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Structural insights into copper and zinc binding to tau protein and the impact of metal binding on amyloid aggregation

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Tau protein is a microtubule-associated protein central to the pathogenesis of Alzheimer's disease (AD) and other tauopathies. While metal ion homeostasis is disrupted in AD, the presence of copper and zinc in neurofibrillary tangles suggests a pathological role for metal–tau interactions. In this study, the metal binding properties of Tau441 were probed using a wide range of spectroscopic tools. Specifically, electron paramagnetic resonance, circular dichroism, nuclear magnetic resonance (NMR) and X-ray absorption spectroscopy (XAS) results point to a single high-affinity Cu²⁺ binding site within the microtubule-binding domain (MTBD), coordinated by the bis-His motif in R3 (His329/His330), possibly His299 and an oxygen-based ligand. This complex can be reduced resulting in a trigonal Cu⁺–Tau441 complex involving Cys322, His299 and a third ligand (likely Cys291 or His329/330), as characterized by XAS and NMR. NMR and XAS results indicate the presence of three Zn²⁺ binding sites: one high-affinity site in the MTBD involving His299, His330, Cys322 and Asp295, and two lower-affinity sites in the N-terminal region, coordinated predominantly by carboxylate and His residues. Moreover, the impact of Cu²⁺ and Zn²⁺ ions on the amyloid aggregation of full length Tau441 was evaluated using thioflavin T fluorescence, electrophoresis and transmission electron microscopy. Both metal ions significantly accelerate aggregation, promoting the formation of amyloid fibrils with distinct morphologies. Our study provides valuable structural insights into the copper and zinc binding sites in Tau441 that provide a rational basis to understand the impact of metal ions on amyloid fibril aggregation and morphology. The current study expands the bioinorganic facet of AD and other tauopathies, and it underscores the importance of metal–tau interactions as potential therapeutic targets in these neurodegenerative diseases.

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Introduction

Tauopathies are a group of neurodegenerative disorders characterized by the abnormal deposition of the microtubule-associated protein tau in the brain.¹ Among these, Alzheimer's disease (AD) is the most prevalent, defined pathologically by the presence of extracellular amyloid-beta (Aβ) plaques and intracellular neurofibrillary tangles (NFTs) composed primarily of

hyperphosphorylated tau.² Increasing evidence suggests that disruption in metal homeostasis, particularly involving copper and zinc, contributes to the pathogenesis of AD and other tauopathies.^{3,4} Indeed, zinc is increased, and copper is decreased in the AD brain, while blood levels of labile copper, *i.e.* not bound to ceruloplasmin, are also increased in a subset of AD patients, as compared to healthy ones.⁵ These metals are essential for normal central nervous system function, as they can regulate different neurotransmitter receptors,^{6,7} and they are cofactors in several important enzymes. For example, superoxide dismutase 1 requires copper and zinc, dopamine-β-hydroxylase is a copper-dependent enzyme critical in the synthesis of the neurotransmitter norepinephrine⁸ and zinc plays structural roles in a variety of transcription factors.⁹ The loss of metal homeostasis can lead to metal accumulation, undesired metal redox activity and metal-induced protein aggregation.^{5,10,11} X-ray fluorescence analysis has demonstrated the presence of both Cu and Zn in senile plaques of AD patients,¹² while metal accumulation has been correlated with Aβ deposits in AD transgenic murine models.¹³ Indirect

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evidence for the presence of copper and zinc in NFTs of AD patients has been derived from copper-dependent oxidase activity¹⁴ and the use of fluorescence zinc chelators.¹⁵

The early discovery that A β is a metal binding peptide¹⁶ and that extracellular amyloid plaques in AD concentrate copper and zinc ions,³ has driven extensive research into the metal binding properties of A β peptide in the last two decades.^{5,17} Copper and zinc can bind A β and impact its aggregation, while copper-bound peptides display redox activity that can contribute to the generation of reactive oxygen species, thereby exacerbating neuronal damage.^{18,19} While the presence of metal ions in NFTs suggests a direct pathological role for metal–tau interactions, the metal binding properties of tau and the impact of metal ions on tau protein misfolding and aggregation have not been investigated as extensively as in the case of A β .

Tau protein exists in six major isoforms in the adult human brain, generated through alternative splicing of the microtubule associated protein tau (MAPT) gene, and characterized by the number of N-terminal negatively charged inserts (0N, 1N, or 2N) and the presence of either three (3R) or four (4R) microtubule-binding repeats.²⁰ Tau441 is the longest isoform comprising a 2N4R structure (Scheme 1), and together with 2N3R and 2N4R, represents the most expressed isoforms in the adult brain. The most studied model fragments containing the microtubule binding domain (MTBD) are TauK18,²¹ encompassing only the MTBD repeats, and TauK32²² that also contains two regions flanking the MTBD (Scheme 1). These models have been used to study microtubule binding as well as protein aggregation. Tau441 undergoes different posttranslational modifications (PTMs)²³ under physiological and pathological conditions. Tau phosphorylation, normally regulated by different kinases and phosphatases, is exacerbated in AD. Under physiological conditions, tau stabilizes microtubules and supports axonal transport,²⁴ and it is also located in the nucleus, where it is involved in nuclear maintenance.²³ Canonically, tau is an intracellular protein, but recent evidence suggests that tau

exists in the extracellular environment under physiological and pathological conditions, although the role that it may play extracellularly remains unclear.²⁵

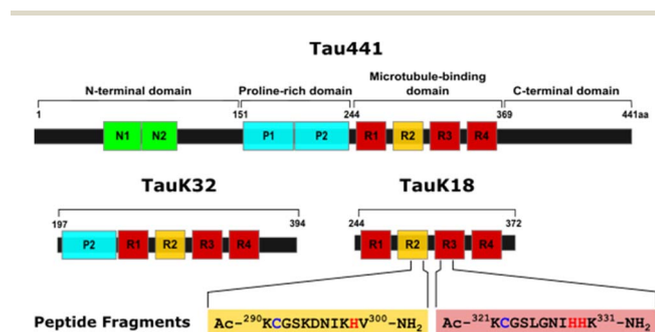
Tau is an intrinsically disordered protein in solution with a very low transient secondary structure.²⁶ Tau is a protein with a low tendency to aggregate; however, in the presence of cofactors such as heparin and negatively charged polymers, the protein forms aggregates.²⁷ Tau has been observed to aggregate in the presence of metal ions. Copper can induce aggregation of R1–R4 peptides,²⁸ and oligomerization of Tau441 *in vitro*.²⁹ In triple transgenic (3xTg-AD) mice, dietary copper administration causes increased tau phosphorylation, aggregation and toxicity.³⁰ Zinc accelerates the aggregation of TauK18³¹ and induces conformational changes and oligomerization of Tau441.^{32–35} Indeed, early pioneering studies suggest that both copper and zinc ions, may bind to tau. Soragni *et al.*²⁹ examined the copper binding properties of TauK32 by nuclear magnetic resonance (NMR) spectroscopy and isothermal titration calorimetry (ITC), finding a single copper coordination site with a metal binding affinity in the micromolar range ($K_d = 0.7 \mu\text{M}$), similar to that of the full-length protein ($K_d = 0.5 \mu\text{M}$); however, structural characterization of the copper binding site in the full length Tau441 has not been further explored. However, zinc binding to Tau441 has been studied by ITC, revealing that it may have up to four metal-binding sites,³³ while ITC and mutagenesis studies of the TauK18 fragment show that this fragment has a single metal-binding site with binding affinity in the micromolar range, and that Cys291 and 322 and His268, 299, 329, 330 and 361 may be important for metal binding.^{31,36} Although these early studies reveal that tau can bind copper and zinc, the molecular and structural details of these metal–tau interactions have not been elucidated.

Given the fact that metals accumulate in NFTs, understanding the metal binding properties of tau should yield insight into the structural and biochemical basis of metal-induced aggregation. In this study, the metal binding properties of Tau441 were probed using a wide range of spectroscopic tools. Specifically, Cu²⁺ binding to Tau441 was studied by electron paramagnetic resonance (EPR), circular dichroism (CD), NMR and X-ray absorption spectroscopy (XAS). Reduction of the Cu²⁺–Tau441 species was probed by EPR and XAS, while the nature of the resulting Cu⁺–Tau441 complex was studied by XAS and NMR. The Zn²⁺ binding properties of Tau441 were examined by NMR and XAS. Finally, the impact of Cu²⁺ and Zn²⁺ ions on the amyloid aggregation of full length Tau441 was evaluated, while the nature of the resulting aggregates was examined by electrophoresis and transmission electron microscopy. This spectroscopic study provides valuable insights into the copper and zinc binding sites in Tau441 and expands the bioinorganic facet of AD and other tauopathies.

Results

Tau has a single Cu²⁺ binding site

Cu²⁺ binding to Tau441 was studied by titrating the protein with 0 to 2 equivalents of the metal ion, followed by electron paramagnetic resonance (EPR) and circular dichroism (CD)



Scheme 1 Tau441 and fragments. (Top) Schematic representation of the full-length tau protein, Tau441. Primary sequence of Tau441 includes an N-terminal, with N1 and N2 that are negatively charged inserts, P1 and P2 are the proline-rich region; the microtubule binding domain (MTBD) encompasses the four pseudo-repeats, R1–R4 and finally the C-terminal domain. (Middle) TauK32 and TauK18 are fragment models of the full-length tau and encompass the MTBD. (Bottom) Sequence of the two short peptides of the R2 and R3 domains that contain Cys291, Cys322, His299 and the bis-His motifs His329 and His330.



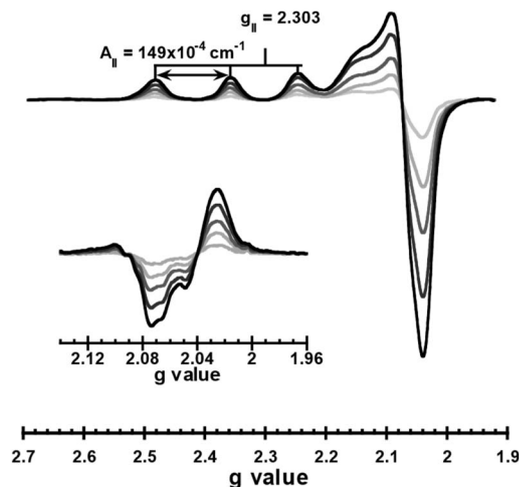


Fig. 1 Tau441 has a single Cu^{2+} binding site. Titration of 0.2 mM Tau441 in 20 mM MOPS buffer at pH 7.4 with Cu^{2+} ions (with 0, 0.25, 0.5, 0.75, 1.0 and 1.5 equiv., represented by lines from light gray to black), followed by electron paramagnetic resonance (EPR) spectroscopy; the inset shows the second derivative of the EPR spectra.

spectroscopies (Fig. 1). Having a d^9 electronic configuration, Cu^{2+} ions have an unpaired electron, and EPR spectroscopy can give insights into the chemical environment around the metal ion. EPR spectra of Tau441 with Cu^{2+} show one set of signals, indicating the presence of a single Cu^{2+} coordination site (Fig. 1). It should be noted that these signals are distinct from those of Cu^{2+} in buffer (Fig. S1), and hence, can be assigned to Cu^{2+} bound to Tau441. The EPR signals of the Cu^{2+} -Tau441 complex ($g_{\parallel} > g_{\perp}$) are indicative of a $d_{x^2-y^2}$ ground state, with $g_{\parallel} = 2.304$ and $A_{\parallel} = 149 \times 10^{-4} \text{ cm}^{-1}$. The second derivative of the EPR spectra in the perpendicular region reveals hyperfine structure, likely corresponding to His nitrogen super-hyperfine couplings (inset Fig. 1). The titration of Tau441 followed by CD (Fig. S2A) results in no signals in the range of 0–1.0 equiv. of Cu^{2+} ; hence the Cu^{2+} -Tau441 complex does not give rise to chiral electronic transitions, suggesting that it might have more than two equivalent ligands in the coordination sphere.

To elucidate the regions of Tau441 involved in Cu^{2+} coordination, TauK18 and the peptide fragments R2 and R3 were studied. TauK18 encompasses residues 244–372 of the full-length protein, containing the four pseudo-repeats of the microtubule binding domain (MTBD). The R2 and R3 peptides correspond to residues 290–300 and 321–331, respectively, and contain residues that have been previously identified as putative ligands for Cu^{2+} binding to TauK32, namely His residues 299, 329 and 330.²⁹ Each fragment was titrated with 0–2 equiv. of Cu^{2+} and followed by CD and EPR (Fig. S2C–F); the CD signals that arise from Cu binding to R2 and R3 fragments saturate below 2 equiv., while EPR shows the presence of one main Cu^{2+} species that is the same at 1 and 2 equiv. For TauK18, no CD signals were observed (Fig. S2B), as in the case of Tau441, indicating that the metal–tau complex is not optically active. A comparison of the EPR and CD spectra of Cu^{2+} bound to Tau441, TauK18 and the R2 and R3 fragments at 1 equivalent

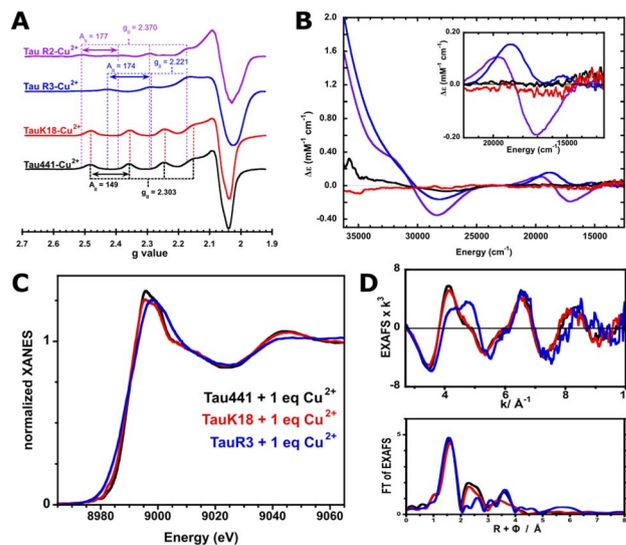


Fig. 2 Spectral comparisons of Tau441 and fragments with Cu^{2+} . EPR (A), CD (B), Cu K-edge XANES (C), EXAFS and FT (D) spectra of Cu^{2+} bound to Tau441 (black), TauK18 (red), TauR3 (blue) and TauR2 (purple) fragments; all with 1 equiv. of Cu^{2+} in 20 mM MOPS buffer at pH 7.4.

clearly shows that only TauK18 replicates the Cu^{2+} coordination site in Tau441 (Fig. 2A and B). Although R2 and R3 display complex metal–peptide speciation (Fig. S2C–F), none of the observed species resemble the Cu^{2+} site in Tau441. Consistently, the Cu K-edge X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) spectra for the Cu^{2+} complexes with Tau441 and TauK18 are identical, while that of R3 are quite different (Fig. 2B–D). While all XANES spectra (Fig. 2C) show a white line at 8990 eV that is a characteristic edge value for tetragonal square-planar Cu^{2+} centers (typically known as type 2 Cu centers),^{37,38} distinct differences are observed in the edge shape and EXAFS spectra of the protein complexes (red and black in Fig. 2D), as compared to the R3 peptide (blue in Fig. 2D). EXAFS yields information about the local structure and nearby atoms around the metal center; hence, the EXAFS spectra shown in Fig. 2D clearly indicate that the coordination sphere around the Cu^{2+} ions in TauK18 and Tau441 is practically the same, while that of the Cu^{2+} -R3 peptide complex is completely different.

Overall, R2 and R3 peptide fragments cannot reproduce the Cu^{2+} binding features of Tau441, while the shorter protein TauK18 can bind this metal ion in a similar fashion to Tau441. Hence, we can conclude that metal binding to tau peptide fragments is different from that to the full protein, underscoring the importance of studying metal binding to physiologically and pathologically relevant forms of tau.

Cu^{2+} binding to Tau441 involves amino acids in R2 and R3

The Cu^{2+} coordination site in Tau441 was explored at single-residue resolution by ^1H - ^{15}N Heteronuclear Multiple Quantum Coherence (HMQC) NMR, which displays cross-peaks for each amide group in the protein, and thus, provides



multiple probes to locate the Cu^{2+} -binding site. Having an unpaired electron, Cu^{2+} causes paramagnetic relaxation effects (PREs), resulting in the differential broadening of the amide cross-peak signals by bond-mediated or through space interactions, providing a means of mapping metal-binding sites in the protein. A titration of Tau441 with Cu^{2+} was followed by ^1H - ^{15}N HMQC spectroscopy. While addition of 0.01 equiv. of the metal ion causes a drastic broadening of all amide resonances, addition of 0.005 equiv. causes moderate broadening of certain residues of the protein, as compared to that of Tau441 with no metal ion (Fig. 3A and B). A ratio of signal intensities before and after the addition of the metal ion (I/I_0) (Fig. 3C) allows for the identification of two regions that are affected by copper binding. The most affected region encompasses residues between 275 and 330 (R2 and R3 domains), containing His299, His329 and His330, which could be participating in copper binding. Other regions affected by copper are in the C-terminal domain, namely: residues 370–380 that contain His374, and the proline-rich domain encompassing residues 210–230. However, these residues are likely not involved in metal binding, since TauK18 lacks the C-terminal domain, and still yields a similar copper complex to Tau441, as assessed by EPR, CD and XAS (Fig. 2). Signal broadening in the 370–380 region might be due to electrostatic interaction between the C-terminal and the R2/R3 domains, which would cause PREs due to the proximity to the copper site. However, the proline rich domain has a propensity to form a transitory secondary structure, which could be affected by metal binding.^{26,39}

However, individual chemical shift perturbations show some changes in the following regions: 52–63, 82–89 and 104–132 in the N-terminal domain; 190–226 in the proline-rich region proximal to the microtubule binding domain; the region flanking the R3 region (285–326); 339–372, 386–410 and 434–435 of the C-terminal regions (Fig. 3D). Of all these, residues 285–326 are within the region that displays the most line broadening due to PREs, where metal binding is likely to occur, hence the chemical shift perturbations at 285–326 are likely due

to copper binding, while changes in all regions might reflect conformational changes caused by metal binding to Tau441.

Overall, the NMR results reveal that the copper binding site of Tau441 is in the MTBD, within the R2 and R3 domains. Residues in the R2 and R3 domains, namely His299 and the bis-His motifs His329 and His330, are putative ligands for the copper site. Furthermore, long-distance interactions exist between the C-terminus and the copper site of Tau441, as has been previously observed for monomeric tau in the absence of metal ions.²⁶

Structural features of the Cu^{2+} -Tau441 complex

To gain insights into the coordination sphere of Cu^{2+} bound to the full-length tau, EPR simulation and EXAFS fitting were performed. The EPR simulation for the Cu^{2+} -Tau441 complex is consistent with having an axial Cu^{2+} center with $g_{\parallel} = 2.304$ and $A_{\parallel} = 149 \times 10^{-4} \text{ cm}^{-1}$ (448 MHz) (Table 1). The parallel copper hyperfine value is at the lower end of the range of A values for typical tetragonal square-planar type 2 Cu^{2+} -sites (120 – $200 \times 10^{-4} \text{ cm}^{-1}$).⁴⁰ The copper hyperfine value is indicative of the interaction of the electron spin with the nuclear spin of the metal ion, and hence, correlates with the covalency of the Cu^{2+} -complex: a lower value is indicative of larger covalency, *i.e.* more delocalization of the spin density toward the ligands. Hence, the A_{\parallel} value for Cu^{2+} bound to Tau441 suggests increased covalency in the site. The g_{\parallel} and A_{\parallel} values of the Cu^{2+} -Tau441 complex indicate a 2N2O equatorial coordination sphere, according to Peisach–Blumberg correlations (Fig. S3).⁴¹ Consistently, the fine structure in the second derivative of the perpendicular region of the EPR spectrum (Fig. 4A) suggests the presence of super hyperfine couplings arising from non-equivalent nitrogens in the coordination sphere. This super hyperfine pattern can be simulated using two or three non-equivalent nitrogen-based ligands (Table 1 and Fig. 4A, red and blue lines, respectively), with $^{\text{N(His)}}A$ values that are typical for imidazole ligands.^{42–44} These results suggest that the Cu^{2+} -Tau441 complex contains at least two nitrogen-based ligands that are likely His residues.

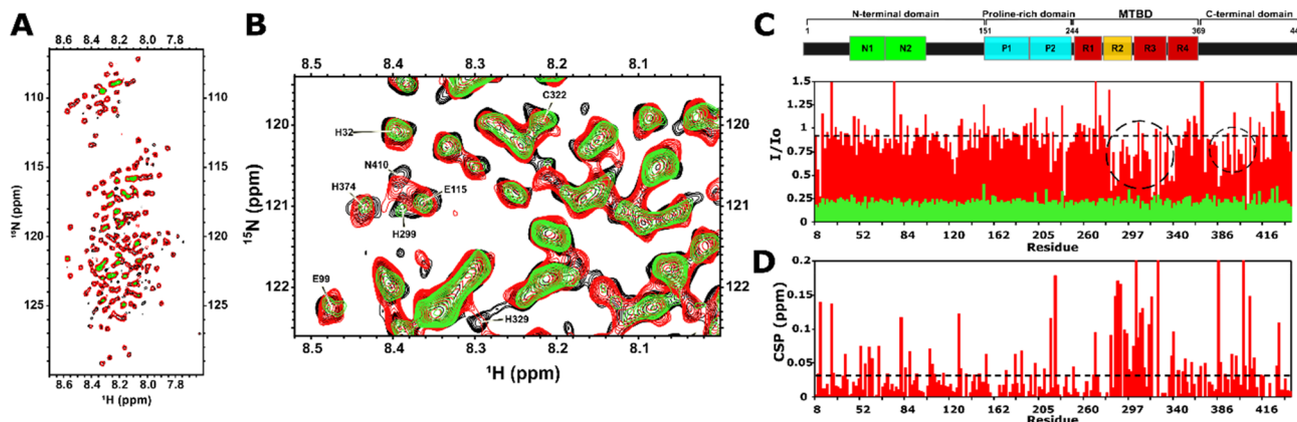
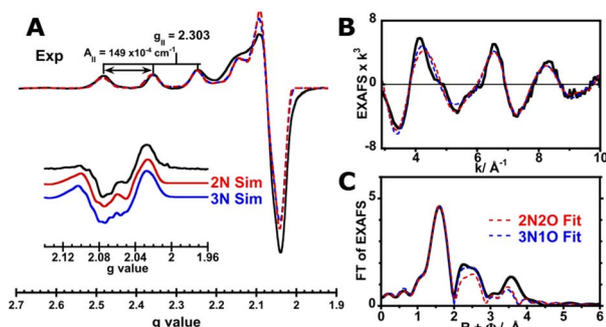


Fig. 3 R2 and R3 domains are involved in Cu^{2+} binding to Tau441 protein. (A) Overlay of ^1H - ^{15}N HMQC NMR spectra of Tau441 100 μM with no metal ion (black), and with 0.005 (red) and 0.01 (green) equiv. of Cu^{2+} ; a zoomed image of the region of ^1H - ^{15}N HMQC (8.00–8.52 ^1H ppm and 119.5–122.5 ^{15}N ppm) spectra is shown in (B). (C) I/I_0 profiles of the ^1H - ^{15}N HMQC NMR signals of assigned residues at different concentrations of Cu^{2+} : 0.005 equiv. (red bars) and 0.01 equiv. (green bars); circles of dotted black lines show the patches due to the PREs of Cu^{2+} . (D) Chemical shift perturbations (CSPs) for the HMQC signals of assigned residues of Tau441 with 0.005 equiv. of Cu^{2+} .



Table 1 Spin Hamiltonian parameters of the EPR simulations for the Cu²⁺-Tau441 complex with different coordination spheres^a

	2N	3N
g_x	2.066	2.068
g_y	2.078	2.078
g_z	2.304	2.304
${}_{\text{Cu}}A_x$	5	10
${}_{\text{Cu}}A_y$	20	20
${}_{\text{Cu}}A_z$	448	448
$\text{N1(His)}A_x$	42	41
$\text{N1(His)}A_y$	37	36
$\text{N1(His)}A_z$	37	36
$\text{N2(His)}A_x$	34	35
$\text{N2(His)}A_y$	39	40
$\text{N2(His)}A_z$	34	35
$\text{N3}A_x$	—	33
$\text{N3}A_y$	—	28
$\text{N3}A_z$	—	28
$g\text{-Strain}_x$	0.012	0.014
$g\text{-Strain}_y$	0.012	0.0095
$g\text{-Strain}_z$	0.020	0.019

^a A values are in MHz units.**Fig. 4** Spectral simulation of the Cu²⁺-Tau441 complex. EPR simulations for the spectrum of Cu²⁺-Tau441 (A) with 2N (red line) and 3N (blue line) coordination spheres using the parameters listed in Table 1; the inset shows the second derivative of the perpendicular region. Fitting of EXAFS (B) and FT (C) of Cu²⁺-Tau441 (black trace) with 2N2O (red dotted line) and 3N1O (blue dotted trace) coordination spheres using the parameters listed in Table 2.

To gain insights into the local coordination environment of the Cu²⁺-Tau441 complex, EXAFS spectra were collected (Fig. 2D and 4B, C black trace) and simulated. EXAFS provides information about the local coordination structure, its nature and distances from the metal center. Considering that EPR results suggest a 2N or 3N coordination sphere for the Cu²⁺-Tau441 complex, likely with His ligands, simulations with two or three nitrogens in the coordination sphere were tested. As for the other ligands, sulfurs from Cys or oxygen-based ligands (water, Asp or Glu) were considered, and hence, the following first coordination spheres were probed: 2N2S, 2N1S1O, 2N2O, 3N1S and 3N1O. Overall, Bond Valence Sum (BVS) analysis⁴⁵ reveals a larger formal charge on the metal center for the coordination models that include sulfur-based ligands (Table S1), achieving values closer to +2 only with the coordination models that include oxygen-based ligands. Moreover, Debye-Waller values for the fits that include sulfur-based ligands (especially 2N2S) are higher (Table S1), further supporting the exclusion of Cu-S interactions. A comparison of the 2N2O and 3N1O fits to the experimental data (Fig. 4B, C and Table 2) shows that they both yield an adequate fit, with reasonable Cu-N distances (1.92–2.02 Å).⁴⁶ In both cases, the Cu-O distances are in the range of 1.88–2.00 Å, which would be consistent with having oxygen-based ligands from a carboxylate ligand, *i.e.* Glu or Asp residues, or water molecules.⁴⁷ Moreover, the scattering pattern at larger distances ($R + \Phi > 2$ Å) that is evident in the FT of the EXAFS (Fig. 4C) is better fitted by including the carbon and nitrogen atoms of His imidazole rings (Table 2). Both, the 2N2O and the 3N1O models yield a reasonable EXAFS fit of the experimental data, although the 3N1O model better fits the scattering patterns at larger distances with a lower R-factor (Fig. 4B and C). Nonetheless, it would be risky to discard one of the two possible coordination models, given the limitations of EXAFS to discern scattering from atoms within a $\Delta Z = \pm 1$ range, *i.e.* to distinguish between nitrogen and oxygen scatterers in the coordination sphere. Hence, from EXAFS data, it can be concluded that the Cu²⁺ site in Tau441 involves two or three His ligands.

The spectroscopic features (EPR and EXAFS spectra) of Cu²⁺ bound to Tau441 are strikingly similar to those reported for Cu²⁺ bound to human carbonic anhydrase II (HCA II),^{48–52} for

Table 2 EXAFS fitting parameters using a 3N1O or 4N coordination mode for the Cu²⁺-Tau441 complex^a

2N2O				3N1O			
Path	N	R (Å)	σ^2 (Å ²)	Path	N	R (Å)	σ^2 (Å ²)
Cu-N	2	1.92(1)	0.006(3)	Cu-N	3	2.02(2)	0.004(2)
Cu-O	2	2.00(1)	0.003(2)	Cu-O	1	1.88(3)	0.005(4)
Cu-C(His)	2	2.92(2)	0.003	Cu-C(His)	3	2.98(2)	0.003
Cu-C(His)	2	2.80(3)	0.003	Cu-C(His)	3	2.82(2)	0.003
Cu-N(His)	2	3.86	0.003	Cu-N(His)	3	3.79(5)	0.003
Cu-C(His)	2	3.74	0.003	Cu-C(His)	3	3.55(5)	0.008(4)
$S_0^2 = 0.9$, $E_0 = -6.03$ eV, $R_{\text{factor}}^b = 17.87\%$, BVS = 2.24				$S_0^2 = 0.85$, $E_0 = -4.47$ eV, $R_{\text{factor}}^b = 7.97\%$, BVS = 2.26			

^a Coordination numbers have been fixed. The number of long-distance Cu-C and Cu-N scattering paths was fixed to the number of His ligands. Numbers in brackets are the estimated uncertainties in the last digit. ^b R-Factor filtered between 1.1 and 3.1 Å in FT.



which X-ray crystallography indicates a binding site involving a bis-His motif, and possibly a third or fourth N or O ligand.⁵² Here, a comparison to HCA II might be useful, since the Cu²⁺ binding site in HCA II is highly sensitive to the substitution of the exogenous ligand. Cu²⁺ bound to HCA II displays an EPR signal with $g = 2.320$ and $A = 156 \times 10^{-4} \text{ cm}^{-1}$ that changes drastically upon binding of a sulphide ligand,⁴⁸ yielding EPR parameters that are quite far from those observed for Cu²⁺ bound to Tau441. This is consistent with ruling out a Cys residue acting as a fourth ligand in the Cu²⁺-Tau441 complex. However, the EXAFS spectrum of Cu²⁺ bound to HCA II was fitted with 2 Cu-N interactions at 1.98 Å that were assigned to the bis-His motif, and 2 Cu-N/O interactions at 1.88 Å; while multiple-scattering analysis shows that the second coordination sphere is best fit by considering 2 or 3 His ligands, with the participation of His64 and the bis-His motif of HCA II. Similarly, the inclusion of multiple-scattering from His ligands is needed to best fit the EXAFS spectrum of Cu²⁺ bound to Tau441, while the Cu-N distances are in the same range (Table 2). The striking similarity of the spectroscopic features of Cu²⁺ bound to HCA II and the Cu²⁺-Tau441 complex further supports the conclusion that a bis-His motif participates in copper binding to Tau441.

Together, EPR and XAS results indicate that Cu²⁺ bound to Tau441 is a covalent square planar type 2 site encompassing two or three His ligands in its coordination sphere. While it is not possible to discern between 2N and 3N equatorial coordination spheres, it is likely that the oxygen-based ligands would arise from Asp or Glu residues, while the participation of Cys ligation can be discarded. NMR results suggest that His299 from R2, and the bis-His motif (His329/His330) from R3 could be the nitrogen-based ligands in the Cu²⁺-Tau441 complex. NMR experiments also point to a possible participation of Asp295 or Asp314, which could act as oxygen-based ligands. Future studies using site-directed mutagenesis will be needed to probe the role of these residues in Cu²⁺ binding to Tau441.

Chemical reduction of the Cu²⁺-Tau441 complex

Given that the principal location of Tau441 is intracellular, where copper ions are in their reduced form, it would be of great interest to probe Cu⁺ binding to Tau441. For this purpose, chemical reduction of the Cu²⁺-Tau441 complex was attempted using ascorbic acid and/or dithiothreitol (DTT) as reducing agents (see text in the SI and Fig. S4).

Full reduction of the complex was achieved using 2 mM DTT and 50 mM ascorbic acid, assuring that the pH of the copper-protein solution was kept at 7.4. Spin quantitation of the EPR spectrum for the reduced sample indicated 2% of residual Cu²⁺ species (Fig. 5A), while the XANES spectrum displayed the typical feature at 8982 eV associated with the electric dipole-allowed 1s → 4p transition for Cu⁺ ions and its position and intensity are correlated with the coordination pattern of Cu⁺ (Fig. 5B).³⁷ The intensity and shape of the Cu⁺ feature peak suggest that the Cu⁺-Tau441 complex would be best represented as a tri-coordinated complex.^{37,53} The FT of the EXAFS spectrum for the Cu⁺-Tau complex is shifted towards longer metal-ligand distances with respect to those for the Cu²⁺-Tau

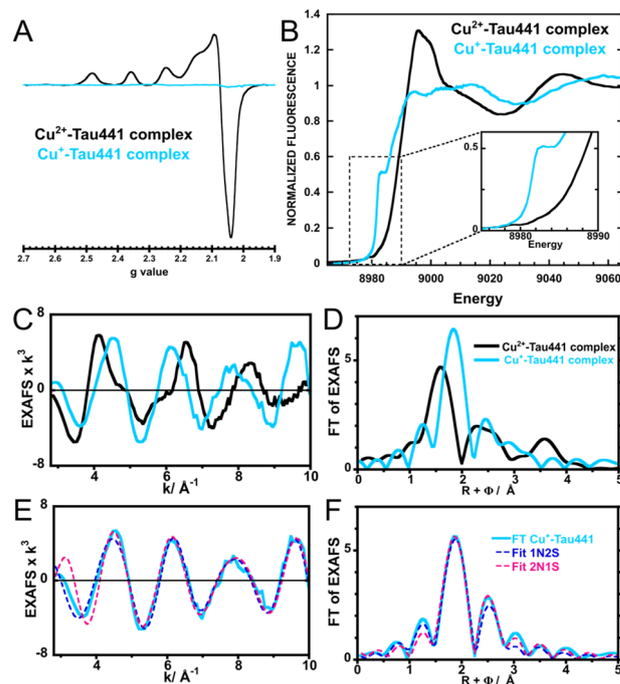


Fig. 5 Spectral comparison of Cu²⁺-Tau441 and Cu⁺-Tau441 complexes. EPR (A), Cu K-edge XANES (B), EXAFS (C) and FT (D) of Cu²⁺-Tau441 (black) and Cu⁺-Tau441 (light blue) complexes. The inset in (B) shows the feature at 8982 eV associated with the electric dipole-allowed 1s → 4p transition for Cu⁺ ions. (E) EXAFS fit and (F) FT for the spectrum of the Cu⁺-Tau441 complex with 1N2S (blue dotted line) and 2N1S (pink dotted line) coordination spheres using the parameters listed in Table 3.

complex (Fig. 5C and D), consistent with having a lower metal oxidation state. EXAFS fitting for the spectra of the Cu⁺-Tau complex was performed considering nitrogen and/or sulfur-based ligands (Table S2 and Fig. S5), revealing that the best results were achieved with a mixture of both types of ligands. The two possibilities for a tri-coordinate Cu⁺ complex were considered, *i.e.* 1N2S and 2N1S coordination spheres (Fig. 5E, F and Table 3), revealing that both models fit adequately (Fig. 5C and D), although BVS analysis for the 1N2S model yields a metal

Table 3 EXAFS fitting parameters using a 1N2S or 2N1S coordination sphere for the Cu⁺-Tau441 complex^a

Path	1N2S			2N1S		
	N	R (Å)	σ ² (Å ²)	N	R (Å)	σ ² (Å ²)
Cu-S	2	2.238(4)	0.003(1)	1	2.267(25)	0.006(8)
Cu-N	1	1.881(23)	0.005(3)	2	2.151(20)	0.002(2)
Cu-C(His)	2	2.714(2)	0.002	4	2.788(22)	0.002
Cu-C(Cys)	2	3.079(15)	0.002	1	3.063(28)	0.002
$S_0^2 = 0.95$, $E_0 = -4.47 \text{ eV}$, $R_{\text{factor}}^b = 13.26\%$, BVS = 1.11			$S_0^2 = 0.95$, $E_0 = 8.57 \text{ eV}$, $R_{\text{factor}}^b = 13.09\%$, BVS = 0.76			

^a Coordination numbers have been fixed. The number of paths of imidazole rings was fixed to the number of His ligands. Numbers in brackets are the estimated uncertainties in the last digit. ^b R-Factor filtered between 1.0 and 4.0 Å in FT.



formal charge (+1.13) that is closer to the true value, as compared to that of the 2N1S model. These fits (Table 3) yield Cu–S interactions in the range of 2.24–2.27 Å and Cu–N distances in the range of 1.88–2.15 Å; the former falls within the range for known Cu–S distances for Cys coordination in Cu⁺ metalloproteins, while the latter is consistent with distances reported for His ligation.⁴⁷ The second sphere contribution in the FT was best fit considering the remote carbons of Cys and His residues (Table 3). Overall, these results support a tri-coordinate model for the Cu⁺–Tau441 complex with one His and one Cys, while the third ligand could be a second His or Cys residue.

Preliminary NMR studies of the Cu⁺–Tau441 complex were performed using an excess of ascorbic acid (166 equiv.) (Fig. S6); under these conditions, 71% reduction of the complex was achieved, as assessed by EPR (Fig. S6B). Nevertheless, NMR analysis (Fig. S6C and D) reveals a distinct result with respect to that observed for the Cu²⁺–Tau441 complex (Fig. 3C and D). Cu⁺ is a diamagnetic metal ion, and hence, no signal broadening due to paramagnetic relaxation effects would be observed upon reduction of the Cu²⁺ complex; consistently, the *I/I*₀ shows very little changes (Fig. S6C). However, individual chemical shift perturbations show some changes in the MTBD region; specifically at residues: I278, I297, Q307, V318, G326, I328 and H329 (Fig. S6D). The most affected residues in the R2/R3 region are I297 and Q307, which flank His299, a residue that also acts as a ligand for the Cu²⁺–Tau441 complex; hence His299 is likely to act as a ligand in the Cu⁺–Tau441 complex. Although the NMR signals for the two Cys residues of Tau441 (Cys291 and Cys322) cannot be discerned, chemical shift perturbations around Cys322 are quite prominent, while residues around Cys291, such as G292, S293, K294 and N296, are not perturbed (Fig. S6D); hence Cys322 is more likely to be acting as a ligand in the Cu⁺–Tau441 complex than Cys291. Additional chemical shift perturbations in all regions might reflect conformational changes caused by metal binding to Tau441.

Altogether, XAS and NMR results indicate that Tau441 can bind Cu⁺ ions and that the Cu⁺–Tau441 complex is tri-coordinate with one His and one Cys, likely His299 and Cys322, while the third ligand could be Cys291 or a His from the bis-His motif. Future studies using site-directed mutagenesis and direct Cu⁺ binding assays with Tau441 are needed to further elucidate the nature of the Cu⁺ binding site in Tau441.

Tau441 has multiple Zn²⁺ binding sites

Zn²⁺ binding to Tau441 was probed at single-residue resolution by NMR, by titrating the protein with different equivalents of Zn²⁺, followed by ¹H–¹⁵N HMQC spectroscopy (Fig. 6A). Minor changes are observed upon addition of 0.1 equiv. of the metal ion (Fig. 6A, left panel), while larger changes in different regions of the NMR spectra become evident upon addition of 0.3 equiv. (Fig. 6A, middle panel), including the Gly (¹⁵N chemical shifts of 105–112 ppm) and His (¹⁵N chemical shifts of 118–124 ppm) regions. Finally, after addition of 1.5 equiv. of Zn²⁺, most protein residue signals are drastically perturbed (Fig. 6A, right panel). An analysis of individual chemical shift perturbations (CSPs) shows

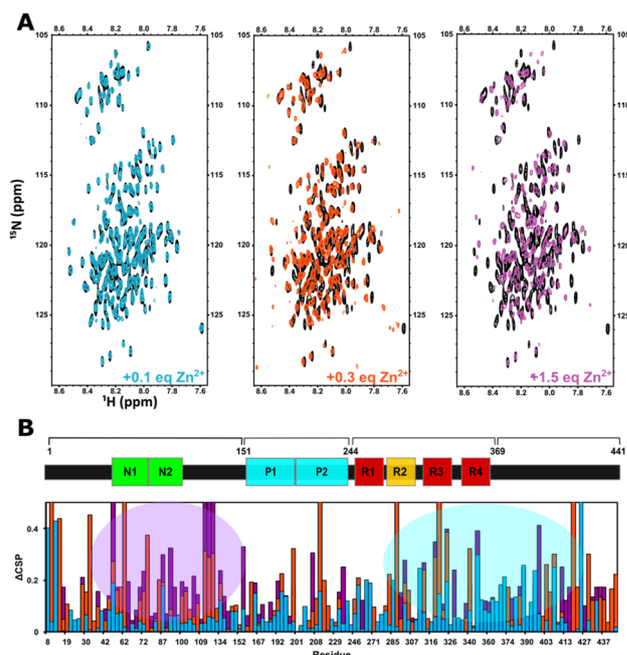


Fig. 6 Domains involved in the Zn²⁺ coordination sites. (A) Overlays of ¹H–¹⁵N HMQC NMR spectra of Tau441 100 μM in the absence (black) and in the presence of 0.1 equiv. (light blue), 0.3 equiv. (orange) and 1.5 equiv. (purple) of Zn²⁺; (B) analysis of the chemical shift perturbations (CSPs) for the HMQC ¹H–¹⁵N signals of assigned residues of tau with 0.1 equiv. (blue), 0.3 equiv. (orange) and 1.5 equiv. (purple) of Zn²⁺. Blue circles highlight perturbed residues with 0.1 equiv. of zinc and purple circles highlight residues perturbed with 1.5 equiv. of zinc.

that the addition of 0.1 equiv. of the metal ion impacts the C-terminal and the microtubule binding domain (light blue region in Fig. 6B), while the rest of the protein remains mostly unchanged.

At 0.3 equiv., the changes at the C-terminal and the MTBD become even more evident, while large perturbations are also observed at the N-terminal region, encompassing residues 43–135 that include His121 (orange bars in Fig. 6B). A couple of residues in the proline-rich region 210–230 known to form a transient secondary structure,^{26,39} are also affected upon addition of 0.3 equiv. of Zn; however, they are unlikely to participate directly in metal binding and the CSP effects may reflect an indirect effect of Zn binding to the local structure. Finally, the N-terminal residues are the most affected region upon addition of 1.5 equiv. of Zn²⁺ (purple bars and region in Fig. 6B).

A more detailed analysis of the NMR spectra reveals that at 0.1 equiv. of Zn²⁺ the most affected residues are Gly120 (Fig. 7A), His94 (Fig. 7B), Ala119, His121, Ile297, Val318 and Ile328 (Fig. 7C). These changes are mapped onto the protein sequence in Scheme 2 (in light blue color), where it is evident that His94 and Ala119/Gly120/His121 flank a region that is rich in Glu residues, which together with His94 and His121 could act as ligands for Zn²⁺. However, Ile297 is in the R2 domain, within the sequence DNIKH, that contains Asp295 and His299, which



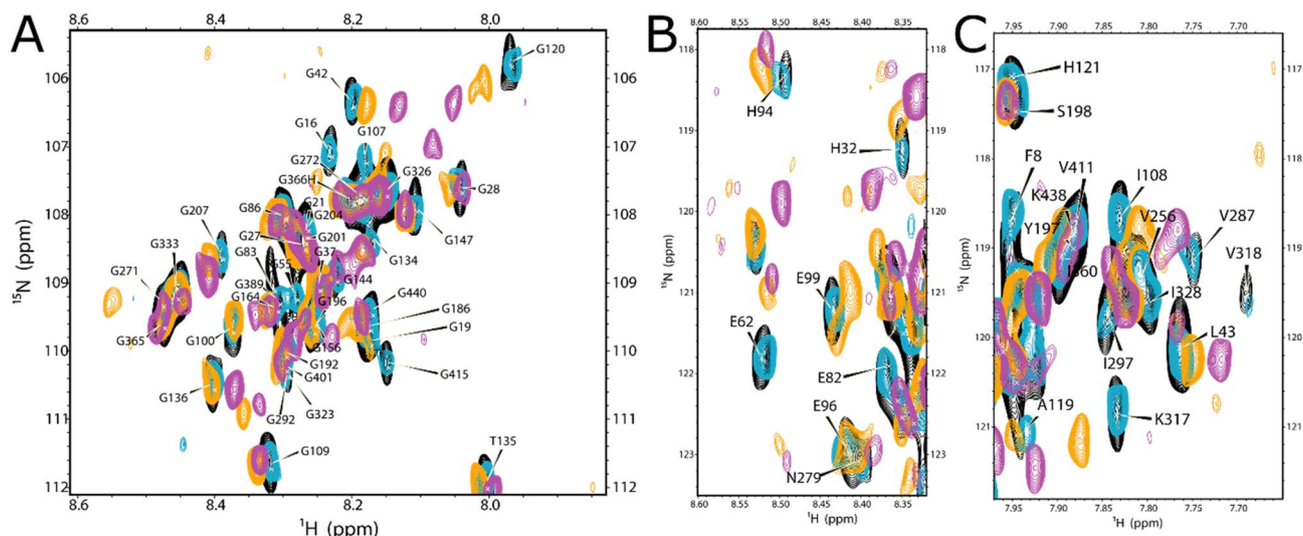
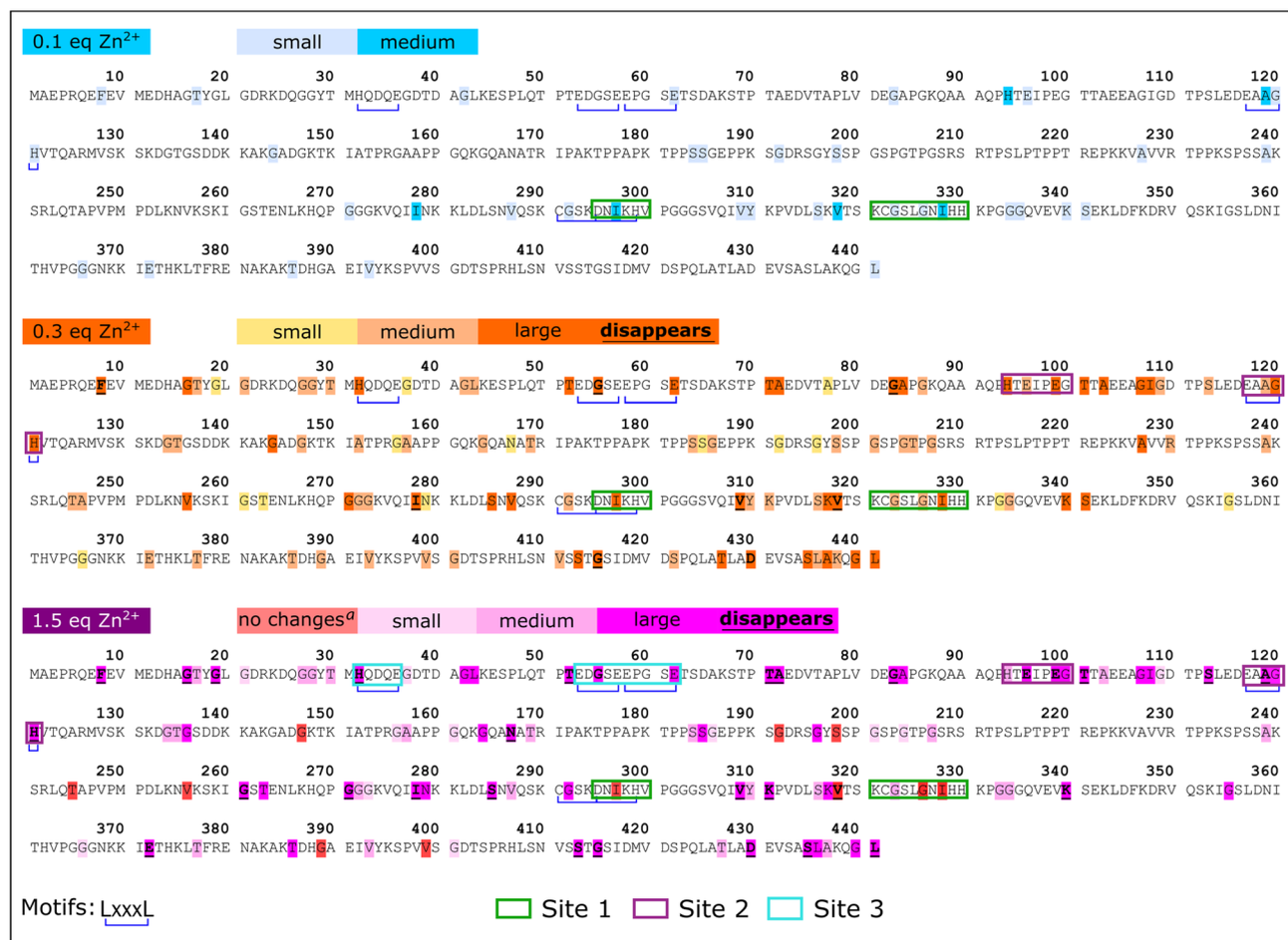


Fig. 7 Residues involved in the Zn^{2+} coordination sites. A zoomed image of the Gly (A), His (B) and Ile-Val (C) regions of ^1H - ^{15}N HMQC overlaid spectra of Tau441 (black) with 0.1 equiv. (blue), 0.3 equiv. (orange) and 1.5 equiv. (purple) of Zn^{2+} .



Scheme 2 Mapping of the NMR perturbations of Tau441 with Zn^{2+} . Mapping of the chemical shift perturbations of the sequence of tau protein with 0.1 equiv. (blue), 0.3 equiv. (orange) and 1.5 equiv. (magenta) of Zn^{2+} . ^aNo changes in the CS between 0.3 and 1.5 equiv. of Zn^{2+} . Motifs LxxxL are underlined in blue. Coordination sites are highlighted in green (site 1), purple (site 2) and light blue (site 3).



could also act as Zn anchoring ligands. Finally, Val318 and Ile328 are in the R3 region, near the bis-His motif (His329/His330), which is also known to bind zinc. It should be noted that all these residues are further perturbed upon addition of 0.3 equiv. of Zn^{2+} (Scheme 2, in orange), where the most drastic changes include the disappearance of the Val318 signal and large displacements of Ile297 and 328 (Fig. 7C), which are residues in the vicinity of the DNKIH motif in R2, Cys322 and the bis-His motif in R3. Since no further changes are observed in these residues at the R2/R3 region upon addition of 1.5 equiv. of Zn (light blue region in Fig. 6B), these anchoring residues might constitute a Zn binding site (labeled as site 1 in Scheme 2) that gets saturated at >1 equiv. of the metal ion. In contrast, drastic changes are observed for the Gly120 (Fig. 7A), His94 (Fig. 7B) and Ala119 (Fig. 7C) signals, which shift significantly at 0.3 equiv. and continue to change at 1.5 equiv. Moreover, the signals for Gly100 and Gly109 (Fig. 7A), Glu96 and Glu99 (Fig. 7B) shift significantly at 0.3 and 1.5 equiv. of zinc or even disappear, as is the case for Glu99 and Gly100. Overall, these results suggest that the region flanked by His94 and His121, containing glutamate residues like Glu96 and Glu99, may constitute a zinc anchoring site (labeled as site 2 in Scheme 2) that gets populated starting from 0.1 and up to 1.5 equiv. of the metal ion.

At 0.3 equiv. of Zn^{2+} , several additional residues at the N-terminal domain of the protein are affected, namely Gly55 (Fig. 7A) and Glu62 (Fig. 7B) disappear, while Gly42 (Fig. 7A), His32 (Fig. 7B) and Leu43 (Fig. 7C) show a large shift at 0.3 equiv., which further changes at 1.5 equiv. of zinc. Mapping these and other shifts onto the sequence of the protein (Scheme 2, orange and purple), it becomes evident that these changes occur in a region containing several putative ligands for zinc; for example, His32 is contained within the sequence HQDQEGDTD, while Glu62 is located in the EDGSEEPGSE motif, both containing several glutamates and aspartates that could act as anchoring ligands for zinc. This region likely constitutes a third zinc binding site of lower affinity, labeled as site 3 in Scheme 2.

In summary, NMR results reveal the presence of at least three putative binding sites for zinc in Tau441: site 1 likely involves the DNKIH motif in R2 containing Asp295 and His299, and Cys322 and the bis-His motif (His329/His330) in R3; site 2 encompasses the region flanked by His94 and His121 at the end of the N2 insert, containing several Glu residues like Glu96 and Glu99; while site 3 is located at the N-terminal domain, in the N1-insert, involving the HQDQEGDTD and EDGSEEPGSE motifs. It should be noted that all these sites contain motifs of the type LxxxL, where L is a putative ligand for Zn^{2+} (Scheme 2 blue underline); this type of zinc binding motif is known to exist in zinc binding proteins, such as zinc fingers and zinc transporters.⁵⁴ Examples of identified LxxxL motifs that may be involved in Zn^{2+} binding to Tau441 are DNKIH in site 1, EAAGH in site 2, and HQDQEGDTD or EDGSEEPGSE in site 3. Finally, it should be noted that several other regions of the protein are affected upon zinc binding that are not located in the vicinity of residues that could act as zinc ligands (Scheme 2), but that

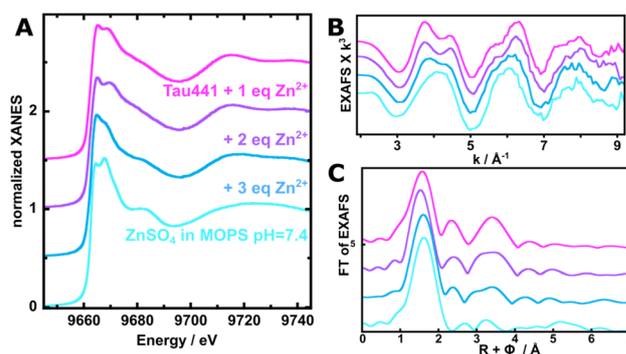


Fig. 8 Tau441– Zn^{2+} binding sites at different equivalents. Titration of Tau441 protein (0.8 mM) in 20 mM MOPS buffer at pH 7.4, with Zn^{2+} (0 to 3 equiv.), followed by XAS. Comparison of the Zn K-edge normalized XANES (A), EXAFS (B), and FT (C) spectra of Tau441 with 1 (magenta), 2 (purple), and 3 (blue) equivalents of Zn^{2+} . The spectrum of $ZnSO_4$ in buffer (turquoise) is included for comparison.

might correspond to protein regions that suffer conformational changes upon metal binding.

To gain insights into the nature of the Zn^{2+} binding sites of Tau441, Zn K-edge XANES (Fig. 8A) and EXAFS (Fig. 8B and C) spectra were collected for the protein with 1 equiv. (magenta), 2 equiv. (purple) and 3 equiv. (dark blue) of the metal ion. The spectrum for Zn^{2+} in MOPS buffer was collected as a control (light blue traces in Fig. 8); the edge intensity in its XANES spectrum (Fig. 8A) is typical of an octahedral site of Zn^{2+} (a d^{10} configuration has no pre-edge feature, regardless of the geometry).⁵⁵ In contrast, the white line intensity and fine structure observed around 9670 eV in the XANES spectra for the protein with 1 and 2 equiv. of Zn^{2+} reveal that the protein bound Zn^{2+} species have a tetrahedral geometry, while the XANES spectrum for 3 equiv. shows a peak shoulder at 9682 eV that arises from a mixture of Zn^{2+} in buffer and Tau441-bound Zn^{2+} species (Fig. 8A). Consistently, the EXAFS spectra (Fig. 8B) for the protein with 1 and 2 equiv. of Zn^{2+} exhibit similar features, while that of 3 equiv. seems to be a combination of Tau441-bound Zn^{2+} species and free Zn^{2+} in buffer solution. Specifically, the double feature centered at 4 \AA^{-1} in the EXAFS spectra (Fig. 8B) is most evident at 1 and 2 equiv. of the metal ion; this feature is typical of Zn^{2+} bound to a bis-His motif, as observed previously for Zn^{2+} binding to the beta-amyloid peptide.⁵⁶ The FT of the EXAFS spectra for the protein with 1 and 2 equiv. of Zn^{2+} (Fig. 8C) shows slight differences in the short metal–ligand distance region, which could not be adequately modelled by the fits. The best EXAFS fits for Tau441 with 1 and 2 equiv. of zinc were achieved assuming a mixture of species: one with the 2N1S1O coordination sphere (site 1) and a second species (site 2) with 2N2O (Fig. S7 and Table S3); however, determining the percentage of each of these two species is difficult, since a variation in the percentage of the species with a sulfur ligand is compensated for by the metal–ligand distances. A BVS analysis⁴⁵ (Table S3) reveals that the fits that do not consider any contribution from a Zn–S interaction yield a formal charge for the metal center that is significantly below +2.0 (+1.47 or +1.51, Table S3), supporting the hypothesis that there might be a Cys



ligand at site 1. However, the EXAFS fits that yield a formal charge of the metal center closer to +2.0 are those having an average coordination number of 0.3 for the S ligand, *i.e.* having 30% of the 2N1S1O species (site 1), with Zn–S distances in the range of 2.25–2.28 Å. The Zn–O interactions at 2.12 Å are likely due to Asp or Glu residues, while Zn–N interactions at 1.96 Å likely correspond to His ligands; indeed, scattering at larger distances can be fitted by including carbon and nitrogen atoms of an imidazole ring (Table S3). These results are consistent with reported Zn–S distances from Cys coordination, which are greater with respect to Zn–N or Zn–O distances arising from His or Glu ligation, respectively.³⁷ Hence, XAS results would be consistent with having a coordination sphere for site 2 with two His and two carboxylate residues, while site 1 likely involves two His residues, one carboxylate and one Cys; the latter could be either Cys291 from R2 or Cys322 from R3, both regions being highly affected by zinc, as observed by NMR (Scheme 2). However, better characterization of the Zn binding sites in Tau441 would be achieved by site-directed mutagenesis and/or by studying fragments or mutants that could help isolate the different Zn species.

In summary, a combination of NMR and XAS studies reveals that Tau441 is likely to have up to three tetrahedral binding sites for Zn²⁺, one of them in the R2/R3 region likely encompasses His299 from R2, the bis-His motif (His329/His330) from R3, and Cys291 or Cys322. The other Zn²⁺ binding sites are located in the N-terminal domain, and they seem to involve mostly His, Asp and Glu residues in LxxxL motifs for zinc binding (Scheme 2).

Cu²⁺ and Zn²⁺ ions impact the aggregation of Tau441 protein

The impact of Cu²⁺ and Zn²⁺ ions on the amyloid aggregation of Tau441 protein was evaluated by Thioflavin T (ThT) fluorescence assays. ThT is a dye whose fluorescence emission centered at 482 nm increases upon intercalation into the β -sheet structure of amyloid aggregates; hence, an increase in the fluorescence intensity reports on the growth of amyloid fibrils.⁵⁸ Heparin was added in a 1 : 4 ratio to achieve Tau441 amyloid aggregation within 20 hours; control experiments with Tau441 without heparin do not display any aggregation in this time frame (Fig. S8). In the absence of metal ions, ThT fluorescence increased significantly over the course of 20 h, displaying a lag phase of 1.2 hours before a sigmoidal growth of Tau441 amyloid fibrils (Fig. 9A and D, black trace). Transmission electron microscopy (TEM) analysis at the end point of the assay shows the formation of long helical Tau441 aggregates (Fig. 9B), as previously observed.²¹ In the presence of 1 equiv. of Cu²⁺ ions, the increase of ThT fluorescence is accelerated and no lag time is observed (Fig. 9A, light blue trace), suggesting that the metal ion accelerates the first steps of amyloid aggregation. Moreover, the ThT trace in the presence of 1 equiv. of Cu²⁺ displays exponential growth of Tau441 amyloid fibrils at a rate that is approximately four times higher than in the absence of metal ions (Fig. S9). TEM analysis at the end point shows the formation of long and intercrossed Tau441 fibrillar aggregates that seem to be less helical than those in the absence of copper (Fig. 9B). Upon addition of 2 or 4 equiv. of Cu²⁺ ions, the exponential growth behavior is conserved; however the maximum intensity of ThT fluorescence is severely diminished

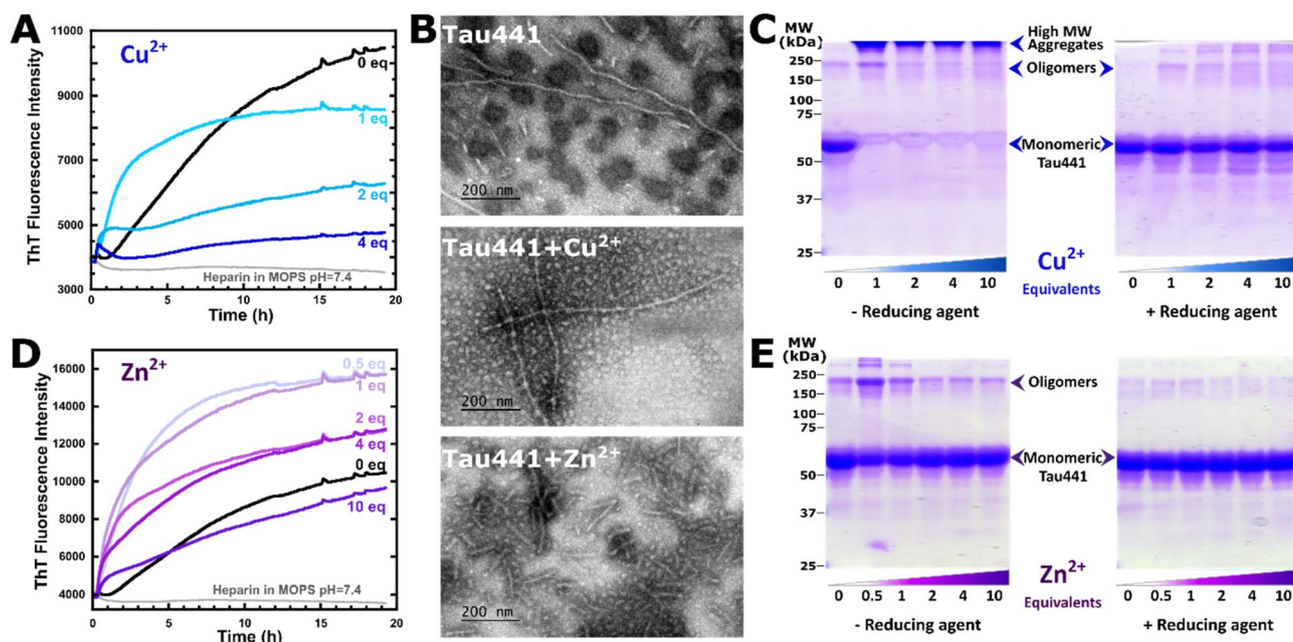


Fig. 9 Cu²⁺ and Zn²⁺ ions impact the aggregation of Tau441 protein. Aggregation assays of Tau441 protein (25 μ M) in 20 mM MOPS buffer at pH 7.4, in the presence of heparin (6.25 μ M), and different equivalents of Cu²⁺ (A) or Zn²⁺ (D) ions, followed by ThT fluorescence at 37 °C. SDS-PAGE analysis of the samples at the endpoint (20 h) of the ThT aggregation assays (A and D) with Cu²⁺ (B) or Zn²⁺ (E) ions, performed in the presence (right) or absence (left) of 2-mercaptoethanol (BME) as the reducing agent. (C) Electron microscopy images of the amyloid aggregates of Tau441 protein, obtained at the endpoint (20 h) of the ThT aggregation assays, in the absence of metal ions (top panel), and with 1 equiv. of Cu²⁺ (mid panel) or Zn²⁺ (bottom panel).



(Fig. 9A, darker blue traces). Being a paramagnetic metal ion, Cu^{2+} is known to quench fluorescence if bound in the vicinity; hence, decreased ThT fluorescence does not necessarily translate into less aggregation. Indeed, when Cu^{2+} ions are added to Tau441 fibrils that were grown in the absence of metal ions, the ThT fluorescence of the fibrils is quenched significantly, while TEM still shows the presence of amyloid aggregates (Fig. S10).

SDS-PAGE analysis of Tau441 aggregates at the end point of the ThT fluorescence assay was performed. Analysis of Tau441 amyloid aggregates in the absence of metal ions reveals mostly monomeric species (band at around 60 kDa) and a small amount of oligomers (band between 150 and 250 kDa) (Fig. 9C, left). Tau441 aggregates formed in the presence of copper ions show mostly high molecular weight species (band at $\text{MW} > 250$ kDa) and almost no monomers, while the band associated with low molecular weight oligomers is more evident at 1 equiv. of Cu^{2+} . These bands are no longer observed when the SDS-PAGE analysis is performed in the presence of a reducing agent, such as β -mercaptoethanol (BME) (Fig. 9C, right), suggesting that these species are linked *via* disulfide bridges.

Similarly to the case of Cu^{2+} ions, Tau441 aggregation in the presence of Zn^{2+} ions shows increased ThT fluorescence with no lag time (Fig. 9D, light purple trace), suggesting that Zn^{2+} ions also accelerate oligomerization. Moreover, the ThT traces in the presence of 0.5 and 1 equiv. of Zn^{2+} display exponential growth of Tau441 amyloid fibrils at a rate that is approximately four and three times higher than in the absence of metal ions, respectively (Fig. S9). Upon increasing the Zn^{2+} equivalents,^{2,4,10} the lag phase is still abolished, but the final ThT intensities decrease, and the aggregation kinetics show biphasic behavior (Fig. 9D, dark purple traces). It should be noted that the final ThT fluorescence with 0.5 and 1 equiv. of Zn^{2+} is significantly higher than that of Tau441 with no metal ions, suggesting the formation of more amyloid aggregates. However, protein quantification of soluble tau in the supernatant of the aggregation reactions reveals similar residual amounts of soluble Tau441 (about 10%) regardless of the presence of Zn^{2+} ions. TEM analysis at the end point with 1 equiv. of Zn^{2+} shows the formation of abundant and shorter Tau441 fibrillar aggregates than those in the absence of metal ions (Fig. 9B). Thus, the increased ThT fluorescence observed in the presence of zinc might be due to a larger fluorescence response of ThT to the short fibrils observed by TEM, which are small straight filaments and lack the helicity of Tau441 fibrils with no metal ions.

SDS-PAGE analysis of Tau441 aggregates in the presence of Zn^{2+} shows the presence of mostly monomers and a small amount of oligomers (Fig. 9E). The band associated with low molecular oligomers (between 150 and 250 kDa) peaks in intensity at 0.5 equiv. of Zn^{2+} (Fig. 9E, left). All oligomeric species are no longer observed when the SDS-PAGE analysis is performed under reducing conditions (Fig. 9C, right), suggesting that these species are linked *via* disulfide bridges. In contrast to the case of copper, no significant amount of high molecular weight species (above 250 kDa) is observed in the aggregates grown with Zn^{2+} , the low molecular oligomer species seems to be more favored, and the aggregate is mostly composed of monomers, regardless of the amount of metal ions

added (band between 50 and 75 kDa). These observations point to a metal-bridging mechanism, where monomers and some trimers could be bridged by Zn^{2+} ions.

Overall, these results suggest that both Cu^{2+} and Zn^{2+} ions accelerate the amyloid aggregation of Tau441, while having distinct effects on the morphology of the aggregates. Copper-induced aggregation of Tau441 yields amyloid fibrils that are intercrossed and less helical than those of the protein with no metal ions, while Zn^{2+} ions favor the formation of short fibrils.

Discussion

Copper and zinc are essential metals that perform important functions in the central nervous system. In this study, we demonstrate that Tau441 can bind Cu^{2+} , Cu^+ and Zn^{2+} ions, at the microtubule binding domain (MTBD); while Zn^{2+} can also bind at the N-terminal region (Fig. 10A). The MTBD region provides residues that act as anchoring ligands for metal ions: His299 and Cys291 from R2, Cys322 and a bis-His motif in R3. Spectroscopic results indicate that, while the Cu^{2+} site has a square planar geometry coordinated to two or three His residues (Fig. 10B), the reduced Cu^+ site is tri-coordinated involving His299, Cys322 and a third ligand, likely Cys291 or His329/330 (Fig. 10C), and the Zn^{2+} site is tetrahedral and likely also includes a Cys and an Asp residue in its coordination sphere (Fig. 10D). Structural models for these metal binding sites, as built by AlphaFold 3.0,⁵⁹ confirm some of these features, depicting a Cu^{2+} ion bound to His299 and His329/His330 from the bis-His motif, a Cu^+ ion coordinated to His299, Cys322 and His 330, and a Zn^{2+} binding site with His299, His330 and Cys322. Although having distinct geometries, the copper and zinc binding sites in Tau441 share some His and Cys ligands; thus, these metal ions could be competing for binding to the MTBD region. It is interesting to note that several proteins related to AD display sites with the ability to bind Cu^{2+} , Cu^+ and Zn^{2+} ions; these include the highest affinity copper-binding site (component 3) in the octarepeat region of the prion protein (PrP),^{60,61} the E2 domain site of the amyloid precursor protein (APP)⁶²⁻⁶⁴ and the N-terminal region of the A β peptide.¹⁷ This feature, together with the fact that the homeostasis of these metal ions is affected in AD,^{3,4} strongly suggests an important interplay of copper and zinc in this neurodegenerative disease that remains to be elucidated. The structural features of the metal binding sites in Tau441 and their potential physiological and pathological implications are discussed below.

Copper binding properties of Tau441

Tau441 displays a single Cu^{2+} binding site. Spectroscopic comparison of Cu^{2+} bound to TauK18 shows that it reproduces the metal site in the full-length Tau441, consistent with having a coordination sphere provided only by the MTBD. In contrast, the spectroscopic features of Cu^{2+} bound to peptide fragments modelling R2 and R3 and those reported for the separate R1, R2, R3 and R4 repeats^{28,65} cannot replicate the metal coordination site of the full-length protein. The fact that metal binding properties of tau peptide fragments are different from those of Tau441



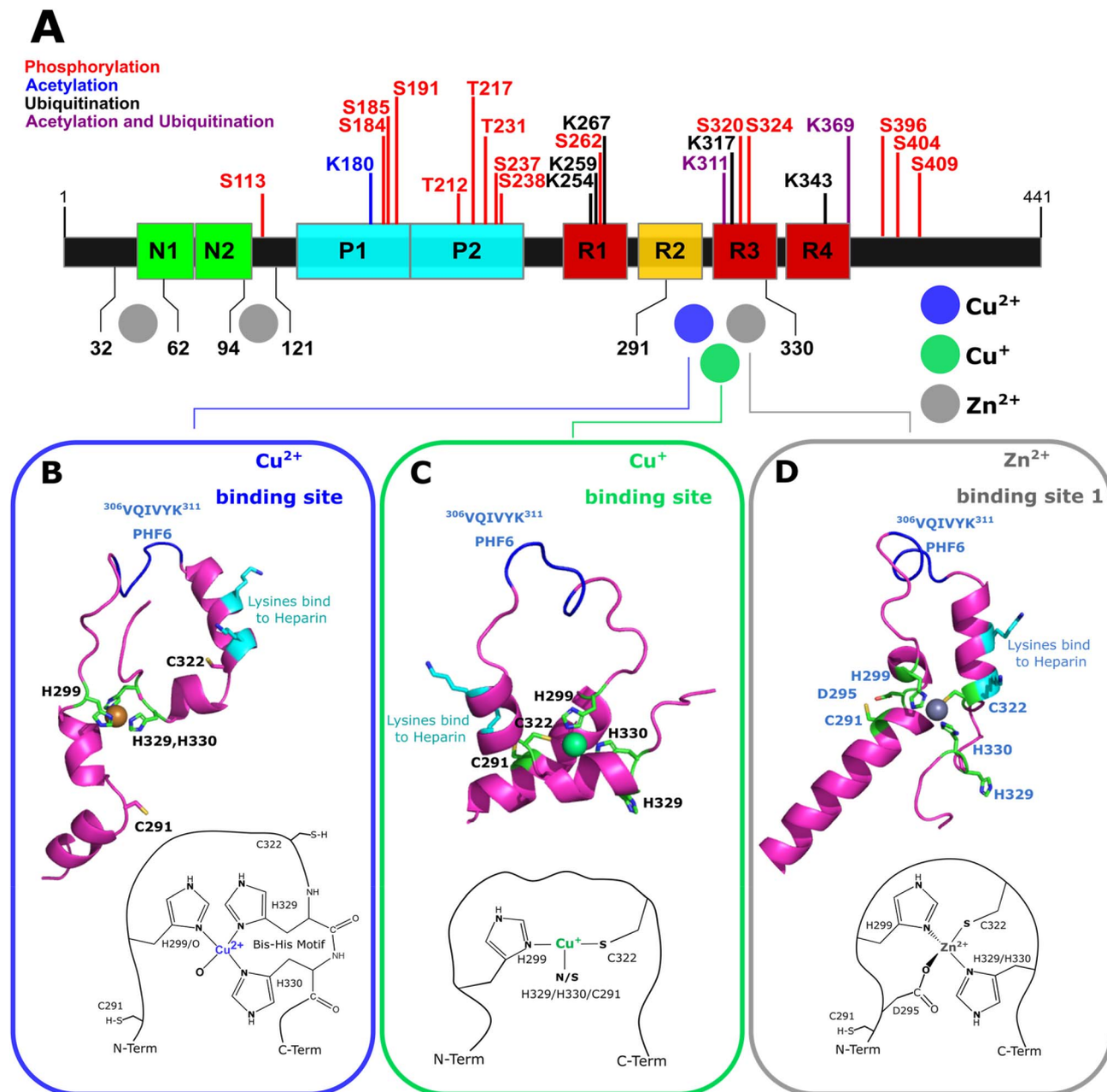


Fig. 10 Metal Binding properties of Tau441 and AD-related post-translational modifications. (A) Cu^{2+} , Cu^{+} and Zn^{2+} coordination sites in Tau441 are identified by blue, green and gray circles, respectively. The location of Tau441 AD related PTMs is shown above: phosphorylation (red), acetylation (blue) and ubiquitination (black) sites. Lys residues that undergo both ubiquitination and acetylation are marked in purple. Coordination modes (derived from spectroscopic results) and structural models generated by AlphaFold 3.0 for the Cu^{2+} (B), Cu^{+} (C) and Zn^{2+} (site 1) (D) binding sites in Tau441 are presented. For the AlphaFold 3.0 models, only the R2 and R3 repeats were used, putative metal binding residues are shown in green, the PHF6 sequence is in dark blue, and Lys residues that interact with heparin are in cyan.

underscores the importance of studying metal binding to the full protein and to physiologically and pathologically relevant forms of tau.

Having a multi-His Cu^{2+} binding site, Tau441 would be expected to have a higher binding affinity (with a K_d in the low nM range), similar to other multi-His Cu^{2+} sites, such as: the component 3 in the PrP with a $K_d = 10$ nM,⁶⁰ and the E2 domain of the amyloid precursor protein (APP) with a $K_d = 28$ nM,⁶² both proteins displaying four-His Cu^{2+} sites. However, ITC studies

reveal a $K_d = 0.5$ μM for Cu^{2+} binding to Tau441.²⁹ The fact that Tau441 is an intrinsically disordered protein and that the bis-His motif and His299 are 30 residues apart, may explain its $10\times$ lower Cu^{2+} binding affinity as compared to those systems. While the His residues acting as ligands for Cu^{2+} in the E2 site of APP are also many residues apart (>50), they are provided by three alpha helix bundles that impose a specific geometry for a tetragonal copper site.⁶³ However, while the multi-His Cu site in the PrP is located in its intrinsically disordered octarepeat



region, each His ligand is provided by an octapeptide with sequence PHGGGWGQ,⁶⁰ where the Pro residues play an important role in facilitating a protein conformation that is amenable to the coordination of the four His residues to copper. Moreover, having the His residues supplied by the same protein chain and spaced by only 8 residues provides an important chelate effect to achieve high Cu binding affinity. It is important to note that, in the MTBD domain of Tau441, the four repeats are each linked by a PGGG motif, similar to that of the PrP repeats. Specifically, the PGGG motif present between the R2 and R3 repeats may provide structural flexibility for copper binding to His299 in R2 and the bis-His motif in R3; however, the distance between these His ligands is greater (30 residues) than that between the His ligands in component 3 of the PrP (8 residues), which would contribute to the lower Cu binding affinity displayed by Tau441.

Beyond metal binding affinity, the fact that Tau441 displays a multi-His Cu²⁺ binding site warrants a comparison with other proteins containing multi-His Cu centers in terms of biological relevance and function. In the case of PrP, component 3 is responsible for stabilizing the *cis* conformation of the protein that is key to its neuroprotective role.^{60,66} For APP, copper binding to the E2 domain increases the thermal stability (T_m) of its α -helical structure.⁶³ A multi-His Cu site found in the S100A12, a protein with a prominent role in regulating inflammatory processes and immune responses,⁶⁷ appears to play a role in stabilizing the multimeric structure of the protein.⁶⁸ The phenoxazinone synthase (PHS), a protein naturally present in *Streptomyces antibioticus* bacteria,⁶⁹ has up to five Cu coordination sites, and one of them is a multi-His type 2 Cu site that plays a role in stabilizing the hexameric form of the protein to achieve greater catalytic activity.⁷⁰ In all these systems, the multi-His Cu²⁺ sites appear to play a structural stabilizing role in the protein. Recent reports indicate the presence of extracellular tau under both physiological and pathological conditions.^{23,25,71,72} Given that the oxidation state of copper in the extracellular milieu is primarily +2, Cu²⁺ binding to Tau441 could stabilize a protein conformation that might be relevant to its function in the extracellular environment. For example, extracellular tau has been shown to interact with membranes, and R2 and R3 appear to be the key domains for this interaction, particularly involving hydrophobic residues located between His299 and His329;⁷³ since copper binding occurs in this region, copper binding to His299 and the bis-His motif could be favoring a protein conformation where such hydrophobic residues located between these His ligands are more readily available for interaction with the lipid membrane. Indeed, NMR studies found that Cu²⁺ binding to TauK32 impacts the region encompassing such hydrophobic residues.²⁹ If this were the case, Cu binding to the MTBD of Tau441 would be playing a structural role in facilitating the interaction of tau with lipid membranes.

However, bis-His motifs are known to facilitate the formation of Cu complexes with a labile coordination position. For example, the CuB site in cytochrome *c* oxidase (CoO),⁷⁴ which activates oxygen in conjunction with an Fe heme a site, involves a bis-His motif and a nearby His residue, leaving a vacant position for O₂ binding. Similarly, the bis-His site in HCA II also displays a fourth

labile ligand that can accommodate an exogenous ligand.⁴⁹ This raises the possibility that Cu²⁺ bound to the bis-His motif and His299 in tau may also display an open coordination position for exogenous ligand binding or for the formation of ternary complexes with proteins that could provide a fourth ligand. Tau is known to interact with other His-containing proteins, such as PrP and A β .^{75–77} Hence, it would not be unreasonable to propose that such protein–protein interactions may be mediated by Cu binding through the formation of ternary complexes, as it has been demonstrated in the case of PrP and A β .^{78,79}

Cu⁺ interaction with Tau441 was studied by chemically reducing the Cu²⁺–Tau441 complex. Upon reduction, Cu⁺ remains bound to the MTBD using His299, Cys322 and a third ligand that can be a His or a Cys residue from the R2 or R3 domain. One of the key features of the Cu²⁺ binding site of Tau441 is that it contains a bis-His motif, which is known to stabilize Cu⁺ binding⁸⁰ and is present in several important copper binding proteins, including the copper transporter Ctr1p and the A β peptide.^{17,81} In the case of Ctr1p, the bis-His motif is thought to bind Cu⁺ and to be auxiliary to its high affinity Cu²⁺ ATCUN binding site (Amino-Terminal site of Copper and Nickel),⁸¹ while in A β , the bis-His motif is thought to play a role in both Cu²⁺ and Cu⁺ binding. The bis-His (His13/14) of A β , together with His6, can yield a multi-His Cu²⁺ binding site that displays a lower binding affinity, as compared to that of the main N-terminal site;¹⁷ while Cu⁺ binding to the bis-His motif is known to yield a linear bi-coordinate Cu⁺ site.⁸¹ In contrast, this does not seem to be the case for Tau441, where the bis-His motif (His329/330) participates as a bidentate ligand for Cu²⁺ binding but apparently not for Cu⁺. This is possibly due to the presence of nearby Cys residues that can coordinate to the Cu⁺ ion; hence, only one or two His residues remain coordinated to the metal ion upon reduction.

Given that ascorbic acid alone does not completely reduce the Cu²⁺–Tau441 complex, its reduction potential must be lower than that of ascorbic acid ($E^\circ < 52$ mV). The reduction potential of the Cu²⁺–Tau441 complex can be thermodynamically related to the reduction potential of free copper ions and the relative binding affinity of the protein for Cu²⁺ and Cu⁺ ions (Scheme S1); from this thermodynamic cycle, a lower limit for the dissociation constant of the Cu⁺–Tau441 complex can be estimated to be $K_d > 32$ μ M. This Cu⁺ binding affinity is strikingly low, considering the proposed coordination sphere for the Cu⁺–Tau441 complex. Indeed, Cu⁺ binding sites with two Cys and one His, as those found in cytosolic copper chaperones Sco1 and Sco2,^{82,83} display K_d values in the pM to fM range. Tri-coordinate type 1 sites with two His and one Cys display K_d in the pM range,^{82,84,85} while *de novo* designed sites with the same coordination set have K_d values in the fM range.⁸⁶ However, it should be considered that these metalloproteins contain Cu⁺ binding motifs, such as CXXXC or CXXXH, in a well-structured region with a nearby His residue. In contrast, Tau441 is an intrinsically disordered protein and the residues that may be anchoring Cu⁺ are 8 to 30 amino acid residues apart, leading to a lower binding affinity for Cu⁺.



Zinc binding properties of Tau441

In contrast to copper, zinc binding to Tau441, not only happens in the MTBD region, but it also occurs in the N-terminal domain, specifically at the N1 and N2 inserts (Fig. 10A). The nature of the N-terminal sites is evidenced by NMR, allowing for the identification of zinc binding motifs of the type LxxxL, that provide coordinating residues for Zn^{2+} (Scheme 2) in a similar fashion to those found in zinc finger proteins. The N-terminal zinc binding sites (sites 2 and 3) likely involve LxxxL motifs with a combination of His and carboxylate (Glu or Asp) anchoring ligands. In contrast, the MTBD site (site 1) is the only zinc binding site in Tau441 that may have a contribution from a Cys ligand, involving also two His from the R2 and R3 repeats. Cys and His residues constitute the most frequent ligands in high affinity Zn^{2+} binding sites in metalloproteins,⁸⁷ while Asp and Glu constitute anchoring residues for sites with lower zinc binding affinity.⁵⁷ Hence, one would expect sites 2 and 3 to display a lower affinity for zinc than that of site 1. Consistently, ITC studies at different temperatures of Zn^{2+} binding to Tau441 indicate the presence of one high affinity zinc binding site ($K_d \sim 0.5 \mu M$), likely associated with site 1 in the MTBD region, and three additional zinc binding sites with lower affinity ($K_d \sim 170 \mu M$)³³ which might correspond to the more oxygen-rich complexes in sites 2 and 3. It is interesting to note that the N-terminal region of Tau441 is involved in its interaction with proteins involved in synaptic transmission, such as synapsin-1 and synaptotagmin-1, and with annexin A5, a calcium-dependent phospholipid binding protein.⁸⁸ Hence, the identification of zinc binding sites in the N-terminal domain of Tau441 prompts to question if zinc may be playing a role in modulating the interaction of tau with some of its protein partners where the interaction involves the N-terminal region.

The highest affinity Zn^{2+} binding site in Tau441 protein likely involves His299, His330 and Cys322 (Fig. 10C), all residues located within the MTBD; this suggests a physiological role for zinc binding in regulating the interaction of Tau441 with alpha- and beta-tubulin.²⁴ Indeed, a recent report suggests a role for zinc in the regulation of cargo transport by microtubules (MTs) and their binding to microtubule-associated proteins (MAPs), including tau protein.⁸⁹ The study by Qin *et al.* demonstrated that binding of tau, and other MAPs (such as DCX and MAP2C), to microtubules is disrupted by the addition of Zn^{2+} , due to a direct interaction of the metal ion with the MTs with a K_d in the low nanomolar range. Structural analysis and modelling identified a putative Zn^{2+} binding site in alpha-tubulin, potentially involving residues H192, E196, E417, E420, E423, and D424. The residue pairs His192/Glu196 and Glu420/Asp424 are part of LxxxL motifs that are provided by two different alpha-helices, making them amenable to a tetrahedral Zn coordination site. Interestingly, these carboxylic residues are also engaged in binding of tau, as they display electrostatic interactions with the Lys residues present in the MTBD. Tau441 is a zinc binding protein with a high nM affinity (500 nM), which is one order of magnitude lower than the binding affinity of MTs for Zn (low nM range);⁸⁹ hence, even though it would be tempting to propose that zinc binding to Tau441 may be playing

a role in the Zn-mediated disruption of the interaction of tau with MTs, this is likely not the case. However, a role for the Zn-Tau441 interaction in the assembly of MTs cannot be discarded, since it has been proposed that Cys291 and/or Cys322 may form disulfide bonds with Cys347 in alpha-tubulin and/or Cys131 in beta-tubulin that contribute to stabilizing the tau-MT interaction.⁹⁰ Given that Cys322 is a putative ligand for Zn^{2+} in Tau441, a regulatory role for zinc in the function of tau during microtubule assembly would be plausible.

Beyond its cytoplasmic localization and interaction with microtubules, tau protein is also known to localize to the nucleus where it can interact with DNA, histones, RNA and nuclear proteins, potentially playing a role in DNA stabilization, chromatin organization and gene expression.^{91,92} The repeats R1–R4 in the MTBD of tau are involved in its binding to DNA, as this region is rich in Lys and Arg residues, known to interact electrostatically with negatively charged DNA.^{93,94} The nucleus is a zinc-rich cellular compartment;⁹⁵ hence, the Zn binding site 1 in tau could be populated, potentially modulating its interaction with DNA. Indeed, it has been reported that the binding mode of tau with DNA changes in the presence of zinc, specifically, going from minor groove to major groove binding for the Zn-bound protein. Such an effect could reflect a change in the conformation of tau in its metal-bound form. While studying how the conformation of Tau441 would be impacted by zinc binding would require a significant investment in structural studies, an AlphaFold 3.0-built model of Zn-bound tau (Fig. 10B) displays the Lys residues in the R3 repeat aligned on one side of an alpha-helix, a conformation that would facilitate the electrostatic interaction with DNA.

Metal-tau interactions: aggregation, toxicity and pathology

Tauopathies are characterized by the accumulation of amyloid fibrils composed entirely of self-assembled tau protein, displaying distinct morphologies and fibrillar structures in various types of tauopathies.⁹⁶ In AD, the NFTs are composed of two types of fibrils: paired helical filaments (PHFs) and straight filaments (SFs), sharing a structural core consisting of residues V308–F378, located in the R3/R4 region.⁹⁷ The core is composed of eight β -sheets that include the hexapeptide motif VQIVYK (PHF6, residues 306–311) and is considered essential for tau self-assembly.⁹⁸ The PHF6 sequence is located in R3, between anchoring residues that bind copper and zinc (Fig. 10); hence, metal binding could promote a conformation that favors intermolecular interactions between PHF6 motifs, facilitating core formation. Indeed, copper- and zinc-induced Tau441 oligomerization was previously demonstrated by dynamic light scattering (DLS).^{29,32,33} Copper is known to promote fibril formation of R2 and R3 peptide fragments,²⁸ while Zn^{2+} binding to TauK18 accelerates its aggregation in the presence of heparin.³¹ However, the lack of structural insights into the metal binding properties of Tau441 has prevented an understanding of the mechanism by which metal ions impact tau aggregation.

Our study demonstrates that Cu^{2+} and Zn^{2+} ions accelerate the amyloid aggregation of Tau441 in the presence of heparin.



Structural models of copper and zinc bound to tau (Fig. 10) show that the Lys residues in R3 are aligned in an alpha helix motif, an arrangement that, upon interaction with heparin, a sulfated polysaccharide,⁹⁹ could facilitate the alignment of metal-bound monomers in spatial proximity along the polysaccharide chain. Such interactions would favor the formation of oligomers and protofibrils, which would explain the suppression of the lag phase observed in the metal-induced Tau441 aggregation assays. Tau441 amyloid fibers obtained in the presence of copper are intercrossed and less helical than those formed in the absence of metal ions, while zinc induces the formation of short fibrils. Such morphological differences could arise from the distinct copper and zinc coordination sites (Fig. 10). CryoEM analysis of AD amyloid fibrils indicates that Cys322 is part of the fibril core, possibly forming intermolecular disulfide bridges that facilitate growth along the fibril axis.⁹⁷ Zinc binding to Tau441 involves Cys322, and it is likely to interfere with intermolecular disulfide bridging (as observed by SDS-PAGE, Fig. 9E), preventing elongation of the fibril and yielding shorter fibrils, as observed by TEM analysis (Fig. 9B). In contrast, since copper does not coordinate Cys322, it would not be expected to interfere with intermolecular disulfide bridging and the extent of fibril growth; indeed, copper-induced aggregates have an important component of disulfide-bridged oligomers (Fig. 9C), and have similar length to amyloid fibrils grown in the absence of metal ions (Fig. 9B). This is consistent with a cryoEM study of the truncated tau dGAE (297–391), showing that aggregates grown in the presence of copper display intermolecular disulfide bridges using Cys322, while zinc does not.¹⁰⁰ However, cryoEM analysis of PHFs shows that His330 is facing inwards in the amyloid fibril core, while His329 is facing outwards to the solvent-exposed area of the fibril.⁹⁷ In monomeric Tau441, copper coordinates to both His residues (Fig. 10); hence, upon fibril formation the copper ion would have to release His330, and it would likely remain bound to His329 on the outer side of the fibril; in such an arrangement, copper would likely recruit other solvent-exposed ligands. A structural analysis of the fibril core region V308–F378,⁹⁷ reveals that beyond His329, His362, Glu338 and Glu342 are other putative metal anchoring ligands facing the outer side of the fibril core. Hence, a His329-anchored copper ion could recruit any of these ligands from another fibril to complete its coordination sphere, possibly disturbing the inter-fibril interactions that cause the characteristic helicity of PHFs. This would explain why copper-induced aggregates, while preserving their fibril length, are inter-crossed and display decreased helicity. Further characterization of metal-bound aggregates of Tau441 should help probe these hypotheses and gain insight into the impact of metals on amyloid fibril morphology.

Disturbed metal ion homeostasis and aberrant metal–protein interactions associated with aggregation are distinct features of several neurodegenerative disorders.⁵ In the case of AD, the homeostasis of copper, zinc and iron is perturbed, and these essential metals accumulate in the NFTs.^{14,15} Specifically, levels of labile copper (not bound to ceruloplasmin) are elevated in blood plasma, while the concentration of extracellular tau is increased by five-fold in cerebrospinal fluid (CSF) of AD patients,^{101–103}

raising the possibility for a potentially toxic extracellular tau–copper interaction that could lead to increased oligomerization and aggregation; although, further studies are needed to determine the pathological consequences of copper-bound tau. However, the role of zinc in pathological models of tau has been explored. A study using SH-SY5Y cells expressing TauΔK280, a pathological mutation associated with frontotemporal dementia (FTD-17), revealed that Zn²⁺ markedly enhances apoptosis and toxicity induced by ΔK280 fibrillation.³⁶ Furthermore, in a *Drosophila* tauopathy model with TauR406W, both increased dietary zinc and genetic manipulation of zinc transporters leading to increased metal uptake, which led to increased tau toxicity.³⁴ In both models, mutation of Cys291 and Cys322 to Ala abrogated the effects of zinc in tau toxicity, strongly supporting a pathological role for zinc–tau interactions.^{34,36}

Moreover, it is interesting to note that the 3xTg-AD mouse model, encompassing the P301L mutation in tau, displays disruption of copper and zinc homeostasis.¹⁰⁴ Pro301 is important for the conformational flexibility of the loop connecting His299 and the bis-His motif (329/330), which are important anchoring sites for both copper and zinc binding to Tau441. Hence, it is plausible that the ability of Tau441 to bind Cu and Zn could be impacted by the P301L mutation, contributing to metal dyshomeostasis in the 3xTg-AD model. Certainly, murine models of AD could represent an excellent platform to further explore the pathological consequences of metal–tau interactions.

Finally, tau protein undergoes several physiological and pathological post-translational modifications (PTMs),²³ including phosphorylation, acetylation and ubiquitination. Some of the PTMs of tau observed in AD patients¹⁰⁵ (Fig. 10A) occur in regions of the protein engaged in metal coordination and could impact the metal binding properties of tau. Specifically, PTMs at the R3 domain could be impacting copper and zinc binding to the MTBD, while phosphorylation at Ser113 could affect zinc binding at the N-terminal. Hence, there may be an important interplay between PTMs and metal binding to tau, as it has been observed for alpha-synuclein, the Parkinson's protein.¹⁰⁶ Further research is needed to elucidate the interplay between all these processes.

Concluding remarks

This spectroscopic study provides detailed structural characterization of the Cu²⁺, Cu⁺ and Zn²⁺ binding properties of Tau441 and explores the impact of metal binding on tau amyloid aggregation. A comparison of the metal binding properties of Tau441 with those of tau fragments underscores the importance of studying metal binding to full-length tau. Tau441 displays a single high-affinity Cu²⁺ binding site located in the microtubule-binding domain (MTBD), involving a bis-His motif in R3 (His329/His330) and His299 in R2 (Fig. 10). Reduction of the Cu²⁺–Tau441 complex yields a Cu⁺ site that is tri-coordinated involving His299, Cys322 and a third ligand, likely Cys291 or His329/330. Tau441 also exhibits multiple Zn²⁺ binding sites: a high-affinity site in the MTBD involving the bis-His motif, His299 and Cys322, and two lower-affinity sites in the N-terminal region (Fig. 10). The fact that both copper and zinc



bind to the MTBD region suggests a physiological role for metal binding in regulating tau interaction with tubulins and/or microtubule assembly that deserves further investigation.

Being an intrinsically disordered protein, metal binding to Tau441 could play a structural role, stabilizing specific conformations to facilitate tau protein–protein and/or protein–nucleic acid interactions that may be physiologically and/or pathologically relevant. In this study, it was found that both copper and zinc, can accelerate the amyloid aggregation of Tau441, promoting distinct fibrillar morphologies that are consistent with the metal coordination models derived from spectroscopic studies. These findings support a pathogenic role for aberrant metal–tau interactions that is consistent with the scenario observed in AD, which includes metal accumulation in NFTs, increased extracellular tau and altered metal homeostasis. Finally, the metal binding sites identified here are located close to important sites for posttranslational modifications, including phosphorylation and ubiquitination; hence, an important perspective is to study the potential interplay between metal binding to Tau441 and its PTMs, both in health and disease. The current study expands the bioinorganic facet of AD and other tauopathies, and it opens new avenues to further explore the nature of metal–tau interactions, while it underscores their importance as potential therapeutic targets in these neurodegenerative diseases.

Author contributions

G. U. J. R.: investigation, writing – original draft; X. S.: investigation, methodology; J. G.: investigation, conceptualization, methodology, funding acquisition, writing – review and editing; C. D. A.: conceptualization, methodology, funding acquisition, writing – review and editing; G. L.: investigation, writing – review and editing; M. N.: methodology, writing – review and editing; R. R.: resources, supervision, writing – review and editing; J. L.: conceptualization, methodology, funding acquisition, project administration, supervision, writing – review and editing; L. Q.: conceptualization, methodology, funding acquisition, project administration, supervision, writing – original draft, writing – review and editing.

Conflicts of interest

There are no conflicts to declare.

Data availability

The supporting data have been provided as part of the supplementary information (SI). Supplementary information: Materials and methods section; EPR spectra comparison of Tau441 and buffer MOPS with Cu^{2+} ; titration of Tau441 and different fragments with Cu^{2+} followed by CD and EPR; Peisach–Blumberg diagram of the EPR parameters for the different fragments of Tau441 with Cu^{2+} ; parameters of the EXAFS fits of the Cu^{2+} –Tau441 complex with different coordination spheres; reduction of the Cu^{2+} –Tau441 complex section; EPR, XANES and EXAFS spectra for reduced Cu^{2+} –Tau441 complexes; EXAFS fitting of

the Cu^{+} –Tau441 complex considering only first coordination sphere interactions; parameters for the EXAFS fits for the Cu^{+} –Tau441 complex considering only first coordination sphere interactions; ^1H – ^{15}N HMQC NMR spectra of the Cu^{+} –Tau441 complex; EXAFS fitting and FT of Tau441 with 1 and 2 equiv. of Zn^{2+} ; parameters of the best fits for Tau441 with 1 and 2 equiv. of Zn^{2+} ; comparison of the Tau441–metal ion aggregation assays with and without heparin; fitting curves of the aggregation kinetics of Tau441 with metal ions; quenching effect in ThT fluorescence intensity upon the addition of Cu^{2+} ; thermodynamic scheme for the reduction of copper–tau complexes. Spectroscopic data will be made available upon request from the authors. See DOI: <https://doi.org/10.1039/d5sc08604c>.

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