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Copper binding and redox chemistry of the Aβ16 peptide and its variants: insights into determinants of copper-dependent reactivity†

Nineveh Yako, Tessa R. Young, Jade M. Cottam Jones, Craig A. Hutton, Anthony G. Wedd and Zhiguang Xiao*‡

The metal-binding sites of Aβ peptides are dictated primarily by the coordination preferences of the metal ion. Consequently, Cu(I) is typically bound with two His ligands in a linear mode while Cu(II) forms a pseudo-square planar stereochemistry with the N-terminal amine nitrogen acting as an anchoring ligand. Several distinct combinations of other groups can act as co-ligands for Cu(II). A population of multiple binding modes is possible with the equilibrium position shifting sensitively with solution pH and the nature of the residues in the N-terminal region. This work examined the Cu(II) chemistry of the Aβ16 peptide and several variants that targeted these binding modes. The results are consistent with: (i) at pH ≈ 7.8, the square planar site in Cu(II)–Aβ16 consists primarily of a bidentate ligand provided by the carboxylate sidechain of Asp1 and the N-terminal amine supported by the imidazole sidechains of two His residues (designated here as component IA); it is in equilibrium with a less stable component IB in which the carboxylate ligand is substituted by the Asp1-Ala2 carbonyl oxygen. (ii) Both IA and IB convert to a common component II (apparent transition pH_B 7.8 for IA and ~ 6.5 for IB, respectively) featuring a tridentate ligand consisting of the N-terminal amine, the Asp1-Ala2 amide and the Ala2-Pro3 carbonyl; this stereochemistry is stabilized by two five-membered chelate rings. (iii) Component IA is stabilized for variant Aβ16-D1H, components I (both IA and IB) are imposed on Aβ16-A2P while the less stable IB is enforced on Aβ16-D1A (which is converted to component II at pH ~ 6.5); (iv) components IA and IB share two His ligands with Cu(I) and are more reactive in redox catalysis than component II that features a highly covalent and less reactive amide N→C ligand. The redox activity of IA is further enhanced for peptides with a His1 N-terminus that may act as a ligand for either Cu(I) or Cu(II) with lower re-organization energy required for redox-shuttling. This study provided insights into the determinants that regulate the reactivity of Cu–Aβ complexes.

Significance to metallomics
Copper binding to Aβ peptides has been subjected to recent intensive studies due to its possible relevance to the aetiology of Alzheimer’s disease. Multiple metal binding ligands are present in the soluble and unstructured N-terminal section of Aβ(x–16), where x > 1 indicates truncation. The nature of the copper binding modes is dictated by the coordination preference of the copper oxidation state and by the medium pH, as is the reactivity of the Cu–Aβ complexes. This study clarifies current controversy regarding several important Cu(n) binding modes and provides an improved understanding of the complex copper chemistry of Aβ peptides.

Introduction
Disruption of metal homeostasis has been linked to Alzheimer’s disease and undesirable interactions between Aβ peptides and copper ions may play roles in the promotion of toxic Aβ aggregation and/or in the catalytic production of reactive oxygen species (ROS). Monomeric Aβ peptides are flexible and unstructured and the copper binding sites and binding modes...
change dynamically with the oxidation state and pH as does the reactivity of the metal centres towards ligand substitution and redox catalysis.

Previous studies have demonstrated that Aβ peptides bind Cu(II) in dynamic linear two-coordinate modes utilizing the three different combinations of the available His-6, -13 and -14 sidechains (Fig. 1a). In contrast, they typically bind Cu(I) in a square planar-based stereochemistry. The N-terminal amine nitrogen acts as a primary anchoring ligand. Depending on the conditions, several distinct combinations of other groups can act as co-ligands. The speciation depends sensitively on the medium pH and on the nature of the available co-ligands including the His residues, the sidechain of Asp1 and the carbonyl oxygen or deprotonated amide nitrogen of the Asp1-Ala2 peptide link (Fig. 1b). Note the presence of stabilizing six- or five-membered chelate rings in each Cu(II) complex.

The three distinct complexes in Fig. 1b are designated as components IA, IB and II in this work. Notably, they each can incorporate different combinations of His ligands as they appear to be conformationally available. Structure IA has frequently been proposed as a coordination mode at lower pH. However, recently IB has been preferred, based mainly on EPR studies. A transition from components IA/IB to component II requires deprotonation of the peptide amide of Ala2 and is promoted by elevated pH. Any modification or truncation to the N-terminal residues, as occurs in native Aβ peptides, can alter the Cu(II) binding modes dramatically. For example, both the substitution of Asp1 with Asn and deletion of Asp1 to impose Ala2 as the N-terminal residue in the Aβ16 peptide lead to a decrease of over one pH unit in the pKₘₐₜ of transition I → II. Deletion of the first three residues, as detected in native Aβ peptides in high proportion, generates a classic N-terminal H₂N-X-X-His₃ motif known as the ATCUN (amino terminal copper and nickel) binding motif: the Cu(II) centre is stabilized by three chelate rings in a tetradeinate ligand (Fig. 2b). The different binding modes sensitively affect the reactivity of Cu-Aβ complexes towards ligand substitution and redox catalysis, especially for Cu(II). This study undertook a range of chemical and spectroscopic studies of the copper chemistry of the Aβ16 peptide to provide improved insight into the relationship between binding mode and reactivity. In particular, several variants that targeted the Cu(II) binding site specifically were examined. The work was facilitated by the recent development of a set of peptide-based fluorescent dansyl probes DP1-4 that are able to quantify Cu(II) binding stoichiometries and affinities from micromolar to femtomolar concentrations on a unified scale. The combined data demonstrated that (i) the Aβ16 peptide binds Cu(II) dominantly as component IA in equilibrium with less stable IB at low pH, while at higher pH, component II predominates (Fig. 1b); (ii) component I is more reactive than component II in redox catalysis.

### Experimental section

**Materials and general procedure**

Materials purchased from Sigma included the ligands Ferene S (Fs), Ferrozine (Fz) and bicinechoninic anion (Bca) (as their sodium salts Na₂Fs, NaHFz and Na₂Bca), and the reductants NH₂OH (as its H₂SO₄ salt) and ascorbic acid (Asc). They were all used as received. A standard Cu(II) solution was prepared by dissolving CuSO₄ in Milli-Q water and the concentration calibrated via reaction with excess of the Cu(II)-binding ligand Bca in Mops buffer containing reductant NH₂OH. Under such conditions, all copper ions are converted quantitatively to the well-defined chromophoric complex anion [Cu(Bca)]⁻ with characteristic absorbance at 562 nm (ε = 7900 M⁻¹ cm⁻¹). The peptides and probes listed in Table 1 were purchased from GL Biochem (Shanghai) except for Aβ16-A2P and Aβ16-D1H/H14A which were synthesized on site by standard solid phase peptide techniques as described elsewhere.
Table 1: Sequences of Aβ16 peptides and probes

<table>
<thead>
<tr>
<th>Peptide or probe</th>
<th>Sequence</th>
<th>Molar mass [Da]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aβ16</td>
<td>DAEFRHDGYEVH0QK</td>
<td>1953.0 1954.5</td>
</tr>
<tr>
<td>Ac-Aβ16</td>
<td>Ac-DAEFRHDGYEVH0QK</td>
<td>1997.0 1997.5</td>
</tr>
<tr>
<td>D1A</td>
<td>DAEFRHDGYEVH0QK</td>
<td>1911.0 1910.7</td>
</tr>
<tr>
<td>D1E</td>
<td>EAEFRHDGYEVH0QK</td>
<td>1969.0 1970.4</td>
</tr>
<tr>
<td>D1N</td>
<td>NAEFRHDGYEVH0QK</td>
<td>1954.0 1954.9</td>
</tr>
<tr>
<td>D1H</td>
<td>HAEFRHDGYEVH0QK</td>
<td>1977.0 1978.0</td>
</tr>
<tr>
<td>D1H/H14A</td>
<td>HAEFRHDGYEVH0QK</td>
<td>1911.0 1911.4</td>
</tr>
<tr>
<td>A2P</td>
<td>DPEFRHDGYEVH0QK</td>
<td>1981.0 1981.5</td>
</tr>
<tr>
<td>DP2</td>
<td>D[K138]HDDH</td>
<td>1240.3 1240.9</td>
</tr>
<tr>
<td>DP3</td>
<td>HK138HDDH</td>
<td>1118.2 1118.0</td>
</tr>
<tr>
<td>DP4</td>
<td>H[K138]H</td>
<td>790.8 790.4</td>
</tr>
</tbody>
</table>

* The residues changed are underlined in bold.

Detailed previously.25 The identity of each peptide was verified by electrospray ionization mass spectrometry (ESI-MS; Table 1) while the purity was confirmed to be >98% by HPLC. Peptide concentrations were estimated from absorbance maxima at ~276 nm using ε_{max} = 1410 M^{-1} cm^{-1} for Aβ16 peptides containing a single tyrosine residue. The concentration of dansyl peptide probes DP2–4 was estimated by absorbance maxima at ~326 nm using the reported ε_{max} = 4500 M^{-1} cm^{-1} for each probe.21 The concentrations obtained matched those estimated from fluorescence titrations with the copper standard assuming the formation of a 1:1 complex.

All ligand stock solutions, reaction buffers, reagents, and solutions were prepared in rigorously deoxygenated Milli-Q water and stored in an anaerobic glove-box (<1 ppm O2). The working solution of ligand Fs was prepared freshly from aliquots of frozen stock solution stored in a freezer at temperature −80 °C.26 All reactions involving Cu(i) were performed anaerobically in the glove-box and transported outside in a capped cuvette for spectroscopic characterization.

Spectroscopic methods

UV-visible spectra were recorded on a Varian Cary 300 spectrophotometer in the dual beam mode with quartz cuvettes. All metal titrations were performed in appropriate buffers and corrected for baseline and dilution. Fluorescence emission spectra were recorded on a Varian Cary Eclipse spectrophotometer with a band pass of 20 nm for both excitation and emission. The excitation wavelength for the dansyl peptide (DP) probes was 330 nm and the emission spectra were recorded between 450 and 750 nm at a scan rate of 600 nm/s. Electron paramagnetic resonance (EPR) spectra were recorded on a Bruker Elexys E 500 EPR spectrometer under the following instrument conditions: microwave frequency ν = 9.456–9.495 GHz but conversion to a single frequency of 9.475 GHz for reporting via relationship g = 714.4775 ν/F, where F is the magnetic field in the Gauss unit, the microwave power is ~1 mW, the modulation amplitude is 4 G, the modulation frequency is 100 kHz, the sweep time is 40 s, the time constant is 20 ms, and the average number of scans is 10. All sample solutions (total 0.2 mL) were prepared in 50 mM buffer (Mes at pH 5.9–6.5; Mops at pH 6.6–8.0 or Ches at pH > 8) containing [Cu(i)]_{tot} = 0.45 mM, [peptide or probe]_{tot} = 0.5 mM and 5% glycerol. The solution pH of each sample was re-checked after recording each spectrum. The samples were snap-frozen in liquid nitrogen and the spectra recorded at 77 K in a liquid-nitrogen finger Dewar. The EPR parameters (g values and A values) were extracted directly from the recorded spectra without simulation.

Quantification of Cu(ii) binding

Cu(ii) binding to Aβ peptides was quantified with a series of DP probes as reported recently.21 These probes feature a dansyl group and emit intense fluorescence at λ_{max} ~ 550 nm upon excitation at ~330 nm. They bind one equiv. of Cu(ii) with varying affinities and Cu(ii) binding to the probes is detected sensitively via proportional fluorescence quenching. The quantification is based on the following relationships:

\[ \text{Cu}^2\text{DP} + \text{Aβ} \rightleftharpoons \text{Cu}^2\text{iAβ} + \text{DP} \] (1)

\[ K_{ex} = \frac{[	ext{Cu}^2\text{iAβ}][\text{DP}]}{[	ext{Cu}^2\text{DP}][\text{Aβ}]} = K_{i} \left( \frac{[	ext{Cu}^2\text{DP}]}{[	ext{Cu}^2\text{iAβ}]} \right) \] (2)

\[ \frac{[	ext{Aβ}]_{tot} - [	ext{Cu}^2\text{DP}]}{[	ext{DP}]} = \frac{K_{D} (\text{Cu}^2\text{iAβ})}{K_{D} (\text{Cu}^2\text{DP})} \left( \frac{[	ext{Cu}^2\text{iAβ}]}{[	ext{Cu}^2\text{DP}]} - 1 \right) \left( 1 - \frac{[	ext{Cu}^2\text{DP}]}{[	ext{DP}]} \right) \] (3)

\[ \frac{[	ext{Cu}^2\text{DP}]}{[	ext{DP}]} = \frac{F_0 - F}{F_0 - F_1} = \frac{\Delta F}{\Delta F_1} \] (4)

where K_{D} (Cu^2\text{iAβ}) and K_{D} (Cu^2\text{DP}) are the respective dissociation constants under the same conditions. Eqn (3) is derived from eqn (2) with incorporation of mass balance relationships for an effective competition described by eqn (1) and is a convenient format for curve-fitting of the experimental data to derive K_{D} (Cu^2\text{iAβ}) with known K_{D} (Cu^2\text{DP}). Under the conditions of the fluorescent emission of the dansyl probes (λ_{ex} = 330 nm; λ_{em} = 550 nm), the Aβ peptides are fluorescence-silent. Consequently, the Cu(ii) speciation in eqn (1)–(3) may be analysed conveniently and reliably by monitoring the response of the probe to binding of Cu(ii) via eqn (4), where F_0, F, and F_1 are the fluorescence intensity of the DP probe upon binding 0, <1.0 and 1.0 equiv. of Cu(ii), respectively. The experiments were performed via reported protocols.21

The K_{D} values for each peptide were determined similarly at a range of pH values. The pH-dependent affinities of the Cu(ii) probes DP2–4 have been reported recently.21 If the Cu(ii) binding site remains unchanged within a pH range and the ligand compositions are known, then the conditional K_{D} within that pH range may be correlated with the absolute dissociation constant (K_{D}^{abs}) and the proton ionization constants (K_{i}) of each ligand function of the site according to eqn (5):

\[ K_{D} = K_{D}^{abs} \left( 1 + \sum_{i=1}^{n} 10^{\left( \sum_{i=1}^{n} pK_{a,i} - \text{pH} \right)} \right) \] (5)

The experiments were conducted in buffers (50 mM) Mes (pH 6.0–6.6), Mops (pH 7.0–7.4), Heps (pH 7.8–8.2) and Ches...
Absorbance A265 of the reduced Asc. The oxidation rate was determined oxidation rate without added Cu has been removed from the background absorbance A265 = 14 500 M−1 cm−1 for Asc.28 The catalytic activity of each Cu–Aβ complex was expressed as molar turn-over number per min (MTN/min) and was obtained from the slope of a linear plot of the Asc oxidation rate (μM min−1) versus the concentration (μM) of each Cu–Aβ complex. The background oxidation rate without added Cu has been removed from the determined oxidation rate.

Formal reduction potential

The formal reduction potential of the copper centres in each Cu–Aβ16 complex was estimated from the relative affinities of the peptides for Cu(i) and Cu(u) under the same conditions via the Nernst relationship of eqn (8):

$$E^0 = E^0 + 59 \log \left( \frac{K_D(Cu^{II})}{K_D(Cu^{I})} \right)$$

where E0 = 153 mV (vs. SHE) is the standard reduction potential of Cu(II)/Cu(I).29

Results and discussion

The N-terminal residues of Aβ peptides have a major impact on Cu(u) binding

Processing of the amyloid precursor protein (APP) by proteases β- and γ-secretase generates the Aβ peptides 1–(36–43) whose interaction with transition metal ions, and with copper, in particular, appears to be a source of toxicity in vivo.30 The possible effects include catalytic generation of reactive oxygen species (ROS) by toxic oligomers.31,32 All major Cu ligands are located within the first 16-residue water-soluble fragment and thus Aβ16 is a valid model for the study of the copper chemistry of full length Aβ.3 In particular, the N-terminal amine nitrogen and nearby residues play a critical role in Cu(u) coordination and in imposing both binding affinity and coordination modes.

In this study, the two N-terminal residues Asp1 and Ala2 of Aβ16 were changed systematically via generation of the following variants (Table 1):

(i) D1A eliminates the sidechain of residue 1 as a potential ligand (cf. IA of Fig. 1b).

(ii) D1E, D1N, and D1H provide alternative residue 1 side-chains as potential ligands; D1H/H14A also removes His14 as a potential ligand.

(iii) A2P removes the amide of residue 2 as a potential anionic ligand (cf. II of Fig. 1b).

In addition, the N-terminus was acetylated (Ac-Aβ16) to perturb its metal binding properties. Detection of fine differences in properties is possible with a set of peptide-based dansyl fluorescent probes DP2–4 (Tables 1 and 2).21

A control titration of standardized CuSO4 solution with the DP2 probe alone generated a titration curve with an endpoint at [Cu(u)]tot/[DP2] = 1.0 (Fig. 3a and b, black trace), confirming that the probe binds a single Cu(u) ion of the highest affinity (log KD = −9.9).21 The equivalent titration into a 1:1 mixture of DP2 and Aβ16 (2.0 μM) in Mops buffer (50 mM, pH 7.4) generated a titration curve with an apparent endpoint at [Cu(u)]tot/[DP2] ~ 2.0 (Fig. 3b, red trace). The essentially linear nature of the curve indicated that the peptide can bind one equiv. of Cu(u) with an affinity similar to that of DP2. Indeed, the Cu(u) binding sequence in the DP2 peptide was designed as a mimic of Aβ16 (Table 1).21

Equivalent titrations for Aβ16-D1A and Aβ16-D1H also displayed apparent titration endpoints at [Cu(u)]tot/[DP2] ~ 2.0 (Fig. 3b, green, blue), consistent with these peptides also binding a single Cu(u) ion. However, the different shapes of the titration curves are consistent with the affinity of Aβ16 being higher than that of Aβ16-D1A but lower than that of Aβ16-D1H. In fact, the probe...
A more quantitative approach is possible with the fluorescence recovery analysis of titrations of apo-peptides into solutions of a copper-probe complex of appropriate affinity. Preliminary testing indicated that the appropriate experiments involved substoichiometric Cu(u) in the systems Cu_{13}–DPx (x = 2 for Aβ16-D1A and 3 for Aβ16-D1H and x = 2 or 3 for Aβ16). Analysis of the results suggested that, at pH 7.4, the Cu(u) binding affinity of Aβ16-D1A is 5-fold weaker than that of Aβ16 while the affinity of Aβ16-D1H is ~100 times stronger (Fig. 3c–f and Table 2).

Equivalent experiments and analyses for the remaining peptides, with their respective optimal DP probes, lead to an increasing order of relative Cu(u) binding affinities at pH 7.4 as follows (Table 2):

\[ \text{Ac-Aβ16 (5.0)} < \text{Aβ16-D1A (5.0)} < \text{Aβ16-D1E (3.2)} \]

\[ < \text{Aβ16-D1N (2.0)} < \text{Aβ16 (1.0)} \sim \text{Aβ16-A2P (1.0)} \]

\[ < \text{Aβ16-D1H/H14A (0.02)} < \text{Aβ16-D1H (0.01)} \quad (9) \]

The numbers in the brackets refer to the dissociation constants normalized to that of Aβ16 and decrease in the order of increasing Cu(u) binding affinity. It is apparent that the residues in the N-terminal region have a major impact on the binding affinity and likely on the binding mode as well. A more complete analysis follows.

### Component IA is favoured by the presence of a Hist1-based chelate ring

The two most significant changes in Cu(u) binding affinity are seen for Ac-Aβ16 and Aβ16-D1H (eqn (9)). The N-terminal amine nitrogen was blocked by acetylation in Ac-Aβ16, leading to a significant decrease in affinity under all conditions (Table 2), confirming the primary anchoring role of the N-terminal nitrogen in Cu(u) binding for each of the components IA, IB and II. Replacement of the carbonyl sidechain of Asp1 with the imidazole sidechain of His1 in Aβ16-D1H led to an increase in affinity of almost two orders of magnitude at pH 7.4 (Table 2).
The presence of His1 allows the formation of a stabilizing bidentate ligand comprising the N-terminal amine and the His1 sidechain within a six-membered chelate ring. Other known examples of this binding mode include the DP3 probe employed in this work (Fig. 2a) and the bacterial copper binding proteins PcoC and CopC. The Cu(II) coordination sphere in CuII(A16-D1H) may be depicted schematically by Fig. 4a(i), noting its similarity to the proposed IA site in A16 (Fig. 1b). There are three other His residues (His-6, -13, -14) in the A16-D1H sequence, but only two of them are required to constitute the proposed Cu(II) site. In fact, replacing one of them (His14) with Ala in A16-D1H/D1A only marginally affects the Cu(II) affinity (Fig. 5a and Table 2).

The Cu(II) sites proposed above for CuII(A16-D1H) and CuII(A16-D1H/H14A) are supported by the fact that the pH-dependence of their affinities for Cu(II) (expressed as pK\textsubscript{a} = \(-\log K_{a}^{\text{Cu}}\)) can be modeled satisfactorily over the pH range 6.0–8.7, assuming that the dependence is driven by the basicity of the proposed ligands (Fig. 5a and eqn (5)). Their derived ligand pK\textsubscript{a} values are listed in Table 3. The I \(\rightarrow\) II transition is not evident in these peptides and the pH dependence of the affinities is comparable to those of several documented examples including DP3 and several CopC proteins that adopt a DP3-type Cu(n) binding mode, shown in Fig. 2a,21,13

In contrast, the absence of His1 in the WT, D1A and A2P peptides leads to very different pH-dependencies that could not be modelled over the same pH range with any single set of ligands or a single binding mode (Fig. 5). The behaviour is attributed to the influence of the I \(\rightarrow\) II transition, even for the A2P peptide at pH > 8.5 (see Fig. 6c). The Cu(II) site of component II features a deprotonated peptide amide ligand (Fig. 1b).

pH-Dependent EPR spectra provide further support for the proposed binding models. The EPR signal of CuII(A16-D1H) shows no change within the pH range 5.9–8.9 and essentially superimposes that of CuII–DP3 (Fig. 6a and b). It differs significantly from the component I EPR signals of CuII(A16) or CuII(A16-A2P) (cf., Fig. 6a–d and Table 4). This supports a Cu(II) site in CuII(A16-D1H) of composition [NH\textsubscript{2}, 3Nim] that features the stabilizing bidentate ligand comprising the N-terminal amine and the His1 sidechain (Fig. 2a and 4a(i)).

This particular His1 sidechain binding mode is suppressed in CuII–DP4. Although its four-residue peptide sequence (His-Lys-His-His) features a His1 residue, the presence of His3 provides the more favourable ATCUN binding mode that includes the N-terminal amine and two intervening deprotonated backbone amides (Fig. 2b).21 This conclusion is supported by the dramatic pH-dependence of its Cu(II) binding affinity and the unique EPR signal: both are readily distinguishable from those of components I and II (Fig. 5b and 6f).21 This same ATCUN binding mode is adopted by the N-truncated A16 peptide, A4-16 (sequence: FRHDSGYEVHHQK) that exhibits Cu(II) affinities and EPR spectra equivalent to those of DP4 (Table 4).6,21–23

Component II is favoured by an Ala1 N-terminals at low pH but is less stable than IA

The variant A16-D1A has a five-fold lower affinity for Cu(II) than A16 at pH 7.4 (eqn (9); Fig. 5a; Table 2). The I \(\rightarrow\) II transition in CuII(A16) with pK\textsubscript{a} ~ 7.8 is characteristic of deprotonation of the peptide amide of Ala2. The non-coordinating methyl side-chain of Ala1 eliminates the IA binding mode, enforcing a IB mode (Fig. 4a(ii)) that is converted to component II at pH ~ 6.5, i.e., > 1 pH unit lower than that for the equivalent transition of CuII(A16) (Fig. 6d and e). However, at higher pH values, the affinities of A16 and A16-D1A for Cu(II) become indistinguishable (Fig. 5a), consistent with a common tridentate chelate [–NH\textsubscript{3}, N\textsuperscript{2+}, CO] as part of the square planar array [–NH\textsubscript{3}, N\textsuperscript{2+}, CO, N\textsubscript{im}], characteristic of component II (Fig. 1b). The binding affinity of the latter shows a different pH dependency and EPR signal to those of DP4, a Cu(II) ligand of the ATCUN binding mode (Fig. 2b, 5 and 6).

The equivalent data for variants D1N and D1E provide further support for a role for Asp1 as an equatorial ligand: their Cu(II) binding properties are intermediate between those of A16 and
Table 3  Thermodynamic data of Cu(ii) sites derived from correlation of log $K_D$ and pH$^a$

<table>
<thead>
<tr>
<th>Peptide or protein</th>
<th>Cu(ii) binding site</th>
<th>Log $K_D^a$</th>
<th>pK$\alpha$ for</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aβ16-D1H</td>
<td>NH$<em>2$ (H1); 3 N$</em>{14}^{14}$ (H1,6,13/14)</td>
<td>$-$12.7</td>
<td>8.3</td>
<td>6.4</td>
</tr>
<tr>
<td>Aβ16-D1H/H14A</td>
<td>NH$<em>2$ (H1); 3 N$</em>{14}^{14}$ (H1,6,13)</td>
<td>$-$12.3</td>
<td>8.1</td>
<td>6.4</td>
</tr>
<tr>
<td>DP3 probe</td>
<td>NH$<em>2$ (H1), 3 N$</em>{14}^{14}$ (H1,5,7)</td>
<td>$-$13.6</td>
<td>8.8</td>
<td>6.6</td>
</tr>
<tr>
<td>Ps-CopC</td>
<td>NH$<em>2$ (H1); 2 N$</em>{14}^{14}$ (H1,91), COO$^-$ (D89)</td>
<td>$-$14.4</td>
<td>8.2</td>
<td>6.1</td>
</tr>
<tr>
<td>Ec-PcoC</td>
<td>NH$<em>2$ (H1); 2 N$</em>{14}^{14}$ (H1,92), COO$^-$ (D90)</td>
<td>$-$14.0</td>
<td>8.2</td>
<td>6.5</td>
</tr>
</tbody>
</table>

$^a$ See Fig. 5 for experiments; parameters obtained by curve-fitting to eqn (5).

Table 4  EPR parameters derived from frozen solution spectra$^a$

<table>
<thead>
<tr>
<th>Ligand for the Cu$^{II}$ complex</th>
<th>Component I</th>
<th>Component II</th>
<th>pK$\alpha$ for I ↔ II</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$g_i$</td>
<td>$g_\perp$</td>
<td>$A_i (G)$</td>
</tr>
<tr>
<td>DP3</td>
<td>2.242$^b$</td>
<td>2.060$^b$</td>
<td>182$^b$</td>
</tr>
<tr>
<td>Aβ16-D1H$^c$</td>
<td>2.243</td>
<td>2.051</td>
<td>183</td>
</tr>
<tr>
<td>Aβ16-A2P$^c$</td>
<td>2.269</td>
<td>2.051</td>
<td>169</td>
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<td>Aβ16$^c$</td>
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<td>Aβ16-D1A$^c$</td>
<td>2.269</td>
<td>2.051</td>
<td>169</td>
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<tr>
<td>Aβ16-2.264$^d,e$</td>
<td>170$^e$</td>
<td>2.226$^e$</td>
<td>153$^f$</td>
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<tr>
<td>DP4</td>
<td>—</td>
<td>—</td>
<td>2.178$^g$</td>
</tr>
<tr>
<td>Aβ16-16$^c$</td>
<td>—</td>
<td>—</td>
<td>2.178$^g$</td>
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</tbody>
</table>

$^a$ Estimated directly from the respective spectrum, see Fig. 6, Fig. S1–S7 (ESI) for details. $^b$ Data adopted from ref. 21. $^c$ Data for Cu$^{II}$(Aβ16-D1H/H14A) were indistinguishable (see Fig. S5 and S6, ESI). $^d$ Data adopted from ref. 16. $^e$ Minor component at pH 7.4.

Fig. 6  Frozen solution EPR spectra of various Cu(iii) complexes recorded at 77 K. The Cu(iii) ligands are: (a) DP3 probe; (b) Aβ16-D1H; (c) Aβ16-A2P; (d) Aβ16; (e) Aβ16-D1A; (f) DP4 probe at various pH values as indicated (see the Experimental section for detail). Insets show enlargements of the spectra scaled up by a factor of 3. Component I is indicated by green dashed lines and component II by red dashed lines.

D1A (Table 2 and Fig. S1–S4, ESI$^f$). The data are consistent with the weaker ligand properties of the sidechains of Asn1 (neutral carboxamide is a poor ligand for Cu(ii)) and Glu1 (which would form a less favoured seven-membered chelate ring). In particular, substitution of Asp1 with Asn or Glu also leads to a decrease in Cu(ii) binding affinity (Table 2) and in the pK$\alpha$ for the I ↔ II transition (Fig. S1 and S4, ESI$^f$).

Several previous studies provide support for this model. Deletion of Asp1 in Aβ16 generates a truncated peptide with an Ala N-terminus (sequence: AEFRHSGYEVHHQK). Its qualitative Cu(ii) binding properties are comparable to those of D1A: weaker binding affinity for Cu(ii) and a lower I ↔ II transition pK$\alpha$. Similar effects were also observed for D1N,17–19 although the decreased pK$\alpha$ was attributed to the change of a putative hydrogen bond to the Cu(ii) ligand.17

Component I is favoured in Cu$^{III}$(A2P)

The I ↔ II transition relies on the availability of the Asp1-Ala2 peptide amide (Fig. 1b). Substitution of Ala2 with Pro eliminates this possibility and all properties observed for Cu$^{III}$(A2P) are consistent with the binding mode being locked in component I. Most obviously, $K_D$ is comparable to that of Cu$^{II}$(Aβ16) at lower pH values but becomes larger (i.e., lower affinity) at higher pH with the point of deviation occurring above pH 7.8, in line with the I ↔ II transition for Cu$^{III}$(Aβ16) around this pH (Fig. 5a and 6d). The EPR signal confirmed that component I of Cu$^{III}$(Aβ16-A2P) was present up to at least pH 8.5 (Fig. 6c).

At lower pH, component IA is preferred for $\beta$ peptides with an Asp1 N-terminus

The Cu(ii) affinities of component I of Cu$^{III}$(Aβ16) and Cu$^{III}$(Aβ16-A2P) are similar. They are about one order of magnitude higher than that of component IB, which is favoured for Cu$^{III}$(Aβ16-D1A) at lower pH values such as pH 6.2 (Fig. 5a and Table S1, ESI$^f$). This implies that component IA rather than IB is the preferred binding mode for both Cu$^{III}$(Aβ16) and Cu$^{III}$(Aβ16-A2P) at lower pH values (Fig. 1b). This is consistent with the...
negatively charged harder carboxylato ligand favouring the harder metal ion Cu(i) over a neutral peptide carbonyl oxygen for similar stereochemistries (a six-membered chelate ring in IA versus a five-membered chelate ring in IB; see Fig. 1b).

However, the possibility for an IB binding mode for either CuH(16) or CuH(16-A2P) at low pH cannot be excluded. The EPR signals for both components IA and IB are indistinguishable (Fig. 6c–e and Table 4) and cannot be used to differentiate the binding modes at low pH. It is possible that components IA and IB are in fast exchange at lower pH for those peptides with an Asp1 N-terminus but with IA being the preferred binding mode. At higher pH, both components IA and IB are converted to component II: the Kd values of CuH(16) and CuH(16-D1A) become indistinguishable at pH 9.4 (Fig. 5a). But the I → II conversion is restricted for CuH(16-A2P), weakening its Cu(i) binding affinity at high pH relative to other Aβ16 peptides (Fig. 5a).

It is unlikely that the differences in the properties of CuH(16), CuH(16-A2P) and CuH(16-D1A) at lower pH values are due to axial coordination by the Asp1 sidechain due to the following reasons:

(i) Given the possibility of axial coordination, mutation of Asp1 by Ala should lower the affinity across the whole pH range. However, there is no difference in the Cu(i) affinity for component II between WT and D1A.

(ii) A bidentate [NH3, COO−] binding mode for peptides with N-terminal aspartyl residues has been well-documented in many cases, including various Aβ peptides.11,36,37

(iii) The IA binding mode is confirmed for many peptides and proteins with a His1 N-terminus including Aβ16-D1H, DP3 probe, CopC and PcoC proteins, as surveyed above.

(iv) Contributions from axial coordination at Cu(i) centres are usually negligible due to the Jahn–Teller effect. There are many possible carboxylate sidechains (Asp1, Asp7, Glu3 or Glu11) which have been proven to be redundant. For example, an E3Q mutation showed no significant effect on the EPR spectrum.3

The IA coordination mode for Aβ16 has been proposed frequently as one of the two possible coordination modes of Asp1 at lower pH for component I (Fig. 1b).5–11 However, component IB has been preferred recently, based mainly on pulsed EPR studies.3,12,13 The chemical and spectroscopic study presented in this work, supported by previous Cu(i) binding studies for those peptides with N-terminal aspartyl residues,11,36 provides strong evidence that the preferred binding mode for component I in CuH(16) is IA, and not IB (Fig. 1b). It is recommended that the possibility of the IB binding mode can be re-investigated by pulsed EPR using CuH(16-D1H) and CuH(16-D1A) as controls.

Impact of N-terminal modification on Cu(i) binding

The first evidence of Cu(i) binding to Aβ peptides was implicated in an early study of Cu(i) potentiation of Aβ neurotoxicity.38 We now know that Aβ peptides bind Cu(i) mainly via a combination of two histidine residues within the first 16 residues.7,8 The Cu(i) binding stoichiometries and affinities of various Aβ16 peptides generated in this work were examined using Cu(i) probe ligands Ferene S (Fs) and Ferrozine (Fx) according to a recent protocol.26

Both probe ligands react with Cu(i) under reducing conditions to yield respective chromophoric Cu(i) complexes [CuL2]3− with different formation constants (log β2 = 13.7 for L = Fs and 15.1 for L = Fx).26 They were used to estimate the binding stoichiometries of Aβ16 peptides according to eqn (6) under weakly competing conditions and the binding affinities according to eqn (6) and (7) under strongly competing conditions. As demonstrated previously for Aβ16,8 most peptides listed in Table 1 bind a single equiv. of Cu(i) under Cu2+ limiting conditions imposed by a slight excess of the weak probe ligand Fs (Fig. 7a). The exception was Aβ16-D1H that appeared to bind more than one equiv. Cu(i) under the same conditions (Fig. 7b).

Increasing Fs ligand concentration or employing the stronger ligand Fx created more effective competing conditions for eqn (6) that allowed analysis and definition of Cu(i) binding affinity of each peptide with eqn (7) (Fig. 7c and d). There are only marginal differences in Cu(i) binding affinities for those peptide variants with substitution of a non-His residue in the sequence. But, the Cu(i) binding affinities of Aβ16-D1H and Aβ16-D1H/H14A are slightly higher than those of the other peptides (Table 5).

The data are consistent with previous observations: (i) the non-His residues in the N-terminal region play little role in Cu(i) binding;38 (ii) two and only two His ligands are required for the flexible Aβ16 peptides to constitute a Cu(i) binding site (Fig. 1a);7,8,39–41 (iii) the Cu(i) bound by Aβ16 peptides is
Ac-A/C0D1H/H14A residues in determining these Cu(II) binding modes and an
study has identified the critical role of the first two N-terminal
Cu(I) is expected to restrict the binding sites to two His pairs only
may be populated among six different combinations of His pairs
Binding of the first equiv. of Cu(I) appears to discourage the
similar affinities, but this does not seem to be the case (Fig. 7b).
ligands,49 but also to the surrounding residues. 50 Indeed, Asc is
radical that can cause damage not only to the direct copper
further reduction
of copper to A
metal dyshomeostasis in the Alzheimer brain may result in binding
Impact of the N-terminal modifications on the catalytic aerobic oxidation of ascorbate
A common feature in Alzheimer’s disease is oxidative stress, a
component of which may result from undesirable catalytic reduction of dioxygen by cellular reductants such as ascorbate (Asc) mediated by Cu complexes of Aβ peptides.5,6,31,32,38,46–48 The dioxygen reduction product H2O2 is liable to undergo further reduction via Fenton-type reactions to generate the hydroxyl radical that can cause damage not only to the direct copper ligands,49 but also to the surrounding residues.50 Indeed, Asc is abundant in the brain as an important neuromodulator and/or a neuroprotective agent and its concentration fluctuates dynamically in the range 200–400 μM in certain synaptic clefts.51,52 In reality, the oxidation rate of Asc by dioxygen is slow and must be catalysed by redox-active couples such as Cu(n)/Cu(i). It has been proposed that metal dyshomeostasis in the Alzheimer brain may result in binding of copper to Aβ peptides to initiate catalytic roles.53
Aβ peptides can bind Cu(i) with the distinct binding modes of components I (including IA and IB) and II at physiological pH (Fig. 1b). Recent studies suggest that the different components available in Cu–Aβ16 systems (Fig. 1) display different reaction kinetics in both metal substitution and redox chemistry.5,6 This study has identified the critical role of the first two N-terminal residues in determining these Cu(n) binding modes and an integrated study of the thermodynamic and kinetic contributions of these binding modes in the generation of H2O2 via catalytic aerobic oxidation of Asc was also undertaken. It should be noted that the reaction scheme in Fig. 8a is presented as an overall two-electron reaction process, but it may proceed via two consecutive one-electron steps with O2•− as an immediate reduction intermediate that may be reduced further to H2O2 by a second Cu1–Aβ complex.54
The experiments were performed in air-saturated buffers (50 mM) at three different pH values (Mops buffer at pH 6.6 and 7.4; Ches buffer at pH 8.7). The catalytic aerobic oxidation of Asc (50 μM) was undertaken with each Cu–Aβ16 complex (prepared with a one-fold excess of apo-peptide) at a range of concentrations (1.0–20 μM). The reaction was followed by the rate of decrease in absorbance at 265 nm for the reduced form of Asc (Fig. 8a). The catalytic activity of each Cu–Aβ16 peptide complex was determined from the slope of a linear correlation

| Aβ16 peptide | log K_D pH 6.2 | log K_D pH 7.4 | log K_D pH 9.4 | Δα/Δβ K_D/Δα K_D (wt) | pH 6.2 | pH 7.4 | pH 9.4 | Probe-affinity std
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α The log K_D values were determined via ligand competition for Cu(i) between each peptide and the specified probe ligands Fs and/or Fz in the respective buffers (50 mM) Mes (pH 6.2), Mops (pH 7.4) and Ches (pH 9.4). See the Experimental section for details.
of the initial oxidation rates versus the catalyst concentrations and expressed as μM Asc oxidized/μM catalyst per minute, i.e., molar turnover number min⁻¹ (see the Experimental section for details).

As observed previously, Asc is stable in metal-free air-saturated buffers within the experiment timescale (1–2 h) and all apo-peptides are inactive (at least after demetallation with EDTA). ‘Free’ Cu²⁺ and Cu⁺ have robust catalytic activities that are suppressed, but not silenced, by apo-peptides. The Cu–Aβ16 complexes are redox-active catalysts, but less active than ‘free’ copper. A one-fold excess of apo-peptide over each Cu–peptide was sufficient to minimize the catalytic contribution from ‘free’ copper and to allow reliable detection of the catalytic activity of the Cu–complex. The results are summarized in Fig. 8, Fig. S8 (ESI†) and Table 6.

At pH 7.4, the molar catalytic activities (min⁻¹) of the different Cu–Aβ16 complexes for Asc oxidation are given below in increasing order of their activity:

D1A (0.08 min⁻¹) ~ D1E (0.10) ~ D1N (0.14) < wt (0.38) < A2P (0.66) < D1H (0.78) ~ D1H/H14A (0.9) (10)

As discussed above, the binding modes and affinities of these peptides are similar for Cu(i) (with minor difference for Aβ16-D1H and Aβ16-D1H/H14A) but differ considerably for Cu(ii). On the one hand, replacement of Asp1 with Ala, Glu or Asn leads to destabilization of component IA inducing the I → II transition at lower pH (Table 4 and Fig. 6). On the other hand, substitution of Ala2 with Pro suppresses the transition while substitution of Asp1 by His1 further stabilizes the IA mode (Fig. 4a(i)). The observed reactivity order of eqn (10) clearly indicates that Cu sites accommodating a component I type Cu centre possess higher redox catalytic activity than those adopting a component II binding mode only. The conclusion applies to both pH 6.6 and 8.7 (Table 6; Fig. 8 and Fig. S8, ESI†).

However, contradictions to these conclusions can be seen in Table 6. For example, while components IA and II dominate at pH 6.6 and 8.7, respectively, for Cu⁶⁺(Aβ16), they are both present in equilibrium at pH 7.4 (Fig. 6d and Fig. S1, ESI†), but the observed activities at pH 7.4 and 8.7 were essentially identical (Table 6). The complications arise from the fact that Asc is more stable towards oxidation at lower pH and less stable at higher pH values. Hence, there are two opposing effects acting on the Asc oxidation rates: (i) a transition from component I at lower pH to component II at higher pH decreases the catalytic rate; (ii) deprotonation of Asc that occurs at higher pH increases its oxidation rate. This latter effect is seen clearly for Cu(Aβ16-A2P), Cu(Aβ16-D1H) and Cu(Aβ16-D1H/H14A). Within the pH range 6–9, component IA dominates in the first complex and is locked in place in the latter two complexes (Fig. 6b and c). However, the catalytic oxidation rate increased with pH for each complex.

To separate these two opposing effects, the observed oxidation rates were normalized to that of Cu(Aβ16-D1H) under the same conditions (Table 6). The normalized data indicate: (i) the redox activities of Aβ16 and variants Aβ16-D1A, -D1E and -D1N decrease uniformly with increasing pH, consistent with the influence of the I → II transition (Fig. 6d, e and Fig. S1–S4, ESI†); (ii) the redox activity of Cu(Aβ16-A2P) is lowest at pH 8.7, consistent with a partial transition of the Cu(n) centre in this complex from component I to other unknown Cu(n) centre(s) (Fig. 6c) at high pH; (iii) the redox activities of the two Cu complexes with a His1 N-terminus are higher than all others and notably, the one with peptide Aβ16-D1H/H14A has the highest activity consistently at all pH values (Table 6). This high redox activity is attributed to a combined redox advantage of the Cu centres in both oxidation states and this is further analysed and discussed below.

Determinants that regulate the redox activity of Cu–Aβ peptides

In order to be an efficient redox catalytic centre for a particular process of oxidation and reduction, the formal reduction

<table>
<thead>
<tr>
<th>Table 6</th>
<th>Aerobic oxidation rate of ascorbate catalyzed by various Cu–Aβ16 complexes and calculated thermodynamic reduction potentials⁺</th>
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<tr>
<td></td>
<td>Catalytic rate as molar velocity (min⁻¹) at pH</td>
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<tr>
<td>Aβ16 peptide</td>
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<td>Aβ16</td>
<td>0.28</td>
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D1A | 0.09 | 0.08 | 0.17 | 0.13 | 0.10 | 0.06 | +277 | +242 | +40.9 |
D1E | 0.09 | 0.10 | 0.12 | 0.14 | 0.13 | 0.04 | +283 | +218 | +35.0 |
D1N | 0.15 | 0.14 | 0.15 | 0.22 | 0.17 | 0.06 | +247 | +206 | +40.9 |
A2P | 0.32 | 0.66 | 1.12 | 0.46 | 0.84 | 0.42 | +247 | +206 | +70.4 |
D1H | 0.69 | 0.78 | 2.70 | 1.00 | 1.00 | 1.00 | +183 | +118 | +46.8 |
D1H/H14A | 0.79 | 0.90 | 3.29 | 1.16 | 1.15 | 1.22 | +194 | +118 | +64.5 |
Ac-Aβ16 | +277 | +283 | +35.0 |

⁺ Conditions: [Asc]₀ = 50 μM; [Cu–peptide]₀ = 1–20 μM with a one-fold excess apo-peptide; buffers (50 mM) of Mops (pH 6.6 and 7.4) and Ches (pH 8.7). The background oxidation rate without the added Cu has been removed, see the Experimental section for details. Calculated from eqn (8) using E⁰ = +133 mV for the redox couple Cu²⁺/Cu⁺ (note: the Cu(i) Kθ values at pH 6.6 and 8.7 were not determined and thus the calculations used data at pH 6.2 and 9.4 instead). Determined by cyclic voltammetry, except a recent theoretical calculation predicted +280 mV for component I and ~370 to ~810 mV for component II.⁵⁻
potential of the copper centre must be lower than that of the oxidant but higher than that of the reductant. At pH 7.4, the formal reduction potentials of ascorbate and dioxygen are $\sim +50 \text{ mV}^{28}$ and $\sim +300 \text{ mV}^{27}$ (vs. SHE), respectively.

The formal reduction potentials of the Cu-Aβ complexes may be estimated via the Nernst relationship of eqn (8) from the respective Cu(i) and Cu(u) affinities of each Aβ16 peptide (Table 6). The reduction potentials are generally more negative for component II than for component I and decrease with increased pH. This is consistent with the more dramatic stabilization effect for Cu(u) coordination at higher pH, especially with the dominant coordination effect of the deprotonated amide nitrogen to the Cu(u) centre in component II.\(^\text{5,43}\) The calculated reduction potentials in this work from the respective dissociation constants generally match those determined experimentally by cyclic voltammetry or predicted from theoretical calculations (Table 6). However, it must be noted that the dissociation constants of the Cu–Aβ complexes are quite large (indicating relatively weak binding). In addition, the estimated values vary considerably with experimental conditions, as do the reduction potentials determined by cyclic voltammetry (Table 6). The redox processes are frequently not well-defined due to the different coordination environments for Cu(i) and Cu(u). Consequently, it is difficult to define the reduction potentials consistently and accurately as can be seen from the data in Table 6. Nevertheless, the statistical data seen there at pH 7.4 suggest that the reduction potentials of most Cu-Aβ complexes studied in this work fall within the potential window for catalytic oxidation of ascorbate by dioxygen (Table 6) and therefore should have no significant thermodynamic impact on their efficiency as catalysts for the aerial oxidation of ascorbate. In fact, there is no clear correlation between the calculated reduction potentials and the observed catalytic rates at the three pH values tested (Table 6).

It is apparent that the difference in the redox activity of the Cu centre in each Aβ16 peptide must be attributed primarily to differences in reaction kinetics that are closely related to the binding modes for both Cu(i) and Cu(u). As an efficient catalytic centre, the metal ion must be able to shuttle between different oxidation states with minimal reorganization energy,\(^\text{4,4,4}\) but this condition is often not met for Cu that prefers different ligands and binding modes for its two biologically accessible oxidation states. Blue copper proteins such as plastocyanin and azurin that accommodate robust redox active Cu centres feature conformational rigidity in the protein structures that impose an ‘entatic state’ to the Cu centre for efficient electron transfer.\(^\text{2,8}\)

Such ‘entatic states’ for Aβ peptides, however, are problematic, given their flexibility and unstructured nature. The coordination spheres for Aβ peptides are decided primarily by the coordination preferences of each oxidation state of the Cu centre, as demonstrated above and in previous studies.\(^\text{3,8}\) Consequently, the Cu centres in Aβ peptides are generally not efficient redox catalysts since only a very small fraction of Cu centres in equilibrium are pre-organized for electron transfer\(^\text{4,4}\) and the catalytic activity of ‘free’ Cu is inhibited upon binding to Aβ peptides.\(^\text{8,5,5,59}\)

At physiological pH, Aβ16 binds one equiv. of Cu(i) or Cu(u) with sub-nanomolar affinities (Fig. 1). Component I in the Cu(u) system is more reactive in the catalytic oxidation of Asc than component II for several reasons: (i) component I shares two His ligands with each of the three possible Cu(i) sites while component II contains only one His ligand; (ii) the main Cu(u) ligands for component I are flexible amino acid sidechains while the dominant Cu(u) ligands for component II are derived from the peptide backbone with rigid fused chelate rings; (iii) a Cu–N– bond arising from coordination to a negatively charged amide N– is highly covalent and any reactions involving such a chemical bond are usually slow.\(^\text{5,21,3,4}\) Consequently, component I is more reactive than component II, not only in redox catalysis but also in ligand substitution.\(^\text{5,6}\) These conclusions are consolidated by the observed reactivity orders for related peptides in ligand substitution\(^\text{6,2,1,3,3}\) and redox turnover (Table 6, eqn (10)).\(^\text{5}\)

In particular, substitution of Asp1 with His1 not only locks the Cu(u) binding mode to the IA component within the pH range 6–9 (Fig. 4a(i) and 6b), but introduces a new Cu(i) ligand in the N-terminus to be shared with the Cu(u) binding site. This leads to a considerable increase in the redox reactivity of the Cu centre. Interestingly, removal of the extra non-N-terminal ligand His14 from Aβ16-D1H leads to a further increase in redox activity (Table 6), presumably attributable to the increased similarity between the binding sites for both Cu(i) and Cu(u) in the Aβ16-D1H/H14A peptide.

**Summary and concluding remarks**

Monomeric Aβ peptides are flexible and unstructured. Both metal binding ligands and binding modes change dynamically with the Cu oxidation state and with reaction conditions. A range of chemical and spectroscopic studies were carried out on copper complexes of several Aβ16 peptides in which the two key residues for Cu(u) binding in the N-terminal region were varied. Key conclusions are as follows:

(i) At lower pH, the Aβ16 peptide binds Cu(u) predominantly as component IA in equilibrium with less stable component IB; at higher pH, these are converted to component II with transition $pK_a \sim 7.8$ (Fig. 1b and 4b).

(ii) Component I sites involve flexible amino acid sidechains common to the Cu(i) binding ligands, but the metal sites for component II are dictated by a deprotonated backbone amide nitrogen that cannot act as a Cu(i) ligand and is slow to dissociate.\(^\text{5,6,43}\) Consequently, component I is more reactive than component II in both metal substitution and redox catalysis.

(iii) Substitution of Asp1 with His provides an extra imidazole ligand for Cu(i) and also allows the N-terminal His1 to be part of a stable six-membered chelate ring for Cu(u) coordination. Consequently, the Cu(u) centre in Aβ16-D1H is locked into component IA with a dramatic increase in its redox activity.

(iv) Replacement of Ala2 with Pro suppresses component II relative to component I and increases the reactivity of the bound copper.

(v) Substitution of Asp1 with Ala restricts component I to the less stable IB form that is readily converted to component II at physiological pH with decreased catalytic reactivity.
This study provides insights into determinants that regulate the reactivity of copper toward Aβ peptides and may guide development towards a better understanding of the factors contributing to oxidative stress present within Alzheimer's disease.

**Abbreviations**

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<th>Abbreviation</th>
<th>Full Form</th>
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<tr>
<td>Asc</td>
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<tr>
<td>Bca</td>
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<td>Bsc</td>
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<td>Ches</td>
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<td>Dansyl peptide 2 (DAE[KDNS]RHDH)</td>
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**Acknowledgements**

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