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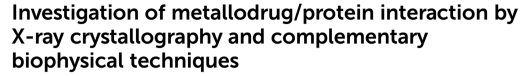
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Giarita Ferraro and Antonello Merlino **

Protein metalation, the process by which a metal compound (or a metal ion) reacts with a protein to produce a metal/protein adduct, is at the basis of many biological events; the knowledge of this process at the atomic level is important in the design and development of new metallodrug-based therapeutic approaches. Recently, single crystal X-ray diffraction experiments have been frequently used to characterize the structures of the adducts formed upon the reaction of Pt, Au, Ru, Rh, Ir, Cu, Mn and V-based drugs with proteins. Although X-ray crystallography is certainly useful to determine the structure of these adducts, the combination with other biophysical techniques provides insights into the system behavior in solution, the reactivity of metal compounds with proteins, fate and stability of the metal/protein adduct and is often helpful for the rationalization of ambiguous or unexpected crystallographic data. Here we describe the results of selected studies carried out in the field of protein metalation, where the structural information achieved by X-ray crystallography has been complemented by data collected using mass spectrometry, vibrational spectroscopy, electron paramagnetic resonance, and computational methods, including density functional theory, docking and molecular dynamics simulations. These works allow us to define the protein metalation process at the molecular level, providing information on the factors responsible for the formation and stability of metal/protein adducts.

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Introduction

The interaction between metal ion/metal complexes and proteins often results in the formation of metal/protein adducts, where metal centers are generally coordinated with specific residue side chains. This process has a major role in biology¹ since it is involved in the correct folding and function of metalloproteins, which are a large fraction of proteins within the cells.² Protein metalation significantly influences the absorption, transportation and storage of metallodrugs within the body and has a major role in the design of synthetic artificial metalloenzymes3 and in the development of protein-based metallodrug delivery systems. The interplay between metal compounds and proteins also contributes to the assessment of adverse reactions associated with the use of metal-based anticancer treatments. Indeed, the exploration of protein metalation induced by anticancer metal-based agents enables the elucidation of fundamental aspects underlying the occurrence of side effects and activation/inactivation mechanisms of these

Department of Chemical Sciences, University of Naples Federico II, Complesso Universitario di Monte Sant'Angelo, via Cinthia, 21, 80126 Naples, Italy. E-mail: antonello.merlino@unina.it

compounds, which are frequently pro-drugs. The formation of metal/protein adducts also alters the catalytic properties of various enzymes.4 Therefore, a detailed understanding of the mechanisms that form the basis of the recognition of metal compounds by proteins has become one of the most stimulating goals in recent years.5-7 X-ray diffraction (XRD) experiments are routinely used for the determination of structures of adducts formed upon the reaction of metal compounds with proteins.8-12 In an XRD experiment, X-rays interact with the electrons of atoms in a crystal, providing an experimental dataset that is related to the electron density (e.d.) map, which allows localizing of electrons of the crystal asymmetric unit and thus determination of the molecular structure. In protein X-ray crystallography, the e.d. map shows a time- and spaceaverage of the e.d. of the atoms in the crystal. When a protein region is highly flexible, it is associated with a poor e.d. map and is difficult to model. Thus, it is sometimes omitted by the final protein structure model. Similarly, molecules that are disordered or present in crystals with low occupancy, i.e. only in a fraction of the unit cell, are associated with weak e.d. and are often difficult to identify. This also happens in the structures of metal/protein adducts. The position of the metal is generally well defined, but metal ligands cannot be unambiguously modelled. Furthermore, in the formation of metal/protein

adducts, the molecular species that bind the protein are not those reacting with the macromolecules in solution or those used to treat protein crystals. Indeed, an important role in the definition of the final metal/protein adduct is played by the reaction that occurs in solution (ligand exchange, hydrolysis, speciation, reduction/oxidation, etc.) or by the reaction occurring upon the metal-containing fragment binding to the protein. The combination of X-ray crystallography with other biophysical techniques helps to define details of the molecular processes that occur before or upon the formation of the metal/protein adduct. In this context, spectroscopic analysis, UV-vis absorption spectroscopy and nuclear magnetic resonance (NMR), for example, are fundamental techniques to study the metal compound behavior in solution alone and in the presence of a protein. On the other hand, circular dichroism or fluorescence can be helpful in unveiling protein secondary and tertiary structural alterations induced by the metal complex binding. Interesting information has also been obtained by using small angle X-ray scattering (SAXS)^{13,14} or perturbed angular correlation spectroscopy. 15-17

Here, selected examples of the combined use of XRD and specific biophysical techniques (mass spectrometry, IR and Raman spectroscopy, electron paramagnetic resonance, dynamic light scattering (DLS) and computational methods, including density functional theory, docking and molecular dynamics simulations) in the definition of the protein metalation process with several different metallodrugs are discussed. The selected works reflect the authors' personal perspective; thus, the paper does not pretend to be a comprehensive study.

Protein metalation studied by a combination of XRD and mass spectrometry

XRD data provide experimental evidence on the metal coordinates and on the protein residues involved in metal recognition. However, sometimes the definition of the e.d. map does not allow us to precisely identify the exact metal-containing fragments that bind a protein. 18 In these cases, mass spectrometry techniques can be used to characterize the metal/ protein adducts formed upon the reaction of metal compounds with proteins. 19,20 Electrospray ionization mass spectrometry (ESI-MS) is one of the most widely used mass spectrometry techniques. It is based on the ionization into the gas phase of polar molecules dissolved in a liquid.²¹ The samples are transferred to the ion source either directly (through a pump²² or a flow-injection unit²³) or indirectly (via a separation system, such as liquid chromatography, LC, high-performance liquid chromatography, HPLC²⁴ or capillary electrophoresis).²⁵ Mass spectra report the intensity values as a function of corresponding mass-to-charge ratios (m/z). ESI-MS is a so-called 'soft ionization' technique, since it produces very little fragmentation, thus allowing the determination of molecular masses and chemical structures of metal/protein adducts, the metal-containing fragment/protein stoichiometry, and the protein binding sites and their number, even preserving non-covalent interactions of metal complexes with proteins.²⁶ ESI-MS is fast, sensitive and specific. It can investigate

the metal compound/protein interaction over time and at different metal to protein molar ratios and under different experimental conditions, 27 also defining the effect of protein post-translation modifications on the metal compound binding.28 The main disadvantages of this technique are the difficulty in ionizing metal adducts of large proteins or of proteins that tend to aggregate forming higher oligomers, causing unclear or multiple mass peaks, and their sensitivity to pH.²⁹ Examples of studies combining ESI-MS and X-ray crystallography to characterize the formation of metal/protein adducts include different metals (Pt, Ru, Au, Ir, and V, among others, see for example complexes shown in Fig. 1) and diverse proteins (examples are shown in Fig. 2). This approach has been already summarized in previous works. 27,30,31 Cisplatin (Fig. 1) binding to proteins has been studied both from the crystallographic and mass spectrometry points of view in a number of cases. It has been shown that Pt2+, [Pt(NH3)2]2+, and [Pt $(NH_3)_2Cl$ ⁺ or the product of its hydrolysis $[Pt(NH_3)_2(H_2O)]^{2+}$ bind Nδ1 or Nε2 atoms of His15 of hen egg white lysozyme (HEWL, Fig. 2).32 Pt binding sites have been found close to Met29 or both Met29 and Gln28,33 and His105 and His119 side chains of bovine pancreatic ribonuclease (RNase A, 33,34 Fig. 2). Cys12 and Cys15 have been identified as Pt binding sites in Atox-135 (Fig. 2), while Met65 and Glu61 are the residues involved in the metal recognition by cytochrome c³⁶ (Fig. 2). Cisplatin fragments bind His68 of ubiquitin²⁹ (Fig. 2), His105, His128, His146, His247, His288, Met298, Met329, His305, His338, His440 + Lys436, and Met548 of human serum albumin³⁷⁻³⁹ (HSA, Fig. 2).

It has been shown that the same metal containing fragments can bind β-lactoglobulin, 40 transferrin 41 and angiogenin⁴² (Fig. 2).

Using the same approach, the interaction of some of these proteins with carboplatin²⁸ and oxaliplatin (Fig. 1)³⁸⁻⁴¹ has also been characterized. In the latter case, $[Pt(dach)(H_2O)]^{2+}$, $[Pt(dach)]^{2+}$ (dach = 1,2-diaminocyclohexane) or the intact oxaliplatin binds proteins^{43–46} close to the side chains of Asp, Met, His or Cys residues, in metal binding sites that are in some cases distinct from those of cisplatin. Using the combined crystallographic/ESI-MS approach it has also been possible to analyze in detail the protein binding of other potential anticancer Pt-based drugs, like picoplatin (Fig. 1),47 the diiodido analogue of cisplatin cis-PtI₂(NH₃)₂³³ (Fig. 1), trans-Pt derivatives (like trans-(dimethylamino) (methylamino) dichloroplatinum(II), [t-PtCl2(dma)(ma)], Fig. 1),48 a tetranuclear Ptthiosemicarbazone complex49 and Pt-compounds containing O,S-bidentate ligands (like compound **9b** in Fig. 1).⁵⁰

Binding of Ru-, 51,52 Au-, 53,54 V-, and homo (Ru-Ru⁵⁵ and Rh-Rh, 56-58 Fig. 1) and hetero-bimetallic-(like Pt-Au⁵⁹ and [Pt $(\mu\text{-NHC}(CH_3)O)_2ClAs(OH)_2]$, AP-1^{60,61} in Fig. 1) based drugs to HEWL and RNase A^{62,63} has also been investigated. In the case of the antimetastatic metallodrug imidazolium (Im) trans-[tetrachlorido(S-dimethyl sulfoxide)(1H-imidazole)ruthenate(III)] (NAMI-A, Fig. 1) and its derivative AziRu (Fig. 1), 64 a complete hydrolysis associated with the release of all the metal ligands has been observed in the reaction with HEWL,51 carbonic

Fig. 1 Chemical structures of metallodrugs and selected metal-based drug candidates.

anhydrase (hCAII),65 and human H-chain ferritin (h-H-Ft)66 (Fig. 2). The HEWL ruthenation mechanism by NAMI-A has been studied in detail by Papakyriakou and coworkers, who solved the X-ray structures of the protein with NAMI-A at

different soaking times.⁶⁷ The structures highlight a series of events that ultimately lead to the final "ruthenated" protein: NAMI-A non-covalently binds the protein (after 1.5 h of soaking, Fig. 3A), then it exchanges all Ru ligands except for

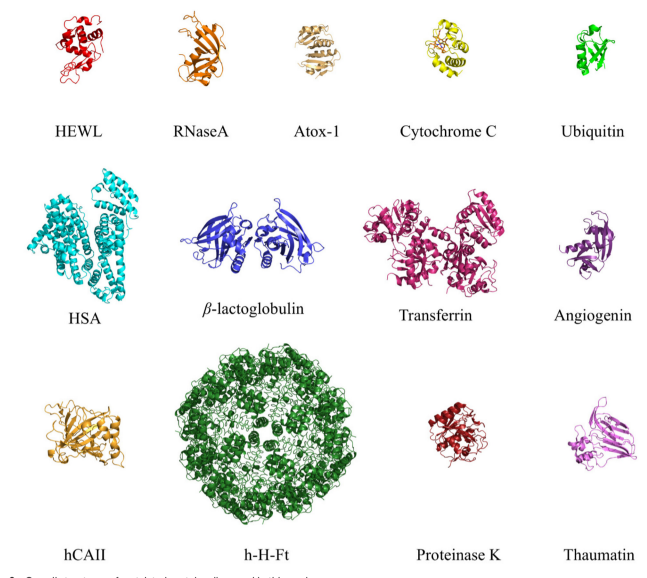


Fig. 2 Overall structures of metalated proteins discussed in this work.

Im (8 h and 26 h of soaking, Fig. 3B and C) and finally the aquated Ru ion coordinates to His15 and Arg14 (98 h) (Fig. 3D). More recently, a comparative structural analysis of the HEWL adducts with two RuIII complexes [HIsq][trans-RuCl₄(dmso)(Isq)] (1, Fig. 1) and [H₂Ind][trans-RuCl₄(dmso) (HInd)] (2, Fig. 1) (where HInd - indazole and Isq - isoquinoline are analogues of NAMI-A), together with in-solution studies, has demonstrated that the hydrolytic release of the N-heterocyclic ligand is one of the main factors in determining the interaction of Ru-containing fragments to proteins.⁶⁸ From these works it emerges that anticancer Ru agents are often prodrugs, which have to be activated before they can react with the final protein target.⁶⁹ In the case of Ru compounds the activation could be a simple aquation reaction.

The selectivity for different binding sites on HEWL of a series of isostructural N-heterocyclic carbene (NHC) compounds with the formula $[M(L)(dmb/dbb)Cl_2]$ (M = Ru/Os/Rh/Ds/R

Ir; L = η^6 -p-cymene [cym], η^5 -pentamethylcyclopentadienyl [Cp*]; dmb = 1,3-dimethylbenzimidazol-2-ylidene; dbb = 1,3dibenzylbenzimidazol-2-ylidene, Fig. 1) was investigated by Sullivan and co-workers⁷⁰ choosing this combined approach. The X-ray structures of the adducts highlighted a diverse behavior of the analyzed metal complexes. Indeed, the Ru and Rh compounds showed a tendency to bind the surface exposed His15 upon loss of the p-cymene or the NHC ligands, while the Ir and Os derivatives were identified close to the electronegative peptidoglycan-binding pocket HEWL, showing a preferential binding for Asn103 and Asp101. In these cases, the binding occurred upon replacement of the chloride ligand with the side chain of the protein residues.

Gold-based drugs also must be activated before reaching their final targets. Crystallographic and ESI-MS studies carried out to unveil the reactivity of anticancer AuIII-based drugs with proteins have suggested that for these compounds the protein

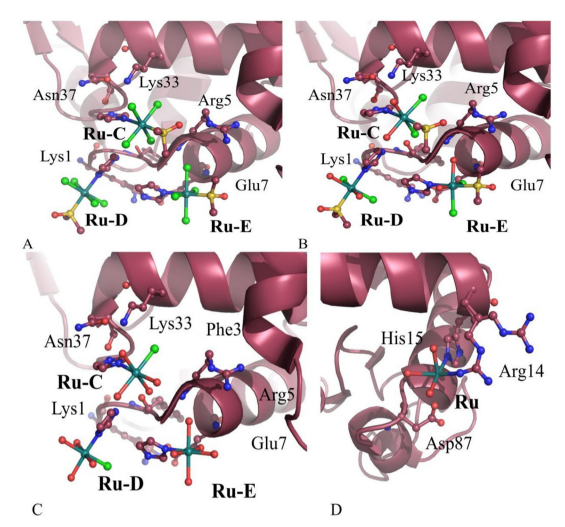


Fig. 3 NAMI-A binding sites in the X-ray structures of a NAMI-A/HEWL system solved after 1.5 h (a), 8.0 h (b), 26 h (c) and 98 h (d) of soaking. Comparison between the three structures show that NAMI-A can bind HEWL non-covalently (see Ru-C and Ru-D molecules in panel A) and covalently, upon the release of the Im ligand (see the NAMI-A moiety, denoted as Ru in panel D, bound to side chains of Arg14 and His15). Data also indicate that chloride ligands and DMSO are progressively replaced by solvent molecules (see panels B and C) and that Im is retained up to 26 h.

binding occurs after or concomitantly to metal compound degradation and metal center reduction, with the coordination of Au^I to His or Met side chains. 54,71 Indeed, in the reaction of Auoxo6 ($[Au_2(bipy^{2Me})_2(\mu-O)_2][PF_6]_2$, where $bipy^{2Me} = 6,6'$ dimethyl-2,2'-bipyridine, Fig. 1), Au₂phen ([Au₂(phen^{2Me})₂(μ- O_2 [PF₆]₂, where phen^{2Me} = 2,9-dimethyl-1,10- phenanthroline, Fig. 1) and AuSac2 ($[NH_4][Au(Sac)_2]$, where Sac = deprotonated saccharinate ligand, Fig. 1) with HEWL, only one Au^I atom bound to His15 has been observed.54 An additional structure of the HEWL adduct with AuSac2, obtained under a different experimental condition, has revealed two Au^I ions close to the side chain of His15, and an additional gold center close to the side chain of Met105, in a buried hydrophobic pocket. These crystallographic data are in good agreement with ESI-MS results indicating that the protein can bind up to three Au^I ions.⁷¹ Degradation of the Au^{III} compound and binding to Au^I to a Gln side chain has been observed in the reaction of HEWL with Aubipy^c ([(bipy^{dmb}-H)Au(OH)][PF₆],⁵³ Fig. 1), in the reactivity of Auoxo6 with RNase A62 and of the medicinal AuIII dithiocarbamato complex AuL₁₂ (Fig. 1) with bovine serum albumin.⁷² ESI-MS/XRD data have been collected also for adducts formed by proteins with vanadium compounds. ESI-MS recorded on the system containing bis(maltolato)oxidovanadium(IV) ([VIVO(malt)2], Fig. 1) and HEWL with a 2:1 metal to protein molar ratio at pH 5.0 and pH 6.5 showed the formation of HEWL-VO-malt adducts, with [VIVO]2+, [VIVO $(\text{malt})^{\dagger}$ and $[V^{IV}O(\text{malt})_2]$ that bind the protein. These results agree well with crystallographic data showing that in the HEWL-VO-malt adduct, cis-[VIVO(malt)2(H2O)] (Fig. 4) and $[V^{IV}O(malt)(H_2O)_3]^+$ bind the protein non-covalently, while $[V^{IV}O(H_2O)_{3-4}]^{2+}$, cis- $[V^{IV}O(malt)_2]$ and other V-containing fragments are coordinated to the side chains of Glu35, Asp48, Asn65, Asp87, and Asp119 and to the C-terminal carboxylate. 73 Both covalent and non-covalent binding modes have also been predicted by ESI-MS and then verified by X-ray crystallography in the adduct formed when $[V^{IV}O(8-HQ)_2]^{74}$ (8-HQ = 8-hydroxy-

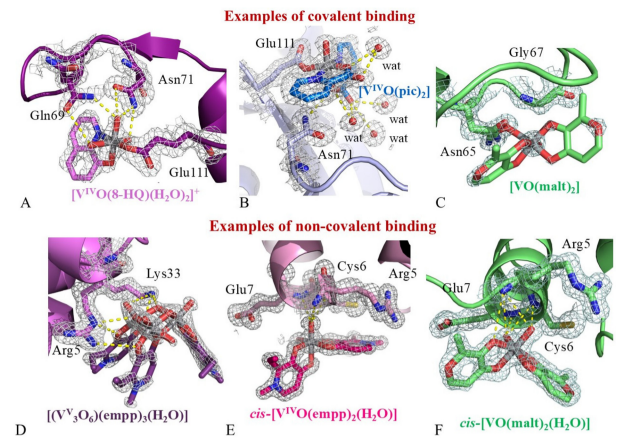


Fig. 4 Examples of covalent and non-covalent binding of V compounds to RNase A (panels A and B) and HEWL (panels C-F).

quinoline) and [V^{IV}O(empp)₂] (where Hempp is 1-methyl-2ethyl-3-hydroxy-4(1*H*)-pyridinone) react with the protein⁷⁵ (Fig. 4).

A similar approach has been used to characterize the interaction of the V^{IV}O-picolinato complex ([V^{IV}O(pic)₂(H₂O)], pic = picolinate) with RNase A (Fig. 4): the structure of the adduct formed when the V complex reacts with the bovine enzyme at acidic pH revealed that [VIVO(pic)2] binds the Glu111 side chain by replacing the water equatorial ligand. ESI-MS confirms this result.76

Metal/protein adducts ionized by electrospray can also be monitored by ion mobility mass spectrometry (IM-MS). In IM-MS, the ions are separated thanks to their different mobility through an inert gas: compact ions move faster than ions with the same m/z values but less compact.⁷⁷ IM-MS can give insights into the effect of metalation on protein folding, stability and dynamics and can distinguish between different protein conformations, which might be indistinguishable by traditional mass spectrometry techniques. The main disadvantage of this technique is its limited availability, since it is not diffuse as the other mass spectrometry techniques. Other disadvantages are related to limits in treating data from large proteins, that often produce peak overlap or broadening.

IM-MS has been used to study the binding of [Ru(cym)Cl₂]₂ (Fig. 1) to HEWL.⁷⁸ X-ray crystallography revealed that [Ru(cym)Cl₂] binds the side chains of His15 and Asp101, after cleavage of the dimeric structure. IM-MS data indicated that the number of higher charge state HEWL ions is increased by "ruthenation", suggesting that metal coordination to HEWL destabilizes the protein structure. Additionally, ion mobilograms indicated possible protein unfolding and a compaction of the unfolded state due to the metal-containing fragment binding. The reduced stability of the Ru/HEWL adduct when compared to metal-free HEWL is confirmed by differential scanning calorimetry (DSC) experiments.⁷¹ Compaction upon metalation was observed in the metalation of other proteins.79-81

Recently, the combined XRD/mass spectrometry approach has also been used for systems of large dimensions. ESI-MS has been used, in combination with proteolytic assays and site directed mutagenesis studies, to identify h-H-Ft residues that bind gold atoms from sodium aurothiomalate, an FDAapproved anti-arthritic drug, in the Au-h-H-Ft adduct formed upon the reaction of the protein with the drug. 82 In this study, the incubation of sodium aurothiomalate with h-H-Ft under near-physiological solution conditions led to the formation of the Au/h-H-Ft adduct with enhanced biological activity when compared to the gold-compound alone. In Au/h-H-Ft, a cluster of gold atoms are bound to Cys90 and Cys102. These residues have been identified as gold binding sites in the adduct formed by h-H-Ft with auranofin. 83,84

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There are other mass spectrometry techniques that have been frequently used to characterize the formation of metal/ protein adducts. As an example, the protein interaction of Pt complexes has been characterized by electrospray ionization Fourier transform ion cyclotron resonance mass spectrometry (ESI-FT-ICR-MS).⁸⁵ Collision-induced dissociation (CID), higher energy collision-induced dissociation (HCD), electron transfer dissociation (ETD) or electron capture dissociation (ECD) have also been used to elucidate specific residues involved in metal compound binding to proteins.86,87

Inductively coupled plasma (ICP)-MS is an element-specific analytical method, which allows quantitative determination of metallodrugs in biological samples through the evaluation of their metal content. 88 The technique features high specificity, in the ng L⁻¹ concentration range, and can be used to determine the metal to protein molar ratio in a metal/protein adduct and its stability over time. 20 Interferences from isotopic patterns and problems with the calibration and standardization of the instrument, which could lead to limited sensing ability towards specific metals, are the main disadvantages of this technique. ICP-MS measurements have been employed to define the presence of Ru within crystals of binuclear trithiolato-bridged arene ruthenium complex (diruthenium-1)/horse spleen L-chain ferritin (hs-L-Ft) nanocages obtained using a protein sample that was disassembled and reassembled in the presence of the anticancer compound diruthenium-1 (DiRu-1 in Fig. 1)89 and within crystals of the adduct of indazolium trans-[tetrachlorobis(1H-indazole)ruthenate(III)] (KP1019, Fig. 1) with HSA.⁹⁰ In the structure of diruthenium-1/hs-L-Ft, Ru atoms were not observed in the e.d. maps, thus suggesting that the anticancer compound was in the bulk within the cage.⁸⁹ In the structure of the KP1019/HSA adduct, Ru atoms were coordinated to His146 and His242. Solvent molecules complete the Ru octahedral coordination sphere, suggesting dissociation of both indazole ligands from the metal center.⁹⁰

ICP-MS measurements have been also used to quantify the amount of Ru in the bionanocomposite formed when recombinant L-chain apo-ferritin from horse liver (hl-L-Ft) was treated with [Ru(CO)₃Cl₂]₂ (CORM-2, Fig. 1)⁹¹ and the amount of Au upon encapsulation of Auoxo3 ($[Au_2(bipy^{Me})_2(\mu-O)_2][PF_6]_2$, bipy^{Me} = 6-methyl-2,2'-bipyridine, Fig. 1),⁹² Auoxo4 $([Au_2(bipy^{nP})_2(\mu-O)_2][PF_6]_2$, $bipy^{nP} = 6$ -neo-pentyl-2,2'-bipyridine, Fig. 1), and Au₂phen⁹³ within hs-L-Ft nanocages. A similar analysis has been also carried out to evaluate the amount of Pt within the cage of the same protein⁹⁴ and the amount of Pt and As within AP-1/h-H-Ft.95 The structure of the RuCO/hl-L-Ft nanocomposite shows Ru atoms coordinated to the side chains of His114, Glu130 and His132;91 the structures of the Au/hs-L-Ft adducts show binding of Au centers to Cys48, His49, His114, His114 and Cys126, Cys126, His132, His147. Pt binds the side chains of His49, His114 and His132.94 AP-1 binds the side chain of His49 of hs-L-Ft.⁹⁵

Notably, ICP-MS has also been used to characterize the Au/ apo-R168H/L169C-hl-L-Ft system where gold nanoclusters are formed.^{96,97} Ten⁹⁷ and 8-12⁹⁶ atoms of Au in clusters were found (see for example Fig. 5). Using the same approach, Fujita et al. 98 studied the role of the R52C mutation in hl-L-Ft when it reacted with Mn(CO)5Br to produce a ferritin-based nanocomposite containing a manganese-carbonyl complex. When the nanocomposite is designed starting from the wildtype protein, only a few Mn atoms could be detected within the cage (about 3 atoms), while the mutated Ft could accumulate about 48 Mn atoms.

A novel approach, developed by Timerbaev et al., 99 matching capillary electrophoresis with ICP-MS enabled a deep characterization of the affinity of cisplatin and two of its novel analogues, 2-ABDP, (SP-4-2)-bis((R)-(-)-2-aminobutanol)dichloroplatinum(II), and 4-ABDP, (SP-4-2)-bis(4-aminobutanol) dichloroplatinum(II) (Fig. 1) for HSA. The coupling of these two techniques turned out to be a useful tool to measure the rate constant of the binding reactions and to determine the number of drug molecules attached to the protein. The binding of cisplatin to HSA was a pseudo-first order reaction. When the protein is incubated in the presence of a huge excess of metallodrug, it can bind up to 10 Pt atoms per protein chain.

ICP-MS is suitable to be combined with different types of techniques. Indeed, the combination of the separation capacity of SDS PAGE with the specificity of ICP-MS has been employed to characterize oligomers of different orders formed upon interaction of HSA with cisplatin. 100 It was found that increasing amounts of cisplatin push the serum albumin to form oligomers bigger than its dimeric form, which is generally the most abundant. In fact, as the reaction proceeds, dimeric HSA is drained from the reaction environment leaving the monomer as the most representative species.

Alternatively, the amount of metals (Pd, 101,102 Rh, 103 Ru, 66 Au,83 Pt104) within Ft nanocages with solved X-ray structures has been evaluated using inductively coupled plasma-optical emission spectrometry (ICP-OES). ICP-OES can detect heavy metal concentrations, even in the range of parts per billion (ppb) or parts per trillion (ppt), with high accuracy. It can analyze a wide spectrum of metals, even simultaneously, but it is a rather expensive technique. Although the ICP-OES is generally resistant to interference, the chemical composition of the sample matrix can affect the results, requiring the use of internal standards or corrections.

Matrix-assisted laser desorption/ionization-mass spectrometry (MALDI-MS) is an ionization technique that is based on the use of a laser energy-absorbing matrix to create ions from large molecules, with minimal fragmentation. 105 MALDI-MS is similar to ESI-MS since both are 'soft ionization' techniques, but it typically produces far fewer multi-charged ions. MALDI-MS can be used for samples of metal/protein adducts treated with denaturants, like urea, or with reducing (like dithiothreitol) and alkylating (like iodoacetamide) agents or using digested samples.

Recently, MALDI-MS combined with chromatographic separation techniques has served for studying an unusual reaction product obtained when HEWL crystals have been treated with [V^{IV}O(malt)₂] in 1.1 M NaCl and 0.1 M CH₃COONa at pH 4.0. 106 In this work, crystals of the adduct formed by the V

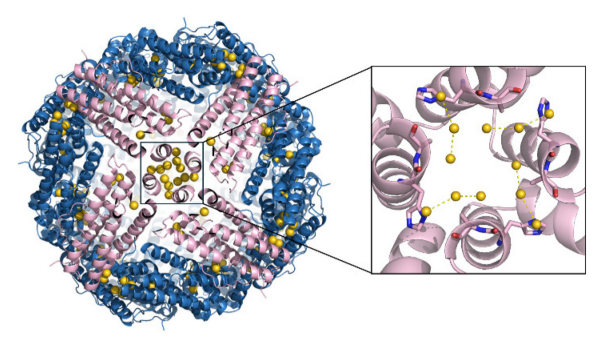


Fig. 5 A gold nanocluster found within a hl-L-Ft nanocage (PDB code 7VIT). Gold atoms are in yellow, and protein chains involved in the recognition of gold atoms in pink.

compound with the protein have been dissolved and analyzed by sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE). The gel showed that HEWL formed an SDS-resistant dimer in the presence of [V^{IV}O(malt)₂] (Fig. 6A), together with a metalated monomeric species. Metalated monomeric

and dimeric protein forms were characterized and found to be in agreement with the crystallographic model: [V^{IV}O(malt)₂] degrades within HEWL crystals, the malt ring is broken, and two V centers are bound to a modified N-terminal Lys side chain, with one of the two metal centers that also acts as a

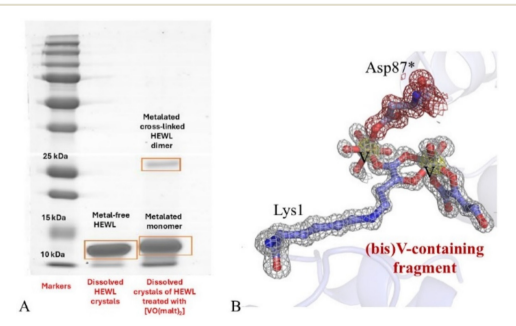


Fig. 6 (A) SDS-PAGE of dissolved crystals of HEWL (lane 2) and its adduct with [VINO(malt)₂] (lane 3). Markers are in lane 1. The gel indicates that upon incubation of HEWL crystals with the V compound, a metalated cross-linked HEWL dimer is formed. (B) Modification of Lys1 in crystals of HEWL grown in 1.1 M NaCl, 0.1 M sodium acetate at pH 4.0 and treated with $[V^{IV}O(malt)_2]$ (PDB code 9FMY). A $2F_0 - F_0$ e.d. map is reported at the 1.0σ level in grey; anomalous difference e.d. map is colored in yellow and reported at the 3.0σ level. *Indicates atoms from a symmetry related molecule. These atoms are in grey; their e.d. map at the 1.0σ level is in red.

crosslinker of two protein molecules within the crystal (Fig. 6B). Gel and mass spectrometry have also been used to test the stability of metal–protein bonds in five platinated proteins by Moreno-Gordaliza et al. ¹⁰⁷

A MALDI-TOF-MS-X-ray crystallography combined approach was used to characterize the adduct formed upon reaction of a series of Cu(II) compounds with 6-methyl-2-formylpyridine-4*N*-substituted thiosemicarbazone compounds (Fig. 1) with HSA previously complexed with palmitic acid (PA). The presence of PA in the IB subdomain of the protein strengthens the affinity of copper for HSA preventing its early release.

Protein metalation studied by dynamic light scattering

The formation of peptide and protein dimers or higher oligomers or a change in the overall conformation of large protein systems upon metalation has been observed in many cases. ^{109–111} To study the formation of metal-based cross-linked oligomeric protein states, SDS-PAGE or size exclusion chromatography can be used.

Cross-linking mass spectrometry (XL-MS) has been widely used in the analysis of metal-induced cross-linked products. Nie *et al.* exploited this methodology to elucidate structural differences between the allosteric enzymes, glycogen phosphorylases (GP) b and a. 112 It is a two-stage analysis which couples a cross-linking experiment, in this case triggered by cisplatin, to an MS experiment. This approach has revealed itself to be worthwhile particularly for the analyses of the N-terminus and the helix interfaces, which were the uncrystallized regions.

Another technique that has been used for this type of characterization is dynamic light scattering (DLS). DLS studies carried out on fatty acid-free and bound forms of HSA, in the absence and in the presence of cisplatin have shown that the tendency of the protein to aggregate increases in the presence of the Pt-based drug, providing a precise estimate of the size of the aggregates.³⁷ DLS analysis has also confirmed the formation of large aggregates of caspase cleaved prostate apoptosis response-4 protein (cl-Par-4) in the presence of cisplatin.¹¹³

Protein metalation studied by X-ray crystallography complemented with spectroscopic techniques

Infrared (IR) spectroscopy and Raman spectroscopy are two variants of vibrational spectroscopy, which provide information on the vibrational transitions which strictly depend on chemical composition, structure, symmetry, electronic environment and bonding of molecules. Infrared spectroscopy has been combined with crystallographic data to characterize the formation of the $[Re(CO)_3(H_2O)_2]^+/HEWL$ adduct¹¹⁴ upon the reaction of $[Re(CO)_3(H_2O)_3]^+$ with the protein and to monitor a time-resolved series of HEWL-fac- $[Re(CO)_3]^+$ -imidazole (HEWL-Re-Im) crystals.¹¹⁵ For these systems, carbonyl stretching frequencies can be used as spectroscopic probes to determine metal coordination.

The presence of $[Re(CO)_3]^+$ within HEWL crystals in the $[Re(CO)_3(H_2O)_2]^+$ /HEWL adduct was confirmed by $\nu(CO)$ stretching bands at 1890, 2001 and 2013 cm⁻¹. The appearance of the

2001 and 2013 cm⁻¹ bands suggests the presence of multiple rotamers of the [Re(CO)₃]⁺ adduct.¹¹⁴

In the IR spectra of HEWL-Re-Im crystals ν (CO) stretching bands (2017, 1896 cm⁻¹) started to appear from week 1, while a slight shoulder appeared at 1896 cm⁻¹ from week 15. Longer soaking (up to 67 weeks) was associated with an increase in the wavenumber (ν (CO) = 2019, 1899, 1883 (shoulder) cm⁻¹), suggesting a continuation of movement of the Re complexes within the protein structure (Fig. 7). 115 Along the same line, carbonyl stretching frequencies in ATR-IR spectra have been used to monitor the presence of [Mn(CO)₃] within HEWL microcrystals soaked with [Mn(CO)₃(acetone)₃](CF₃SO₃) for 4-5 days in the dark 116 and the presence of CORM-2 derivatives in the RuCO/hl-L-Ft adduct formed when hl-L-Ft was treated with the Ru carbonyl complex.⁹¹ Fujita et al.⁹⁸ employed ATR-FTIR spectroscopy to confirm the presence of CO ligands in the ferritin composite containing a manganese-carbonyl complex designed reacting the R52C-hl-L-Ft mutant with Mn(CO)5Br previously described. In the crystal structure there was no electron density map assignable to the CO ligands around the metal center. However, the ATR-FTIR spectrum of MnCO·R52C-hl-L-Ft showed the peaks at 2028, 2011, and 1917 cm⁻¹ in the CO-stretching vibrational region which were assigned to a fac-Mn(CO)3 structure. Similarly, for the $[Mn(CO)_3]/HEWL$ system, the $\nu(CO)$ stretching bands at 2030 cm⁻¹ and 1938 cm⁻¹ have been noted. In the RuCO/hl-L-Ft adduct, the observed bands were at 2038 and 1956 cm⁻¹.

Raman spectroscopy uses the inelastic diffusion of monochromatic incident radiation to gain knowledge about molecular vibrations of the molecules of the sample. In confocal Raman microscopy, an optical microscope focuses the laser beam onto the sample. ¹¹⁷ The technique is very efficient in the analysis of the distribution of chemical moieties within several matrices, including protein crystals. ¹¹⁸ For example, Raman microspectroscopy has been used to evaluate the kinetics of the binding of platinochloride ions within HEWL single crystals treated with a tetrachloroplatinate salt (K₂PtCl₄).

Raman spectra were collected at different times from the HEWL crystal during 14 h of soaking. The spectra revealed growing characteristic Pt–Cl stretching bands in the 305–330 cm⁻¹ region (Fig. 8). ¹¹⁹ The crystal structure then confirmed the presence of four anomalous map signals at the surface of the protein, attributed to Pt atoms.

A similar approach has been used to study the release or the persistence of CO bound to $[IrCl_5CO]^{2-}$ (Fig. 1) within HEWL or RNase A crystals.

Raman spectra have been collected during [IrCl₅CO]²⁻ soaking into HEWL crystals (647 nm line), using spectra of Cs₂[IrCl₅CO] powder, of the protein and of the crystallization conditions as references.¹²⁰ The diffusion of the Ir complex into HEWL crystals has been followed by observing the appearance of low frequency bands (around 300 cm⁻¹) in the spectra of the crystal, during the first hours of soaking. As in the case of IR spectra, Raman carbonyl frequencies can be conveniently used as spectroscopic probes for metal compound binding to the protein. In the spectrum of Cs₂[IrCl₅CO], one signal at

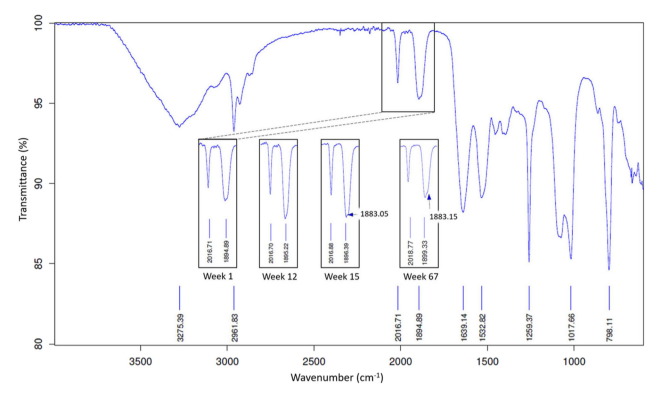


Fig. 7 IR spectrum of HEWL-Re-Im crystals. ν (CO) stretching bands have been highlighted. Insets of this region for crystals at weeks 1, 12, 15 and 67 after crystallization have been added to the spectrum. Reproduced from ref. 115 with permission from IUCr Journals, copyright 2024.

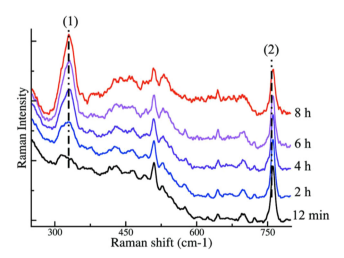


Fig. 8 Raman spectra of a HEWL crystal soaked in 10 mM solution of K₂PtCl₄ for 12 min (dark), 2 h (blue), 4 h (violet), 6 h (pink) and 8 h (red). The Pt-Cl stretching band at 330 cm⁻¹ ((1) in the spectrum) and Trp ring breathing band at 760 cm⁻¹ ((2) in the spectrum) are shown. Reproduced from ref. 118 with permission from IUCr Journals, copyright 2007.

about 2060 cm⁻¹ is found. When [IrCl₅CO]²⁻ soaks into HEWL crystals, two new bands (2173 and 2076 cm⁻¹) and a new shoulder (at 2122 cm⁻¹) appear. The two bands have been attributed to free CO trapped into the protein crystal and to a mixture of unbound $[IrCl_5CO]^{2-}$ and $[IrCl_4CO(H_2O)]^{-}$ ions. The shoulder at 2122 cm⁻¹ has been attributed to the metal containing fragment coordinated to the His15 imidazole.

A similar analysis has been carried out monitoring the diffusion of [IrCl₅CO]²⁻ into RNase A crystals. In this case, the entrance into the crystal of the Ir compound was verified by the appearance of a band at 2073 cm⁻¹, attributed to CO bound to the metal center. 121

Hunter et al. 122 coupled NMR spectroscopy to X-ray crystallography to obtain structural information on the interaction of the anti-HIV drugs Cu-cyclam and Cu2-xylyl-bicyclam (Fig. 1) with HEWL, taken as the model protein for CXCR4 coreceptor, which is the entrance way for HIV into cells. The X-ray structures revealed that Cu-cyclam and Cu2-xylyl-bicyclam protein recognition occurs through polar and non-polar interactions, involving H bonding between the cyclam ring and protein carboxylate groups, as already reported, 123 and hydrophobic interactions between the cyclam ring and the indole ring of tryptophan. The latter interaction was confirmed by the decrease of line-broadening of tryptophan indole NH resonances in the ¹H NMR spectra induced by Cu-cyclam.

Protein metalation studied by a combination of X-ray crystallography and electron paramagnetic resonance

The impact of other biophysical techniques to support X-ray crystallography in the definition of the protein metalation process is illustrated using EPR in achieving the definition of the interaction of VIV, RuIII and CuII compounds with HEWL. EPR can be in principle used to elucidate electronic structures

of transition metal complexes with one or more unpaired electrons. 124,125

The EPR spectra of a monometallic paramagnetic compound with one unpaired electron are characterized by two spin Hamiltonian parameters: the g factor and the hyperfine coupling (HFC) constant between the unpaired electron and the nucleus ($g_{\rm iso}$ and $A_{\rm iso}$ in an isotropic spectrum, and g_x , g_y , g_z and A_x , A_y , A_z in an anisotropic spectrum). When the crystal field has axial symmetry, the splitting between two spin energy states is described by g_\perp and g_\parallel . $g_\perp = g_x = g_y$, $g_\parallel = g_z$. The g and A tensors provide information on the metal chemical environment.

 V^{IV} (d¹ configuration), Ru^{III} (d⁵ configuration) and Cu^{II} are paramagnetic and thus can be monitored by EPR, while V^{V} , Ru^{II} and Cu^{II} are diamagnetic and thus do not exhibit EPR signals. For this reason, the reduction of Ru^{III} and Cu^{II} and oxidation of V^{IV} can be detected by the loss of EPR signal intensity.

EPR data have been collected for several $V^{\rm IV}$ compounds in the presence of proteins (see for example, ref. 127 and 128) but only in a few cases these experiments have been combined with crystallographic data.

Crystal structure of the adducts formed by $V^{IV}OSO_4$ and phen/bipy (L) with HEWL when protein crystals have been treated with these molecules showed binding of the $V^{IV}OL^{2+}$ moiety to the protein. EPR data collected on these systems confirmed the presence of the adducts with spin Hamiltonian parameters in agreement with equatorial (Nbipy/phen, Nbipy/phen; COO-Asp; COAsn) coordination. ¹²⁹

EPR experiments also verified the formation of adducts between $[V^{IV}O(pic)_2]$ and $[V^{IV}O(8\text{-HQ})_2]$ with RNase A. ¹³⁰ For these systems, crystallographic data have been collected at acidic pH, indicating the binding of V to Glu111. EPR data indicated the binding of V centers to carboxylate groups, stemming from Asp or Glu residues, at acidic pH and of an imidazole nitrogen from His residues at physiological pH, ^{76,130} (Fig. 9). Spectroscopic data also suggested that His-N or Ser/Thr-O $^-$ are possible metal binding sites at basic pH (Fig. 9). ¹³⁰

EPR has been used to confirm the formation of mixed valence polyoxidovanadate species in HEWL crystals treated with $[V^{IV}O(acac)_2]$ (acac is acetylacetonato) 131,132 (Fig. 1). During soaking of this compound in HEWL crystals, under different experimental conditions, $[V^{IV}O(acac)_2]$ degrades, acac ligand is released and at least a part of the V^{IV} centers oxidizes to V^V forming $[V_4^VO_{12}]^{4-}$, $[V_{20}O_{54}(NO_3)]^{n-}$, based on the $\{V_{18}O_{46}\}$ cage, 131 $[V_7^VV_8^{IV}O_{36}(OH_2)]^{5-}$, $[V_7^VV_8^{IV}O_{33}(OH_2)]^+$ and $[V_{20}O_{51}(OH_2)]^{n-}$ (the latter is based on an unusual $\{V_{18}O_{43}\}$ cage). 132 Details of the interactions formed by these unusual compounds with HEWL are shown in Fig. 10.

These data enrich the repertoire of structures of polyoxome-talate/protein structures reported in the PDB. 133-137 Since POVs can act as catalysts in a wide range of reactions and can act as antiviral, antibacterial, and antitumor agents, these studies open the way to future research on the reactivity of POV/protein adducts.

EPR has also been used to investigate the binding of [dichlorido(dmb)(cym)ruthenium(II)] compound (where dmb = 1,3-dimethylbenzimidazol-2-ylidene and cym = p-cymene)(3, Fig. 1) to HEWL. ¹³⁸ The structure reveals the formation of a bidentate [Ru(dmb)(OH $_x$)Cl $_2$]/HEWL adduct (x is not determined) close to the side chain of Arg14 and His15, the binding of [Ru(dmb)(OH $_x$) $_2$ Cl $_2$] to the side chain of Lys33 and of a [RuCl(OH $_x$) $_4$] fragment close to the carbonyl oxygen of Ala107, thus suggesting that the arene ligand is lost upon the HEWL binding, while the dmb remains coordinated to the metal. EPR has been used to evaluate the effect of the cym ligand exchange on the Ru oxidation state. ¹³⁸ The adduct formed by 3 with HEWL is paramagnetic, consistent with the presence of a mononuclear Ru^{III} species, ¹³⁹ as observed also in the case of HEWL crystals treated with AziRu. ⁶⁴

EPR sensitivity to structural changes around the CuII centers has been used to obtain structural information on the binding to HEWL of the Cu^{II} complex with a threonine derivative (CuST), a superoxide dismutase mimic. EPR spectra collected on this system have been compared to those registered for CuST-Im, an imidazole(Im)-bound Cu(II) complex formed when CuST reacts with Im. The g_{\parallel} and g_{\perp} values for CuST-Im were 2.25 and 2.07. The adduct with HEWL (CuST@HEWL) exhibited g_{\parallel} and g_{\perp} values of 2.24 and 2.06, respectively. These data suggest that the Cu(II) centers of CuST-Im and in the adduct adopt a square planar or square pyramidal geometry. The comparison between g_{\parallel} and $|A_{\parallel}|$ values of CuST@HEWL and CuST-Im indicates smaller distortion of the metal coordination sphere in CuST@HEWL than in CuST-Im, in agreement with the crystal structure, where the Cu center binds the His15 side chain.140

Protein metalation studied by a combination of X-ray crystallography and quantum chemistry (density functional theory)

Although the combined use of spectrometric and spectroscopic techniques with XRD data permits us to collect a lot of information about the metal compound/protein interactions, computational methods were often necessary to define the origin of the products obtained by the reaction of metal compounds with proteins, fate and reactivity of the metal/protein adducts when in contact with additional ligands. Density functional theory (DFT) is a quantum-mechanical (QM) method that is used to calculate electronic and structural properties of multi-electron system modeling structures up to several hundreds of atoms.141 Proteins are too complicated systems for this computational technique. For this reason, simplified structural models,142 like those obtained using the cluster approach recently reviewed by Himo and co-workers, 143,144 have been used to study the metalation process by computational methods. Using DFT calculations and a larger cluster model able to represent the protein environment of the metal binding site, T. Marino and coworkers studied the metalation process of RNase A by AP-1145 (Fig. 1), comparing the computational results with those obtained solving the structure of the metal/protein adduct.61 According to these authors, the

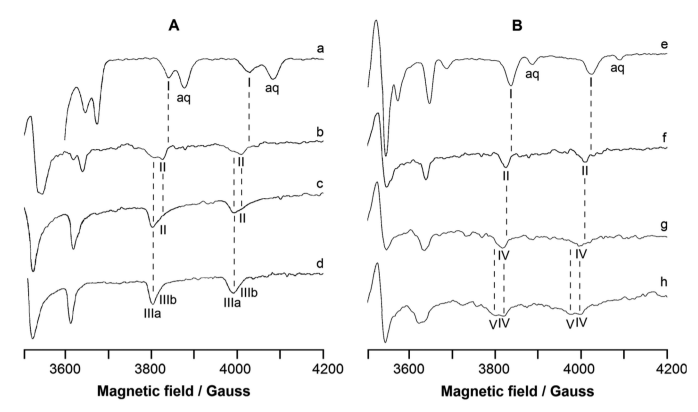


Fig. 9 In solution EPR spectra at 120 K of the systems: (A) $[V^{IV}O(8-HQ)_2]$ -RNase A (metal: ligand: protein molar ratio = 1:2:x with x = 0-1) and (B) $[V^{IV}O(8-HQ)_2]$ -RNase A (metal: ligand: protein molar ratio = 1:1:y with y = 0-1). The spectra were recorded with $V^{IV}O^{2+}$: 8-HQ: RNase A 1:2:0 at pH 3.0 (trace a), VIVO²⁺: 8-HQ: RNase A 1: 2: 1 at pH 5.1 (trace b), at pH 6.4 (trace c) and pH 7.4 (trace d); VIVO²⁺: 8-HQ: RNase A 1: 1: 0 at pH 4.0 (trace e); VIVO2+: 8-HQ: RNase A1:1:1 at pH 5.1 (trace f), at pH 7.4 (trace q) and at pH 8.4 (trace h). V concentration is 1.0 mM in traces a and e, and 0.8 mM in traces b-d and f-h. I indicates MI = 5/2, 7/2 resonances of $[V^{IV}O(8-HQ)(H_2O)_2]^+$, II of the adduct $[V^{IV}O(8-HQ)(H_2O)]^+$ -RNase A with Asp/ Glu-COO-coordination, IIIa and IIIb of the two isomers cis-[VIVO(8-HQ)₂(H₂O)], IV of the adduct [VIVO(8-HQ)(H₂O)]⁺-RNase A with His-N coordination, and V of the adduct [VIVO(8-HQ)(OH)]-RNase A with His-N coordination or [VIVO(8-HQ)(H₂O)]⁺-RNase A with Ser/Thr-O⁻ coordination. while ag denotes MI = 7/2 resonances of the agua ion $[V^{IV}O(H_2O)5]^{2+}$. The positions of MI = 5/2, 7/2 resonances of I, II, IIIa, IIIb, IV, and V species are also indicated by the dotted lines. Reproduced from ref. 130, which is licensed under a Creative Commons Attribution 3.0 Unported Licence, with permission from RSC, copyright 2023.

binding to histidine of the AP-1 fragment found in the structure of the AP-1/RNase A adduct takes place by the aquated AP-1 form and not by AP-1 itself, following the scheme reported in Fig. 11.145 DFT has also been used to gain insights into the reactivity of the metal/protein adduct with additional ligands. It has been shown that metal/protein adducts within protein crystals can still react with ligands. The post-protein binding reactivity of the [Re(CO)₃(OH₂)₂]⁺ fragment bound to His15 of HEWL has been investigated by Zobi and Spingler. 146 These authors have shown that the [Re(CO)3(OH2)2]+/HEWL adduct reacts with Im, pyridine-2-carboxylic acid or L-serine leading to metal ligand substitution.146 Along the same line, XRD data have demonstrated that when dirhodium(II,II) tetraacetate ([Rh₂(μ-O₂CCH₃)₄], Fig. 1) reacts with RNase A an adduct with the dirhodium center bound to the side chains of His105 and His119 is formed. 12,56 This adduct can react with Im, leading to the formation of an unexpected product with Im that binds the dirhodium center at an equatorial site rather than at the expected axial site.147 The origin of this surprising experimental evidence has been studied using DFT. A small portion

of the structure of the dirhodium/protein adduct around the His105 residue has been used as a model (cluster model) to investigate the reactivity of Im to the three available sites: (1) the only free axial site (Route 1 in Fig. 12), (2) the equatorial position to the Rh atom that was not coordinated to His105 (r2) (Route 2 in Fig. 12), and (3) the equatorial position to the Rh atom that was directly bound to the His (r) (Route 3 in Fig. 12). A corresponding minimal model has been used for comparison. The results suggest that Im favourably replaces the water molecule at the (r) equatorial site of the hydrated species $[Rh_2(Im)(\mu-O_2CCH_3)(H_2O)_7]^{3+}$, where Im mimics the His105 side chain. The electronic properties of [Rh₂(Im) $(\mu - O_2CCH_3)(H_2O)_7^{3+}$, where the two metal centers are not equivalent, are responsible for such unexpected Im binding.

The binding of [Ru₂Cl(D-p-CNPhF)(O₂CCH₃)₃] (D-p-CNPhF-= N,N'-bis(4-cyanophenyl)formamidinate, Fig. 1) to His105 of RNase A has been recently studied using a similar approach. The structure of the adduct formed upon reaction of the diruthenium compound with the protein showed that the diruthenium center anchors to the His side chain at the axial

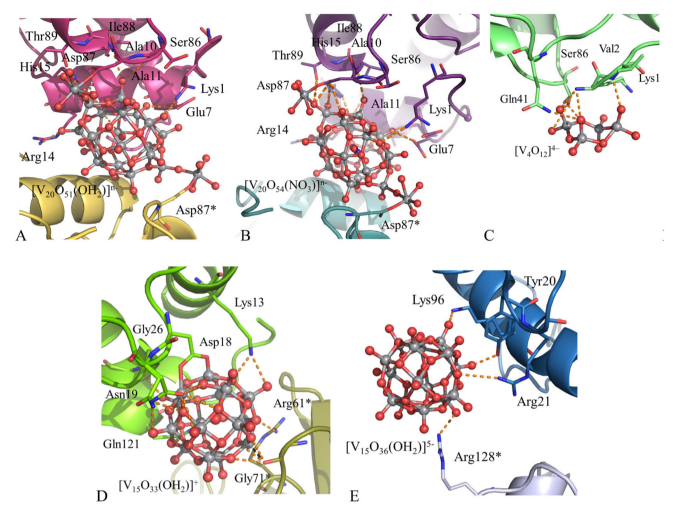


Fig. 10 Interactions of $[V_{20}O_{51}(OH_2)]^{n-}$ (panel A), $[V_{20}O_{54}(NO_3)]^{n-}$ (panel B), $[V_4^VO_{12}]^{4-}$ (panel C), $[V_7^VV_8^NO_{33}(OH_2)]^+$ (panel D) and $[V_7^VV_8^NO_{36}(OH_2)]^{5-}$ (panel E) with HEWL residues in crystals of the protein treated with [VO(acac)₂].

Fig. 11 Proposed mechanism of the metalation process of RNase A by AP-1. TS stands for transition state. His in red is used to mimic His of the protein.

site. 148 In the adduct, two water molecules replace an acetate equatorial ligand. To understand the origin of the formation of the hydrolyzed species, quantum chemical calculations have been carried out. 148 QM calculations have also been performed to estimate the O- disproportionation mechanism catalyzed by CuST@HEWL140 and to study phenylacetylene polymerization by Rh complex encapsulated in hl-L-Ft. 103,149

Protein metalation studied by a combination of XFEL and QM/ MM

In recent years, we have seen the advancement of X-ray-free electron lasers (XFELs) and their use in various scientific domains. XFELs produce powerful X-ray femtosecond (10⁻¹⁵ s) pulses that can be used to solve the structure and dynamics of

Fig. 12 Substitution reactions with inclusion of Im for the dirhodium/protein adduct (cluster model).

matter. 150 The structures can be merged into "molecular movies", which can provide information on the biological macromolecules in action. Recently, time-resolved serial femtosecond crystallography (TR-SFX) using XFEL has been used to characterize a light-induced CO-release from Mn to HEWL within porous metal/protein adduct microcrystals. 116 In this study, conformational variations of the metal/protein adduct during the CO release from the metal center upon light exposure have been evaluated. In darkness, crystals of HEWL in the presence of [Mn(CO)₃(acetone)₃](CF₃SO₃) (Fig. 1) show a $[Mn(CO)_3(wat)_2]^{2+}$ fragment bound to the side chain of His15. At this site, the metal adopts an octahedral geometry, with two CO and two water ligands equatorially coordinated (Mn-CO_{eq} and Mn-OH2,eq bonds) and Im of His and the third CO that are axially coordinated (Mn-COax bond). The structures refined using data collected after 10 ns, 100 ns, and 1 µs of light excitation using a 20 µJ pump laser indicate a greater susceptibility to radiation of the Mn-CO_{ax} bond when compared to the Mn-CO_{eq} bond. Comparison between the solved structures permits them to capture the real-time intermediate formation. Interestingly, the paper also describes the structure of the protein after the complete release of CO. This structure does not show the presence of Mn coordinated to His15. After complete CO release, the Mn-aqua species that probably forms is unstable and immediately releases from the protein. CO release from metal-based CO releasing molecules bound to

proteins had previously been described by collecting snapshots from different crystal structures of Ru-CORMs/HEWL adducts.52,151

QM/molecular mechanics (QM/MM) calculations has been used to elucidate the detailed reaction processes in photodissociation. 116 The QM/MM calculations combine the accuracy of ab initio QM calculations and the speed of the classic approaches, thus allowing the study of chemical processes in large systems as proteins. 152 The QM/MM results predict that the release of the Mn(CO)-unit is faster than the exchange of third CO in the equatorial position with water molecules, in agreement with crystallographic observation.

Protein metalation studied by a combination of X-ray crystallography and other computational approaches

Molecular docking is the computational method that allows the prediction of small molecule binding sites on a biological macromolecule structure. Docking is frequently used in structure-based drug design, but it has been recently combined with DFT calculations and crystallographic experiments for a better definition of the stability of the metal/protein adduct and to highlight the role of specific interactions on the metal compound/protein recognition process. For example, in the structure of the adduct formed upon the reaction of [VIVO(8- $HQ)_2$ with RNase A, a $[V^{IV}O(8-HQ)(H_2O)]^+$ ion binds the protein close to the side chain of Glu111. 130 Docking experi-

ments have been used to refine the metal binding site revealing the reasons behind the ligand exchange process, not observed for similar complexes like [V^{IV}O(pic)₂]⁷⁶ and not predicted by thermodynamic considerations. The computational data show that the [V^{IV}O(8-HQ)(H₂O)]⁺/RNase A adduct forms from [VIVO(8-HO)₂]/RNase A and that its formation is facilitated by the presence of the Gln69 side chain.

Similarly, docking experiments have been performed to study the binding of the same compound to HEWL. X-ray diffraction analyses showed that [VIVO(8-HQ)(H2O)]+ coordinates to the Asp119 side chain. The binding is stabilized by interactions with Asn103 and Trp62 from a symmetry-related molecule.⁷⁴ First, the structure of $[V^{IV}O(8-HQ)(H_2O)_2]^+$ has been optimized by DFT. In the second step, the coordinative interaction of [V^{IV}O(8-HQ)(H₂O)]⁺ with the protein has been investigated to unveil the molecular reasons why the binding of [V^{IV}O (8-HQ)(H₂O)]⁺ to HEWL occurs at the Asp119 side chain and not at other potential binding sites such as Glu35, Asn46, Asp48 and Asp52 side chains. To assess the effect of the crystal lattice or, in general, of protein-protein interactions on the binding of $[V^{IV}O(8-HQ)(H_2O)]^+$ to the protein, two different docking approaches have been applied: (1) An ensemble docking, denoted as the "classic model", which uses one metal-containing fragment and the three superimposed structures of HEWL; and (2) a docking denoted as the "symmetry mate model", which uses one metal-containing fragment with two symmetry-related protein chains. This calculation has been carried out to model the crystal lattice or protein-protein interactions. According to ensemble docking results, [VIVO(8-HQ)(H₂O)]⁺ should interact with Glu35 or Asp52 side chains. Thus, protein-protein stabilization has to be invoked to explain the experimental data. Indeed, the coordination of the V center to the carboxylate-O of Asp119 is observed using the "symmetry mate model". Overall, these data indicate that protein-protein stabilization may facilitate unexpected binding of metal-based compounds to proteins, offering new insights into the interaction mechanisms of metal-based drugs with proteins.74

A series of docking simulations have also been performed to gain insights into the V compound chiral discrimination by HEWL. The computational data indicate that the formation of the HEWL-VO-bipy adduct is driven by the presence of structured water molecules, suggesting the pivotal role of microsolvation for chiral discrimination of the binding region and in stabilizing the final structure of the metal/protein adduct.129

Molecular dynamics (MD) simulations also gave a contribution in defining the interaction of metal compounds with proteins, in combination with X-ray crystallography. Recently, for example, the structures of HEWL adducts with isostructural NHC complexes with Ru, Os, Rh, and Ir were further studied using constant-pH molecular dynamics (CpHMD) simulations. The X-ray structures of these adducts reveal that [Ru(dmb) (H₂O)Cl₂] binds His15 and Arg14, [Os(cym)(dmb)Cl] interacts with Asn103; [Rh(Cp*)Cl₂] binds His15, while [Ir(Cp*)(dmb)Cl] the side chain of Asp101.

CpHMD simulations allow us to evaluate pK_a values of the metal-binding residues. CpHMD indicates that His15 and Lys33 have pK_a of 6.67 and 9.91, respectively, while Asp87 and Asp101 have p K_a s of 2.73 and 0.44. This result suggests that these two Asp have pK_a values significantly lower than those observed in the case of solvent-exposed Asp $(pK_a = 4)$ and provides an explanation for the hydrogen bonds that Asp87 forms with His15. This interaction probably affects the electropositivity of His15, providing insights into the observed specificity of His15 for Ru and Rh compounds vs. Os and Ir, which prefer electronegative binding pockets. 70

Classic MD simulations have also been used in other studies. The X-ray structures of HEWL, proteinase K and thaumatin (Fig. 2) in the presence of Tb-Xo4 (G1, Fig. 1) and three variants (Tb-Xo4-NMet2, Tb-Xo4-OH and Tb-Xo4-SO3, denoted as V1, V2 or V3, Fig. 1) have been solved. To gain insights into the role of pendant arms in the structure of G1, V1, V2 or V3 on the binding to proteins, MD simulations (200 ns) have been carried out on systems containing metal compounds and four HEWL molecules. These starting models mimic the protein crystal packing. Results indicate that V1, V2 or V3 can bind one protein chain through interaction formed by carboxylates with the metal; the pendant arms can interact with surface residues of the other protein molecules with ammonium or sulfonate moieties that form salt bridges with acid residues and the OH group that forms hydrogen bonds, both as H donor or acceptor.153

Computational studies have also been used to help the refinement process of crystal structures of metal/protein adducts. In the adduct formed upon soaking of VIVOSO4 in HEWL crystals, a metal center binds a water molecule, an O_{oxido}, a Cl⁻ ion, and the side chains of Asp46 and Asp52. The sixth ligand, needed to complete the octahedral coordination sphere, is not unambiguously identified by the authors, since it cannot be seen in the electron density map at the 1.0σ level. Refinement of this site addressed by docking and QM/MM calculations suggests that the sixth coordination position is probably occupied by a water molecule. 129

Conclusions

In the last 15 years, X-ray crystallography has provided a detailed description of the protein metalation process in several cases. However, significant progress can be made in this field since many aspects of the protein metalation process remain still unknown.8 The combined use of X-ray crystallography and biophysical techniques, as described in this work, has certainly allowed substantial progress in this specific research area. Of course, there are many other biophysical techniques that can be used to complement X-ray crystallography in the definition of the structure of metal/protein adducts. Examples are synchrotron terahertz and neutron scattering spectroscopy, 154 and capillary electrophoresis. 155 Putting together the results obtained using various physicochemical methods, molecular mechanisms underlying the metallodrug/

protein process and structural/functional features of metal/protein adducts can be elucidated with high accuracy.

Data that have been collected so far demonstrate that the binding of metallodrugs to proteins does not alter the overall in-crystallo conformation of the biological macromolecules¹⁵⁶ and only in a few cases it alters their stability. 38,78 However, the binding can significantly alter the catalytic activity of enzymes⁶² and their recognition by other protein partners. The binding of metal containing fragments to proteins can result in a modification of the metal local environment, significantly affecting its reactivity with additional ligands and sometimes creating a novel catalytic activity. Although there are many potential metal binding sites on protein surfaces, the coordination of metals to residue side chains takes place preferentially only in a few positions, suggesting that the protein metalation is a selective process. The preferential choice of some residues with respect to others is determined by different parameters: solvent accessibility and flexibility of the protein residues involved in the first steps of the metal compound/protein recognition process, steric/electronic effects due to metal ligands or protein residues, and potential interactions that metal ligands can form with protein surface patches. Hardhard and soft-soft interactions of course contribute to driving the metallodrug towards its final target. In this respect, it is worth noting that the reactivity of metal complexes with proteins is certainly regulated by the experimental conditions used for the analysis. Thus, the pictures that emerge using different techniques that require different solutions and conditions can sometimes diverge.

Further studies will greatly accelerate the comprehension of reaction mechanisms of metallodrugs with protein targets, thus aiding the design of novel therapeutics. In this respect, recent reviews have summarized the methods that can be used identification by metal-based agents. 157,158 The accumulated knowledge will also facilitate the design of novel protein-based drug delivery systems, allowing the development of systems with precise mechanism of metallodrug release, and the production of new artificial metalloenzymes, allowing the development of systems that catalyze new reactions. Artificial intelligence and machine learning are groundbreaking tools that can be integrated with existing methods to accelerate this process even further. The study of the interaction of metal compounds with proteins in real cellular systems is certainly enormously more difficult. There are thousands of proteins within a cell and a significant number of additional low molecular weight metabolites at variable concentrations. 159 Nonetheless, we believe that studying the protein metalation process using different techniques can certainly represent a useful basis for the understanding of this process within cells.

Data availability

Data are all available in the literature or deposited in the PBD.

Conflicts of interest

There are no conflicts to declare.

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