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Properties of the major Zn²⁺-binding site of human alpha-fetoprotein, a potential foetal plasma zinc carrier†

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The foetal plasma protein α -fetoprotein (AFP) harbours a highaffinity zinc binding site that is likely involved in transport and delivery of essential zinc during foetal development. Based on a recent electron microscopy structure of AFP and aided by biophysical studies on an AFP-derived peptide, we present a refined 5-coordinate model for this site.

Human α -fetoprotein (AFP) is produced by the foetal liver and is the most abundant protein in foetal plasma (ca. 1-10 mg mL⁻¹) during certain stages of development. 1,2 Although AFP has long been used as a diagnostic tool in prenatal care and liver cancer detection, the physiological roles of this protein are only partially understood.3-5

Human AFP shares 39.4% sequence identity with human serum albumin (HSA), with each composed of three homologous domains (I-III).3,4 The plasma concentration of AFP decreases sharply after birth to trace amounts, with HSA replacing it as the major plasma protein.6 Both AFP and HSA bind many of the same ligands, which include fatty acids,^{7,8} bilirubin,9 and divalent metal ions,10,11 suggesting overlapping functions. The affinities of AFP for these cargoes are typically one to two orders of magnitude higher than those of HSA; this is also the case for the binding of Zn²⁺. ^{11,12}

An adequate Zn²⁺ supply is vital for normal foetal development in mammals, as demonstrated by the teratogenicity of maternal zinc deficiency. 13,14 However, the mechanisms by which Zn2+ is delivered to the foetus and taken up by foetal tissues are poorly understood, and no comprehensive model has been proposed. Several studies have reported an association between Zn²⁺ and AFP in amniotic fluid. ^{15,16} This supports

the hypothesis that AFP - like its relative HSA - binds Zn²⁺ under physiological conditions. The significance of HSA as the most critical player in regulating the availability of zinc in blood plasma is becoming increasingly clear. 17-19 We hypothesise that AFP plays a similar role in foetal plasma. The zincbinding ability of AFP has been confirmed in vitro, 11,20 with the more recent study reporting a single high-affinity Zn²⁺ binding site $(K_{\rm D} < 10^{-8} \text{ M})$ and at least four lower-affinity sites $(K_{\rm D} <$ 10^{-5} M). Given that the pH-independent K_D value for the highaffinity Zn²⁺ site on HSA is reported to be ca. 10⁷, ²¹ AFP is a plausible foetal plasma Zn²⁺ carrier and may participate in Zn²⁺ homeostasis during foetal development.

In 2023 the first 3D-structure of human AFP was elucidated using cryo-EM.22 More recently a second AFP structure was reported with Zn2+ and fatty acids bound, identifying the location and structure of a single Zn²⁺ binding site on AFP.²³ The Zn²⁺ site appears to be 4-coordinate with ligands provided by the sidechains of His4, His246, His250, and Asp262 residues (the amino acid numbering used relates to the mature protein (Fig. 1). Three of the four residues (His246, His250, and Asp262) reside within a helix-loop-helix region in domain II of the protein.

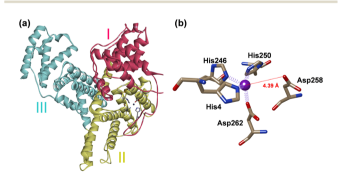


Fig. 1 (a) Cryo-EM structure of human AFP coloured by domain (I-III drawn from PDB: 8X1N). 23 (b) Close-up of the Zn2+-binding site. The site is composed of N-ligands from His4 (domain I), His246 and His250 and an O-ligand from Asp262 (domain II).

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A closer inspection of the Zn²⁺ binding site modelled in the cryo-EM structure (Fig. 1(b) and Fig. S1, ESI†) indicates that its coordination sphere is incomplete or distorted (the authors suggested that this may be due to the site being occupied by multiple different metal ions in their sample²³). In particular, the angles between the coordinating atoms more closely resemble a trigonal bipyramidal geometry, with His246 at one of the apices, and the other apex left empty. In addition, the angles between the aromatic ring planes of His4 and His250 and their bonds to the Zn²⁺ ion also deviate significantly from the expected 180°, and all four Zn-ligand bond lengths are much longer than expected. We attempted to re-model the site with a rotated Asp258 sidechain, and whilst this brought one of the O atoms within binding distance, comparison with the electron microscopy dataset (EMD) map did not support this model (not shown).

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To explore the Zn²⁺-binding properties of AFP further, we have studied the Zn²⁺ affinity of recombinant AFP in a system previously used for HSA to allow direct comparison. In addition, we have also characterised a synthetic peptide corresponding to the helix-loop-helix region. It was necessary to treat the expressed protein with EDTA and extensively dialyse it to remove metal ions (Zn²⁺, Cu²⁺ and Fe^{2+/3+}), presumably picked up during expression and/or purification. Isothermal titration calorimetry (ITC) was employed to examine Zn2+-binding to recombinant human AFP (Fig. 2). The binding of Zn²⁺ was exothermic (Table S1 and Fig. S2, ESI†) and the resultant data could be fitted using a "two sets of sites model" with the stoichiometry of site 2 fixed to 1 (Fig. 2). Models involving a single site or with varying site 2 stoichiometry gave unsatisfactory fits (Table S1, ESI†).

The data gave an apparent binding constant of $\log K_{\text{appITC}}(\text{Zn-}$ AFP) = 7.4 \pm 0.1 with a stoichiometry of 0.93 \pm 0.01 for site 1 and a $\log K_{\rm appitc}({\rm Zn\text{-}AFP}) = 4.8 \pm 0.1$ for site 2. Correction for competition with 50 mM Tris gave an apparent binding constant of $\log K_{\rm app}({\rm Zn\text{-}AFP})$ = 7.8 \pm 0.1, valid at pH 7.4, which translates to $K_{\rm D} = 1.9 \times 10^{-8}$ M. This is two orders of magnitude lower than the analogous apparent K_D for HSA (2.9 × 10⁻⁶ M, also corrected for competition with Tris). 12

The helix-loop-helix region that harbours three of the proposed Zn²⁺-binding ligands has additional residues with Zn²⁺-binding ability, including His248, Glu249, Asp255 and

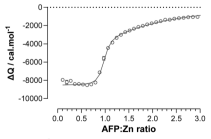


Fig. 2 $\,$ ITC data for ${\rm Zn}^{2+}$ binding to recombinant human AFP. Using a MicroCal VP-ITC instrument, 32 injections of 5 μ L of 300 μ M ZnCl₂ were delivered to samples of 20 μM AFP in buffer containing 50 mM Tris, 140 mM NaCl, pH 7.4 (298 K). The solid line represents the best fit to experimental points using a model involving two sets of sites.

Asp258. In the cryo-EM structure, a carboxylate oxygen of Asp258 is at a distance of only 4.4 Å to the Zn2+ ion (Fig. 1(b)). To explore whether alternative binding modes might be possible in this region, we designed a synthetic peptide corresponding to residues 245-263 of mature AFP. The peptide was synthesised commercially (95% purity; Genscript, Piscataway, NJ, USA). The N-terminus was acetylated, and the C-terminus was amidated to prevent the amine and the carboxylate from participating in metal binding. Cys251 and Cys259 form a disulfide bond in full-length AFP, whilst Cys252 forms a disulfide bond with Cys206. Therefore, Cys252 was substituted with a serine, to avoid formation of erroneous disulfide bonds and the remaining third Cys acting as a metal ligand (Fig. 3(a)). The identity and oxidation state of the AFP-peptide was confirmed using electrospray ionisation mass spectrometry (ESI-MS). The mass data (Table S2 and Fig. S3, ESI†) are consistent with those expected for oxidised peptide (i.e. disulfide bond formed), with the mono-isotopic neutral mass observed at 2128.99 Da, in excellent agreement with the theoretical mass (2128.92 Da). The close match between experimental and theoretical isotopic distributions (Fig. S3, ESI†) suggests that no reduced species were present.

The Zn²⁺-binding ability and stoichiometry were assessed using native ESI-MS and inductively-coupled plasma optical emission spectroscopy (ICP-OES). ICP-OES analysis gave a Zn-to-peptide ratio of 1.2 \pm 0.1. Native ESI-MS revealed the presence of two Zn-peptide species. The most abundant isotopic peak for the predominant species in the deconvoluted mass spectrum of the Zn²⁺-peptide sample had a neutral mass of 2193.10 Da (Fig. 3(b)), which matches the theoretical most abundant isotopic peak for the Zn₁-peptide metalloform (2192.83 Da). A second species, Zn₂-peptide (most abundant neutral mass 2257.07 Da; theoretical mass 2256.76 Da) was also observed in the mass spectrum as a minor constituent. This agrees with the ICP-OES analysis; thus, both methods indicated that a 1:1 species is formed predominantly, but also that more

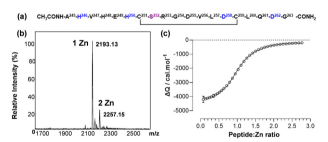


Fig. 3 (a) Amino acid sequence of the synthetic peptide designed to mimic the helix-loop-helix motif that forms the major part of the Zn²⁺ site of AFP. The superscript numbers give the positions of the corresponding amino residues in mature AFP. Residues proposed to be involved in Zn²⁺-binding are coloured in blue and the Ser substitution (in place of Cys252) is in purple. (b) Deconvoluted ESI-MS spectrum (MicrOTOF, Bruker Daltonics) of the Zn²⁺-loaded AFP-peptide (10 mM ammonium acetate, pH 7.4, 10% methanol). The signals correspond to two distinct forms of the metalated-AFP-peptide, with one or two Zn²⁺ ions bound. (c) ITC data for the addition of ZnCl $_2$ (333 μ M) to AFP-peptide (25 μ M) in 50 mM Tris buffer with 50 mM NaCl, pH 7.2. The solid line represents the best fit to experimental points using the model involving two sets of sites.

than one equivalent of Zn^{2+} can bind to the AFP-peptide with appreciable affinity.

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ITC was used to determine the affinity for Zn2+ binding to the AFP-peptide (Fig. 3(c)). As observed with recombinant AFP, the binding of Zn²⁺ to the AFP-peptide was exothermic (Fig. S4, ESI†). Using a "one set of sites" model to fit the data gave an apparent binding constant of $log K_{appITC}(Zn-peptide) = 5.7 \pm$ 0.1 with a stoichiometry of 1.05 \pm 0.01. Since ICP-OES and ESI-MS data indicated the possible existence of a second binding site, models involving two (sets of) binding sites were also explored. Using these models, apparent binding constants of $\log K_{\rm appITC}({\rm Zn-peptide}) = 6.1 \pm 0.1$ for site 1, and 4.6 ± 0.1 for site 2 were obtained. This model improved the curve fitting (goodness-of-fit improved from 75% for the single-site models to 91%), consistent with the existence of a second, much weaker binding site beside a strong primary site. It is however also important to note that the inclusion of a second site had only a relatively small effect on $\log K_{\text{appITC}}$ for the first site. Correction for the competition for Tris gave $\log K_{\text{app}} = 6.0$ or 6.4, respectively, translating to an apparent $K_D = 1.1 \times 10^{-6}$ M or 3.9×10^{-7} M. Hence, Zn^{2+} binding to the peptide was 20 to 60 times less strong than to the full-length AFP protein, but in a similar range as that to HSA. Given that HSA harbours a site with only three protein-derived ligands (two His and one Asp residue),24 binding to the peptide is also likely to occur through (at least) three ligands, also considering that the fourth proposed ligand, His4, is absent.

The interaction between Zn²⁺ and the AFP-peptide was further studied by 1D and 2D ¹H NMR spectroscopy. A series of ¹H NMR spectra were acquired for the apo-AFP-peptide at pH 7.1 and various temperatures between 278 K to 310 K (Fig. S5, ESI†). At 278 K, there was a significant increase in the dispersion of the backbone amide NH resonances, with sharper and more resolved peaks being discernible, presumably because of slowed-down structural dynamics. All subsequent NMR experiments were carried out at 278 K. 2D TOCSY and NOESY data were acquired for the apo peptide as well as for a sample after addition of 1 mol. eq. of Zn²⁺, at pH 7.1. An overlay of the fingerprint region of the TOCSY spectra is shown in Fig. 4.

Strikingly, upon Zn^{2+} addition, the dispersion of NH resonances increased greatly, with new NH signals appearing in both the low-field (*e.g.* 3 signals at >9 ppm) and the high-field (several signals below 7.5 ppm) region. This indicates that the binding of Zn^{2+} drives significant structural changes in the peptide, with most of the NH and $H(\alpha)$ protons having moved into a different electronic environment.

Sequential assignment for both apo-AFP-peptide and the peptide in presence of 1 mol. eq. Zn^{2+} were established through the 2D TOCSY and NOESY spectra (Tables S4 to S6, ESI†). The chemical shifts for $H(\alpha)$ protons in the apo AFP-peptide were compared with random coil shifts (Fig. S6(a), ESI†). All $H(\alpha)$ protons possess chemical shifts that are substantially below random-coil shifts; this is consistent with α -helical character²⁵ and in line with expectations based on the 3D structure of the full-length AFP protein. Sequential assignment of the spectra recorded in the presence of Zn^{2+} revealed the existence of two

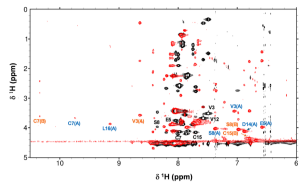


Fig. 4 Overlay of fingerprint region of 2D TOCSY NMR spectra of the AFP peptide in absence (black) and presence (red) of 1 mol. eq. $\rm Zn^{2+}$ (700 MHz, 50 mM [D₁₁]Tris–Cl, 50 mM NaCl, pH 7.1 and 10% D₂O, 278 K). Selected residues are labelled in black (apo form), blue (species A) and orange (species B).

distinct species (designated A and B). Despite the substantial changes in chemical shifts compared to the apo form (Fig. S7, ESI†), α -helical character is largely maintained in both species (Fig. S6(b) and (c), ESI†). Since the volumes of the cross-peaks for these two species do not differ significantly, it can be assumed that they exist at broadly similar concentrations. As only 1 mol. eq. of Zn^{2+} had been added, and no resonances for the apo-form remained, it follows that both species have just one Zn^{2+} ion bound. Whilst in theory, the two species could just be conformers, it appears more likely that the peptide harbours at least two Zn^{2+} binding sites with similar affinity, but (with consideration of the ITC results which indicated only one strong binding site) that they are mutually exclusive.

To explore possible binding site compositions, AlphaFold modelling was used to generate starting models for 1:1 complexes of the AFP peptide with Zn²⁺. Analysis of these models suggested that in the peptide, a site comprising His248, His250 and Asp262 may be most favourable, whilst another model showed Zn²⁺ in proximity to His250, Asp258 and Asp262. Three of these starting models were selected for further refinement. To generate final models, tetrahedral coordination was completed by a water molecule in all cases, and geometries were optimised by energy minimisation (UCSF Chimera v. 1.11, Amber forcefield). The models (Fig. S8(a)-(c), ESI†) suggest that the zinc-bound water molecule could form a hydrogen bond to the carboxylate group of either Asp255 or Asp258. In addition, we used the cryo-EM structure to create two starting models harbouring a site composed of His246, His250 and Asp262 and either one or two waters. Whilst the model with one water failed to yield reasonable geometries, the 5-coordinate model was viable Fig. S8(d), ESI†). Attempts to generate 4-coordinate models involving His246, His250, Asp255 and Asp258 simultaneously did not generate satisfactory geometries. Together, ITC, NMR and modelling results point towards the existence of two alternative binding sites with similar affinities, involving three peptide-derived ligands.

Finally, given the apparently incomplete coordination sphere in the cryo-EM model, we generated a model of full-length AFP

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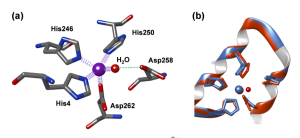


Fig. 5 Optimised geometry for the Zn²⁺-binding site in AFP. (a) The model was optimised using pdb 8X1N as starting point, and after filling the empty coordination site with a water molecule. Angles and distances are summarised in Table S5 (ESI†). (b) Overlay of original structure (blue) and optimised model (red). The model and images were generated in Chimera v. 1.11.26

with the original four ligands plus a water molecule in the empty site (Fig. 5). Refinement of this model did not require any discernible changes of the protein backbone, and only minor adjustments of the sidechain conformations, mainly to improve distances and angles (Table S7, ESI†). The resultant model has a distorted trigonal-bipyramidal geometry and features an H-bond from the zinc-bound water to Asp258.

The work presented supports the recent identification of a novel Zn²⁺ binding region on human AFP that may be important for the transport of zinc in the foetal circulation. In comparison to the major Zn²⁺-binding site in adult plasma, the site in AFP features an additional histidine residue, which is consistent with its higher binding affinity ($K_D = 1.9 \times 10^{-8}$ M vs. 2.9×10^{-6} M under identical conditions). The AFP concentration in foetal plasma (15-150 µM) is 4 to 40-times lower than that of HSA in adult plasma (ca. 650 µM). Using a typical total [Zn] of 14 μM^{27} and [AFP] = 15 μM , [Zn]_{free} = 196 nM can be calculated (based on $\log K = 7.77$). This reduces to less than 2 nM when 150 µM AFP are present. Although it is clear that other plasma proteins will further reduce [Zn]_{free}, these simple estimates have several implications for zinc distribution to and within the foetus: transport through membranes depends on $[Zn^{2+}]_{free}$; 19 therefore, at low AFP concentrations, it may be expected that transfer from foetal plasma into cells is enhanced. This will be advantageous for rapidly dividing cells and developing tissues within the foetus, which have a high requirement for zinc. At high [AFP], [Zn2+]free in the foetal circulation will be lower than that in adult plasma (At 650 μM HSA and 14 μM total Zn, [Zn]_{free} = 63 nM). It is possible that lower [Zn²⁺]_{free} on the foetal side of the placental syncytiotrophoblast promotes mother-to-foetus zinc transfer - somewhat analogously to the higher affinity of foetal haemoglobin toward oxygen.

It is also worth noting that the Zn2+-binding ability of albumin is modulated by fatty acid (FA) binding. 18,28,29 The AFP structure, 23 which has Zn2+ and FAs bound simultaneously, provides strong evidence that Zn²⁺-binding to the primary site on AFP is unlikely to be affected by FA binding. While FA

concentrations in foetal plasma are around 70-80% lower than in adult plasma,³⁰ owing to its lower concentration the average FA loading of AFP likely is significantly higher than that of HSA. The physiological reason behind the allosteric interaction between zinc and FA binding on albumin has remained enigmatic, but it would appear that this crosstalk is not part of foetal physiology, and that foetal plasma zinc speciation and trafficking is independent of FA status.

A deeper understanding of AFP and how it affects zinc speciation and trafficking may aid our understanding of how aberrant AFP levels or alteration in plasma Zn²⁺ concentrations may contribute to developmental disorders.

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Data availability

Data supporting this article are included in the ESI.†

Conflicts of interest

There are no conflicts to declare.

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