the intermediate stages, featuring the emergence and transformation of broad peaks, originate from a dynamic exchange between at least three coexisting species (see e.g. the $C_{\epsilon 1}H$ resonances at a $Zn^{II}:HS$ ratio of 0.75:1 or the $C_{82}H$ resonances at a ratio of 0.5:1), the free peptide, the fully loaded ZnII-peptide complex, and a species with a plausible 0.5:1 ZnII: HS stoichiometry, i.e. a bis-ligand complex.

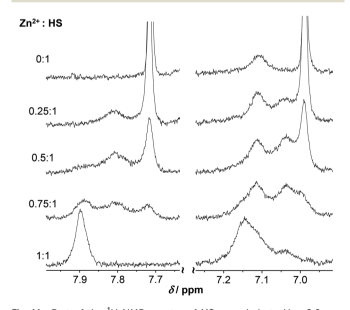


Fig. 11 Part of the ^1H NMR spectra of HS recorded at pH ~ 8.0 as a function of the metal ion to peptide ratio (H₂O/D₂O = 90:10% v/v, $c_{\text{HS}} = 1.3 \times 10^{-3}$ M, T = 298 K). The selected regions reveal changes observed on the $C_{e1}\text{H}$ (left) and $C_{\delta2}\text{H}$ (right) resonance of the His-imidazole moiety of the ligand. Note that the peak seen at $\delta \sim 7.11$ ppm on the free HS spectrum is one of the remaining amide resonances still observable at pH ~ 8.0 .

This provides strong support for the potentiometric data. The species are in slow to intermediate exchange rate relative to the NMR timescale. The emerging signals in the range of $\delta \sim 7.0$ –7.08 and 7.75–7.85 ppm suggests that the His-imidazole moiety of at least one of the two ligands plays a role in Zn^{II}-coordination in some or all of the bis-complexes.

Only two sets of relatively sharp $C_{\epsilon 1}H$ and $C_{\delta 2}H$ resonances are detected at pH \sim 11 in the Zn^{II} : HS 0.5:1 system (Fig. 10) which is in excellent correlation with the equilibrium and UV results, *i.e.* the presence of only the $ZnH_{-1}L$ complex and free ligand. Besides, the notable chemical shift changes observed on the spectra of Zn^{II} : HS 1:1 from pH \sim 7.0 up to 11.0 (see *e.g.* the range of $\delta \sim$ 3.5–2.5 ppm (Fig. 9) or the $C_{\delta 2}H$ signals (Fig. 10), clearly reflect the conversion of the ZnL parent complex to $ZnH_{-1}L$.

Conclusion

The 12-mer HS peptide, inspired by the C-terminal metal ion binding domain of a bacterial metalloregulator CueR, is shown to efficiently bind HgII and ZnII, two metal ions with significantly different coordination preferences. HgII is demonstrated to form a loop-like structure in a {HgS₂} coordination fashion via binding to the two cysteinates of the ligand, but there is no sign for the participation of any other side chain donors in HgII-coordination. The kinetic lability of HgII is manifested in line broadening on the ¹H NMR spectra affecting mostly the resonances of the bound Cys residues and those of the neighbouring units. In contrast to HgII, ZnII dictates the peptide to a more structured form in its ZnL complex via binding to at least three side chain donors, the two Cys thiolates and the His imidazole. Indeed, the SRCD spectra above neutral pH might reflect an increasing helical content in the Zn^{II}-bound HS, although a contribution of the thiolatemetal ion chromophore to the observed CD-pattern may also be present in the same wavelength range. In addition to monomeric species, bis-ligand HS-ZnII-HS complexes are also formed, unlike with HgII. The line broadening in 1H-NMR is pronounced for most of the resonances, indicating that exchange dynamics between different conformers occurs on the NMR time scale (ms-s) and that, in contrast to HgII, ZnIIcoordination notably affects the internal dynamics of the entire peptide chain. The results obtained demonstrate that the conformational and coordination flexibility allows HS to adopt diverse structures, favoured by different metal ions, which is a property that may be utilized for metal ion sequestration in practical applications. Experimental studies on the interaction of a flexible peptide like HS with metal ions are a challenge as speciation may be diverse and the system dynamic. In the span between coordination compounds to metalloproteins, dynamic systems like HS, may yield insight into the underlying mechanisms of metal ion exchange that are necessary to account for transport and distribution of essential trace elements in biological systems. In a more fundamental perspective, also the potential role of metal ions for

protein folding (and misfolding) through transient binding may also be elucidated by interrogating such peptide-metal ion interactions.

Experimental

Materials

The investigated peptide N-acetyl-Ser-Cys-His-Gly-Asp-Gln-Gly-Ser-Asp-Cys-Ser-Ile-NH $_2$ (Ac-SCHGDQGSDCSI-NH $_2$, **HS**) was synthesized, as described earlier. Chemicals and solvents were obtained from Sigma-Aldrich and used without further purification unless otherwise described. The solutions of $\text{Zn}(\text{ClO}_4)_2 \cdot n\text{H}_2\text{O}$, $\text{Hg}(\text{OAc})_2$ (Aldrich) were standardized complexometrically while precise weights of high purity HgCl_2 (Aldrich) was used to prepare metal ion stock solutions. pH-Metric titrations were performed with NaOH (Aldrich) solutions standardized using potassium hydrogen phthalate (Sigma-Aldrich).

Electronic absorption and SRCD measurements

UV-Visible (UV-Vis) spectra were measured on a Shimadzu UV-3600 UV-VIS-NIR spectrophotometer using a cell with 1 cm optical pathlength. Concentration of the ligand was 1.0×10^{-4} M and the metal ion concentration varied between 5.0×10^{-5} and 2.0×10^{-4} M.

The synchrotron radiation CD (SRCD) spectra of the free ligand and the metal complexes were recorded at the SRCD facility at the CD1 beamline on the storage ring ASTRID at the Institute for Storage Ring Facilities (ISA), University of Aarhus, Denmark. All spectra were recorded with 1 nm steps and a dwell time of 2 s per step, using 0.1 mm quartz cells (SUPRASIL, Hellma GmbH, Germany), for the wavelength range of 175–260 nm. The substances were initially dissolved in 1.0 × 10^{-2} M HCl in order to avoid the eventual oxidation process. The pH of the samples ($c_{\rm peptide} = 1.0 \times 10^{-3}$ M) were adjusted by adding the appropriate amount of NaOH solution. From the raw spectra the water baseline was subtracted and spectra were normalized to 1.0×10^{-3} M peptide concentration (to eliminate the effect of dilution).

Perturbed angular correlation of γ-rays

All perturbed angular correlation (PAC) experiments were performed in ISOLDE/CERN with a setup using six BaF $_2$ detectors keeping the samples at a temperature of 1 °C. Production and purification of the radioactive ^{199m}Hg is described in the literature. ⁸¹ The ^{199m}Hg solution (150 μ L) was mixed with solutions of nonradioactive mercury(II) chloride, sodium perchlorate and buffer if needed. TRIS and CAPS buffers were used for adjusting the pH of samples to pH \sim 8.0 and 10, respectively. The peptide was dissolved in 0.01 M perchloric acid and amounts of this stock solution were added to the buffered Hg^{II}-containing solutions to reach the desired final concentrations. Finally, sucrose was added to 55% w/w. The pH of the solutions was adjusted with NaOH and HClO4. In order to avoid contamination of the samples, small volumes were taken for pH

measurements. The temperature dependence of the pH in the TRIS/CAPS-buffered solutions was taken into account and pHvalues were corrected to 1 °C.82 The buffers and the peptide stock solutions were purged with argon. The final volume of the samples was 210 μ L with $c_{peptide} = c_{Hg^{II}} = 8.03 \times 10^{-5}$ M and $c_{\text{buffer}} = 1.60 \times 10^{-2} \text{ M}.$

NMR experiments

¹H NMR measurements were performed on a Bruker Avance DRX 500 spectrometer operating at 500.132 MHz. The spectra were recorded at T = 298 K in a mixture of $H_2O/D_2O = 90:10\%$ v/v and in a few cases in pure D₂O applying the zgpr or zgcppr pulse sequences in order to presaturate the H2O/HDO resonances. In a typical sample the concentration of the peptide was 1.3×10^{-3} M. The chemical shifts were referenced to TSPd4 at 0.0 ppm. Spectra were recorded using a recycle delay of 5 s, an acquisition time of 1.64 s, a spectral width of 5 or 10 kHz and 64-128 scans. In D₂O, the pH* (pH-meter reading uncorrected by the deuterium effect) was adjusted to the desired values with NaOD. The recorded spectra were processed by the ACD/Spectrus Processor software.83

pH-Potentiometric measurements

The protonation and coordination equilibria were investigated in aqueous solutions ($I = 0.1 \text{ M NaClO}_4$, and $T = 298.0 \pm 0.1 \text{ K}$) under argon atmosphere with a special care to avoid the oxidation of the peptide. The potentiometric titrations were carried out by an automatic titration set including a PC controlled Dosimat 665 (Metrohm) autoburette, an Orion 710A precision digital pH-meter equipped with an Orion 8103BNUWP Ross Ultra semi micro pH electrode (165 × 6 mm). Conversion of the relative mV values as pH-meter readings to hydrogen ion concentrations was done as described earlier.⁵⁰ The protonation and complex formation processes were characterized by the following general equilibrium process:

$$pM + qH + rL \rightleftharpoons M_pH_qL_r$$

$$eta_{\mathrm{M}_p\mathrm{H}_q\mathrm{L}_r} = rac{[\mathrm{M}_p\mathrm{H}_q\mathrm{L}_r]}{[\mathrm{M}]^p[\mathrm{H}]^q[\mathrm{L}]^r}$$

where M denotes the metal ion, L the deprotonated ligand molecule, and H the protons. Charges have been omitted for simplicity but can be easily calculated taking into account the composition of the fully protonated dodecapeptide (H₅L⁺). Please note, that this simplified notion is used generally throughout the text and on the figures. The corresponding formation constants $(\beta_{M_pH_qL_r} \equiv \beta_{pqr})$ were calculated using the PSEQUAD computer program. 84 The protonation constants were determined from 3-4 independent titrations (70-80 data points per titration), with a peptide concentration of 1.0×10^{-3} M. The complex formation constants were evaluated from 8 independent titrations (70-80 data points per titration). The applied ratio of Zn^{II} and the ligand was 0.5:1, 1:1 and 2:1 with the $\mathrm{Zn^{II}}$ concentration varied between 5.2 \times 10⁻⁴ and

 2.04×10^{-3} M. Due to precipitation in the presence of metal ion excess in alkaline pH-range, titration data for the Zn^{II}: HS 2:1 samples were evaluated only up to pH 7.1. (The individual fitting parameter of titrations performed with ligand excess dropped by ca. 40% when considering differently protonated bis-ligand species (ZnH₂L₂) besides monomeric ones, and accordingly such species were also included in the final model.

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