

RESEARCH ARTICLE

# Engineering a zinc binding site into the *de novo* designed protein DS119 with a $\beta\alpha\beta$ structure

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## ABSTRACT

Functional proteins designed *de novo* have potential application in chemical engineering, agriculture and healthcare. Metal binding sites are commonly used to incorporate functions. Based on a *de novo* designed protein DS119 with a  $\beta\alpha\beta$  structure, we have computationally engineered zinc binding sites into it using a home-made searching program. Seven out of the eight designed sequences tested were shown to bind  $Zn^{2+}$  with micromolar affinity, and one of them bound  $Zn^{2+}$  with 1:1 stoichiometry. This is the first time that metalloproteins with an  $\alpha$ ,  $\beta$  mixed structure have been designed from scratch.

**KEYWORDS**  $\beta\alpha\beta$ , *de novo*, design, folding, zinc-binding

## INTRODUCTION

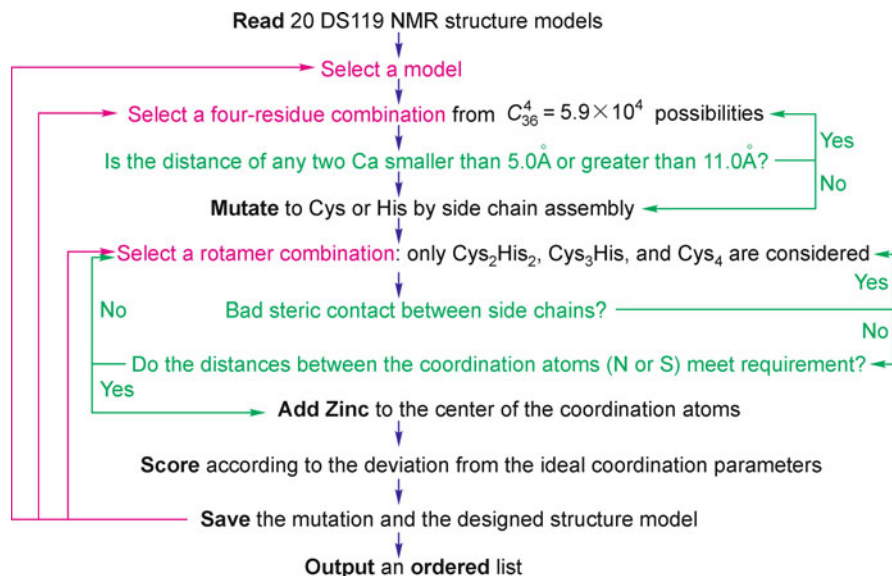
The important goal of protein design is to explore relationships between protein structures and functions. Various approaches can be used to incorporate functions in the designed protein scaffolds (Smith and Hecht, 2011), and metal binding sites have been a particularly interesting target for this purpose because the biological chemistry of metals is extremely rich (Holm et al., 1996; Lu et al., 2009). For example, by engineering different cofactors like  $Zn^{2+}$  (Handel et al., 1993),  $Fe^{2+}/Fe^{3+}$  (Kaplan and DeGrado, 2004), heme (Choma et al., 1994) and abiological chromophore (DPP) Zn (Fry et al., 2010) into the *de novo* designed four-helix bundles,

functions like phenol oxidation, electron transfer, and non-linear optical properties have been obtained.

Zinc binding sites are of special importance for both structural and functional reasons, as in zinc fingers (Berg and Shi, 1996), carbonic anhydrase (Christianson and Fierke, 1996), and zinc metalloproteases (Meinell et al., 1996). Using rational design and *in vitro* evolution, researchers have introduced zinc binding sites into different protein structures such as four-helix bundles (Regan and Clarke, 1990), antibody light chain (Wade et al., 1993), triple-strand coiled-coil (Kiyokawa et al., 2004), and retinal binding protein (Muller and Skerra, 1994). Recently, zinc binding proteins have been engineered for more sophisticated tasks, for example, protein conformation switches (Cerasoli et al., 2005; Ambroggio and Kuhlman, 2006), biosensors (Shults et al., 2003), purification tags (Pasquinelli et al., 2000), control of oligomerization states (Phillips et al., 2010), catalytic zinc sites (Nomura and Sugiura, 2004), or even regulating a bacterial signal transduction pathway (Dwyer et al., 2003). And zinc fingers combined with restriction endonucleases or recominases have emerged as an important tool for molecular biology for their application in editing genomes (Miller et al., 2007; Wu et al., 2007; Proudfoot et al., 2011). So the functions of zinc binding sites are widely explored in designed proteins.

To establish a valid and simple method for engineering zinc binding sites, we chose a designed small protein DS119 as the scaffold to incorporate zinc ions and developed a computational approach. DS119 was a *de novo* designed protein with a unique  $\beta\alpha\beta$  structure (Liang et al., 2009). It is highly thermally stable and folds quickly into a monomeric structure, providing an attractive target for functional

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**Figure 1. Computational process for identifying zinc binding sites in DS119.**

engineering. Similar to the  $\beta\beta\alpha$  zinc finger structure, the  $\beta\alpha\beta$  motif also contains one  $\alpha$ -helix and two  $\beta$  strands, though the two  $\beta$  strands are parallel compared to the anti-parallel  $\beta$  structure in zinc fingers. Zinc finger structures are commonly used in DNA binding proteins, which have inducible structure by binding to zinc ions. In the present study, we explored the possibility of introducing a zinc binding site into DS119 and studied the influence of the zinc binding on its structure and folding properties.

## RESULTS

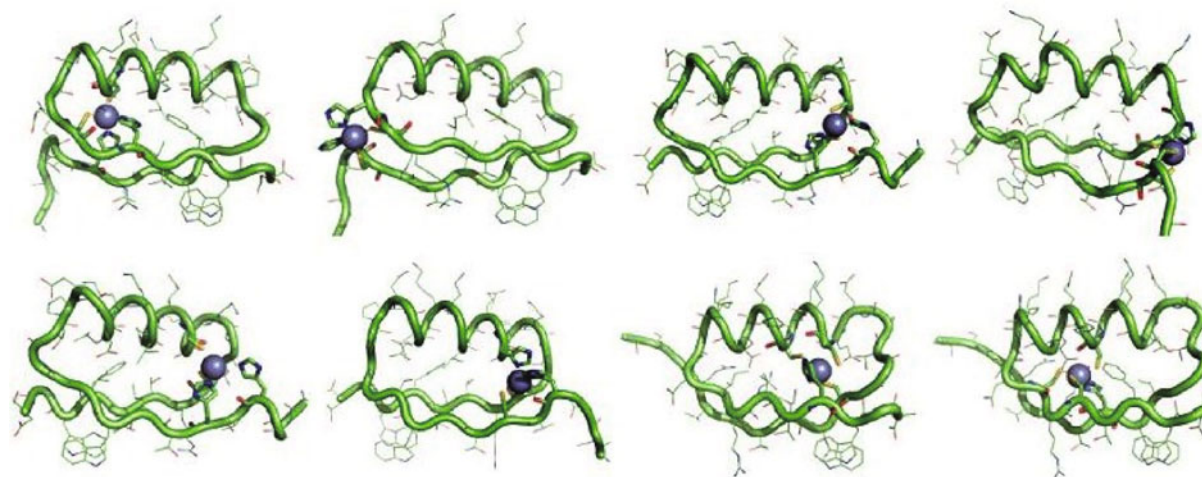
Twenty NMR structure models of DS119 (PDB code: 2K10) which embody the backbone flexibility were used as scaffolds to identify potential zinc binding sites. This approach is different from previous metalloprotein design programs which only search in one rigid conformation (Hellinga and Richards, 1991; Clarke and Yuan, 1995). Three types of zinc coordination sites were considered: Cys<sub>4</sub>, Cys<sub>3</sub>His and Cys<sub>2</sub>His<sub>2</sub>, and their ideal coordination parameters were obtained by statistical analysis of natural zinc binding proteins in the Protein Data Bank (Berman et al., 2000). The average values are as follows: S-Zn bond length is 2.32 Å, N-Zn bond length is 2.09 Å, C<sub>β</sub>-S-Zn angle is 107.3°, C<sub>γ</sub>-N<sub>δ</sub>-Zn or C<sub>ε</sub>-N<sub>ε</sub>-Zn angle is 128.4°, C<sub>α</sub>-C<sub>β</sub>-S-Zn dihedral angle can have three values: 111.1°, 184.4° and 269.4°, C<sub>γ</sub>-C<sub>ε</sub>-Zn-N<sub>δ</sub> or C<sub>δ</sub>-C<sub>ε</sub>-Zn-N<sub>ε</sub> is coplanar (Table S1). The computational process of identifying zinc binding sites consisted of three steps (Fig. 1). First, for each conformational scaffold the residues were substituted by Cys and His using the basic rotamer library of DeMaeyer et al. (1997), and the substitutions with steric contact were pruned. Next all the four-site combinations, in which the Ca

distances between any two residues were in the range of 11.0–5.0 Å, and were carefully examined to determine whether they could accommodate the Cys<sub>4</sub>, Cys<sub>3</sub>His or Cys<sub>2</sub>His<sub>2</sub> binding coordination. The examination used several rules: (1) Zn<sup>2+</sup> was placed on the center of four coordination atoms, (2) the coordination structure was scored according to its deviation from ideal parameters (see supplementary material), and (3) the structure models were ranked by the score. Finally the models with similar sequences were clustered and 20 different design modes (Table S2) were obtained.

According to the top-scoring solutions, eight DS119 mutants (called BABZ) were expressed, purified and tested for their zinc binding affinities and structural properties (Table 1 and Fig. 2). Isothermal titration calorimetry (ITC) was used to check for possible Zn<sup>2+</sup>-protein interactions, and seven out of eight BABZ proteins were shown to bind Zn<sup>2+</sup> with a medium tendency. Their dissociation constants (K<sub>d</sub>) were in the range of 2–20 μmol/L (Table 2). Based on the ITC results, BABZ5 was chosen for further investigation as the titration curve indicated it bound Zn<sup>2+</sup> with 1:1 stoichiometry (Fig. 3B). BABZ5 was designed to bind Zn<sup>2+</sup> with two N-terminal His, one Cys on the helix, and another Cys on the  $\beta$  strand. It showed good secondary structure on CD, preserving the  $\beta\alpha\beta$  fold of DS119 (Table 2 and Fig. 3C). In gel filtration chromatography the peak of BABZ5 without Zn<sup>2+</sup> shifted towards the aggregation state, and upon adding Zn<sup>2+</sup> in the running buffer it moved to the monomer position (Table 2 and Fig. 3D). The peak of BABZ5 also showed trailing edge boundary and expansion towards the monomer state. We thought this was due to the equilibrium between the bound and unbound state, since the binding affinity of BABZ5

Table 1 Sequences of DS119 and BABZ proteins

DS119	GSGQV	RTIWV	GGTPE	ELKKL	KEEAK	KANIR	VTFWG	D
BABZ1	GSGQV	HTIWV	GGTPE	ELKKL	KECAK	KANCR	HTFWG	D
BABZ2	GSGHC	RTIWV	GGTPE	ELKKL	KEEAK	KAHCR	VTFWG	D
BABZ3	GSGHV	HTIWV	GGTPE	ELKKL	KEEAK	KCNCR	VTFWG	D
BABZ4	GSGCC	RTIWV	GGTPE	ELKKL	KEEAK	KHNCR	VTFWG	D
BABZ5	GSGHV	HTIWV	GGTPE	ELKKL	KECAK	KANCR	VTFWG	D
BABZ6	GSGHV	CTIWV	GGTPE	ELKKL	KEHAK	KANCR	VTFWG	D
BABZ7	GSGQV	RTIWC	GGTPE	ECKKC	KEEAK	KANIR	VTHWG	D
BABZ8	GSGQV	RTHWV	GGTPE	ELKKC	KEEAK	KANCR	CTFWG	D



**Figure 2.** Structure models of BABZ proteins. Upper row from left to right: BABZ1, 2, 3 and 4, lower row: BABZ5, 6, 7 and 8.  $Zn^{2+}$  is indicated as a silver sphere, and coordination residues (histidine or cysteine) are shown as sticks.

**Table 2** CD, gel filtration and ITC results of BABZ proteins

	BABZ1	BABZ2	BABZ3	BABZ4	BABZ5	BZBZ6	BZBZ7	BZBZ8
Aggregation state without $Zn^{2+}$	1.41	0.86	0.90	0.89	1.24	NA	1.42	1.40
Aggregation state with $Zn^{2+}$	1.41	NA <sup>a</sup>	NA	NA	1.03	NA	1.41	1.40
Secondary structures without $Zn^{2+}$ (%)	42	91	92	99	98	94	43	34
Secondary structures with $Zn^{2+}$ (%)	40	91	95	100	98	63	45	32
$N^b$	0.65	0.51	NA <sup>c</sup>	0.55	0.93	0.40	0.40	0.72
$K_d$ ( $\mu\text{mol/L}$ )	22	15	NA	7.5	2.2	8.8	4.5	2.3

<sup>a</sup> NA means no elution peak was observed in gel filtration.

<sup>b</sup> N is the molar ratio indicating how many  $Zn^{2+}$  ions interacted with one protein.

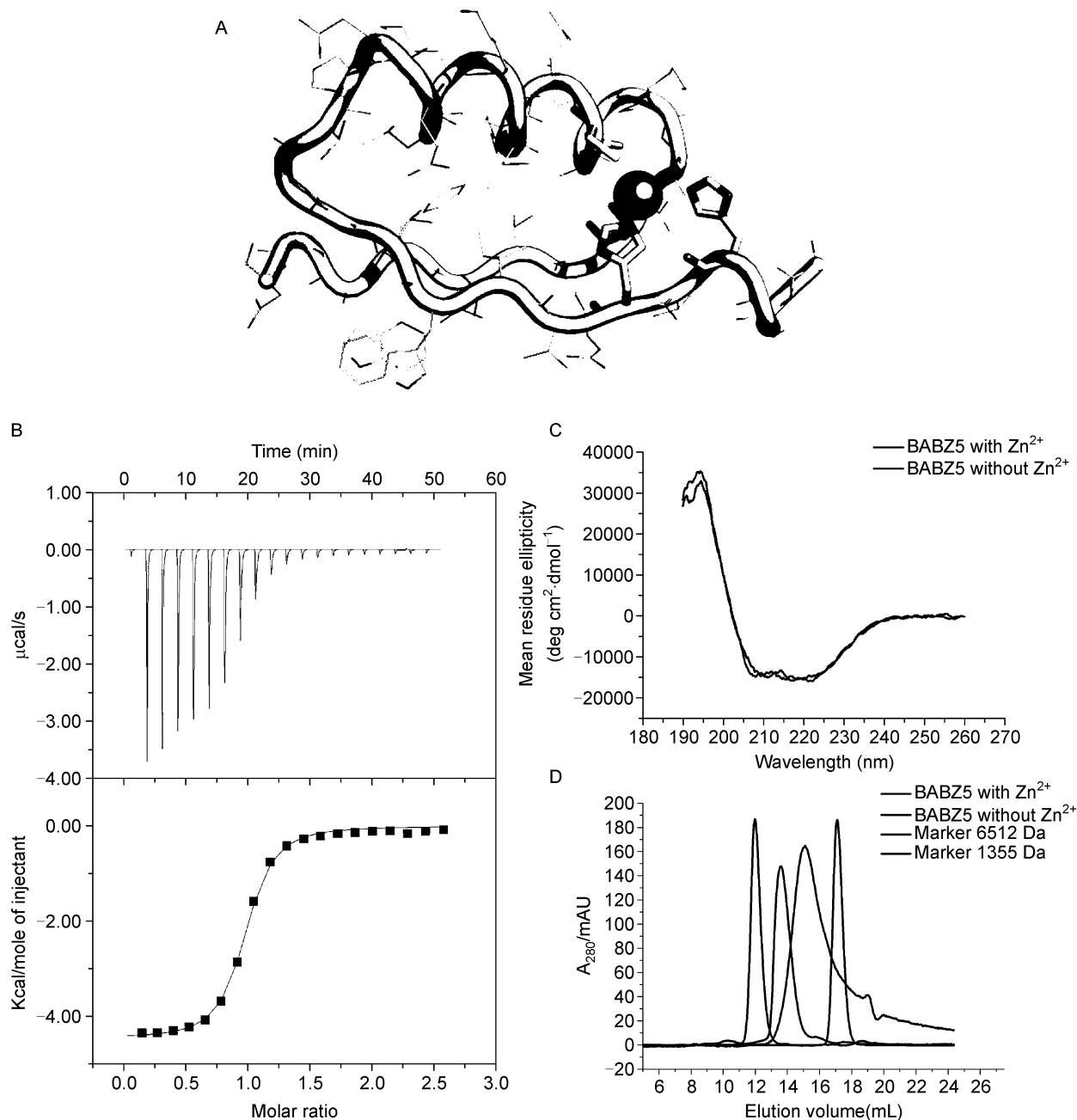
<sup>c</sup> NA means the interaction of BABZ3 and  $Zn^{2+}$  is too weak to be measured by ITC.

for  $Zn^{2+}$  was not extremely high ( $K_d$  was 2.2  $\mu\text{mol/L}$ ). The trailing edge is the result of a systematic increase in the migration rate across the advancing boundary, which in turn, is due to rapid self-association equilibrium. This phenomenon has been used in large-zone chromatography to determine the association constants (Stevens, 1986; Winzor, 2003).

There was no significant difference between the secondary structures of BABZ5 with and without  $Zn^{2+}$ , as indicated by CD in the far-UV region (Fig. 3C). However, their thermodynamics showed dramatic differences (Fig. 4). The CD

signal at 222 nm was used to monitor the unfolding and refolding process of BABZ5. In the absence of  $Zn^{2+}$ , partially unfolded BABZ5 could not fold back to its original thermodynamic state, as indicated by the differences between its unfolding and refolding curves. After BABZ5 binds to  $Zn^{2+}$ , these two curves were identical, indicating the guiding role of  $Zn^{2+}$  during its folding process. Thus, BABZ5 binding with  $Zn^{2+}$  facilitated smooth transition from the unfolded state to the folded state.

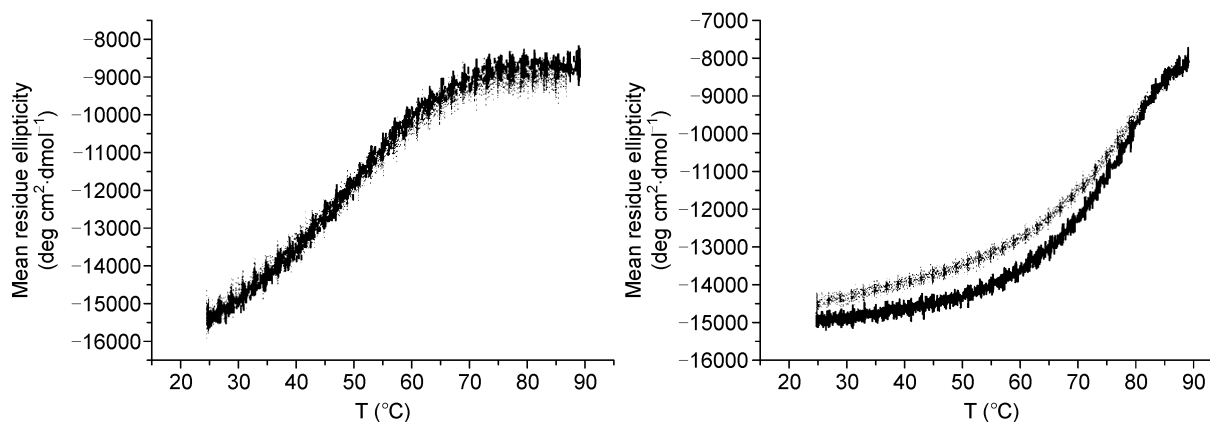
Other BABZ proteins, which also showed zinc binding



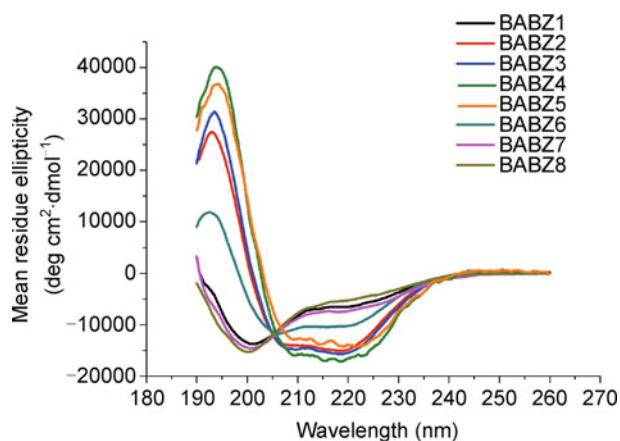
**Figure 3. Structural and Zn<sup>2+</sup>-binding properties of the designed BABZ5 protein.** (A) Structure model of BABZ5 protein. Zn<sup>2+</sup> is indicated as a silver sphere, and coordination residues (histidine or cysteine) are shown as sticks. (B) ITC measurement of BABZ5 and Zn<sup>2+</sup> interactions. (C) CD spectra of BABZ5 with and without Zn<sup>2+</sup>: BABZ5 in ZnSO<sub>4</sub> buffer (20 mmol/L Tris-HCl pH 7.3, 2 mmol/L ZnSO<sub>4</sub> and 100 µmol/L TCEP) (—), BABZ5 in EDTA buffer (20 mmol/L Tris-HCl pH 7.3, 1 mmol/L EDTA and 100 µmol/L TCEP) (—). (D) Gel filtration analysis of BABZ5 with and without Zn<sup>2+</sup> using the same buffer as in CD, but adding 100 mmol/L NaCl. ITC, isothermal titration calorimetry.

affinities, falls mainly into two categories: BABZ2 and BABZ4 preserved the βαβ fold, and BABZ1, BABZ7 and BABZ8 were unstructured (Fig. 5). For the unstructured BABZ proteins, zinc-binding changed neither their secondary structure nor aggregation states. For BABZ2 and BABZ4, the molar ratios

for the interactions between BABZ proteins and Zn<sup>2+</sup> were not 1:1 as designed, according to ITC (Fig. S3). Since their coordination residues are located on flexible N-terminals, it is possible that the coordination sites were composed of residues from two monomeric proteins and the binding of



**Figure 4.** Thermal denaturation curves of BABZ5 with and without  $Zn^{2+}$ . Left: BABZ5 in  $ZnSO_4$  buffer. Right: BABZ5 in EDTA buffer. Thermal unfolding was measured from 25°C to 90°C (—), and refolding was measured from 90°C to 25°C (—). CD signals at 222 nm were monitored.



**Figure 5.** CD spectra of BABZ proteins with  $Zn^{2+}$ . Experiments were carried out in 20 mmol/L Tris-HCl buffer (pH 7.3) containing 2 mmol/L  $ZnSO_4$  and 100  $\mu$ mol/L TCEP.

$Zn^{2+}$  could induce their oligomerization. So compared to BABZ5, they cannot be regarded as successful designs.

## DISCUSSION

Although nature has used zinc fingers with  $\beta\beta\alpha$  structure to incorporate  $Zn^{2+}$ , the  $\beta\alpha\beta$  motif with the same number of secondary structure elements has never been observed for this function. By using the previously designed DS119, we engineered zinc binding proteins with a  $\beta\alpha\beta$  structure. Compared to the binding induced folding process of zinc fingers, the conformation changes of BABZ5 after binding  $Zn^{2+}$  were not significant, partly due to the highly thermally stable structure of DS119 (Liang et al., 2009). It is challenging to design functional sites in a small *de novo* designed protein, and during the process of zinc binding sites engineering,

much can be learned about its sequence, structure and folding relationship.

In order to engineer a zinc binding site in DS119, two to four mutations must be made. Which residues should be replaced is vital for a successful design. For example, BABZ1 resembled the most perfect tetrahedral coordination sites among all design modes, but the  $His_2/Cys_2$  mutations damaged the protein structure, and binding to  $Zn^{2+}$  could not recover the  $\beta\alpha\beta$  fold. Similarly, in BABZ7 and BABZ8, the mutations essentially destroyed the hydrophobic core. So the overall folding involves a delicate interplay of hydrophobic packing, hydrogen bonding, and metal binding (Li et al., 2007, 2008). In addition, if the coordination residues are located on flexible terminals or loops, as in BABZ2 and BABZ4, zinc binding-induced aggregation may occur, especially for the small protein like  $\beta\alpha\beta$ , so the entropy cost to fix the coordination residues at specific conformations should be considered, as zinc binding in proteins is an entropy-driven process (Reddi et al., 2007). It has been demonstrated that the primary interactions are enough to produce potentially useful zinc binding affinity (Wade et al., 1993), and this agrees with our design work.

It is worth noting that the  $T_m$  value for the unfolding process of BABZ5 with  $Zn^{2+}$  was clearly lower than that without  $Zn^{2+}$ , which indicates zinc-binding stabilizes the unfolded state more than it does to the folded state. The unfolded state is hard to be considered apparently and is not included in our computational program. Nonetheless this phenomenon provides opportunities to study the folding mechanism of the  $\beta\alpha\beta$  motif, since the  $T_m$  values were reduced to a reasonable range (about 50°C) compared to DS119 (above 80°C), and the unfolding process is cooperative, which makes it more suitable for fast kinetic experiments like laser induced T-jump (Dyer et al., 1998).

Different methods have emerged for engineering novel

metalloproteins. The simplest way is to analyze protein scaffolds empirically and introduce cysteine-rich coordination sites. Several  $\alpha$ -helical coiled-coil metalloproteins capable of binding Co(II), Fe(II), Zn(II), Cd(II), Hg(II), and As(II) have been designed this way (Matzapetakis et al., 2002; Petros et al., 2006). More complex coordination sites were engineered with the aid of computational searching, for example, the Cys<sub>2</sub>-His<sub>2</sub> zinc binding sites and Fe<sub>4</sub>-S<sub>4</sub> sites (Lu et al., 2001). Retrostructural analysis for designing diiron binding protein is the most complex approach by far (Lombardi et al., 2000). The natural diiron binding site was analyzed first to obtain the desired chelation geometry, the symmetry of the structure, and other rules for forming the binding site. Then the design was accomplished in a hierarchical manner: coordinated side chains were placed first, the residues for coordination stabilization and hydrophobic core came next, and last the polar residues and loops were fixed. Backbone conformation variation was considered in this computation process. Our work of designing zinc-binding  $\beta\alpha\beta$  small proteins shares some characteristics with the third method.

In conclusion, we have successfully engineered a zinc binding site in the *de novo* designed protein with a  $\beta\alpha\beta$  structure. The small zinc binding proteins provide valuable targets for protein folding research, and can be further engineered to target DNA sequences, or catalyze chemical reactions. This work provides an approach to engineering functional proteins with more complex structures.

## MATERIALS AND METHODS

### Cloning, protein expression and purification

The genes encoding BABZ proteins were synthesized by Invitrogen (Beijing, CN) and cloned into pGEX<sub>4T-1</sub> vectors (GE Healthcare). The proteins were expressed in *Escherichia coli* Rosetta cells for 6 h at 30°C and purified using standard GST-fusion proteins purification protocols. Using PBS (100 mmol/L NaCl, 40 mmol/L Na<sub>2</sub>HPO<sub>4</sub>/NaH<sub>2</sub>PO<sub>4</sub> pH 7.3) containing 2 mmol/L DTT as running buffer and 10 mmol/L glutathione in Tris-HCl pH 8.0 as elution buffer, solutions containing the GST-fusion BABZ proteins were collected, and thrombin (Sigma) was added at 4°C to remove the N-terminal GST protein.

The samples containing BABZ proteins, glutathione, and thrombin were loaded onto an Agilent Zorbax 300SB-C18 column and purified by reversed-phase HPLC. Gradient-elution using water and acetonitrile was performed on the Agilent 1100 series HPLC. The peak corresponding to BABZ proteins was eluted at 45%–50% acetonitrile, and then collected for lyophilization. The purity of each protein sample was verified by high resolution mass spectrometry (Fourier Transform Ion Cyclotron Resonance Mass Spectrometer, Bruker APEX IV, USA).

### Circular Dichroism

Circular Dichroism spectra (CD) were measured on a MOS 450 AF/CD (Biologic, France) at room temperature, using 1 mm quartz cuvettes for the far-UV region (190 nm to 250 nm). Band width and

scan speed were set as 0.5 nm and 100 nm/min. Protein samples were dissolved to a final concentration of 0.2 mg/mL in 20 mmol/L Tris-HCl buffer (pH 7.3) containing 2 mmol/L ZnSO<sub>4</sub> or 1 mmol/L EDTA, and 100  $\mu$ mol/L TCEP.

The secondary structure content was calculated using DS119 as standard. The normalized CD signal of DS119 at 222 nm was  $-15,410 \text{ deg cm}^2 \cdot \text{dmol}^{-1}$  and this value was set as 100%. The CD signal of BABZ proteins at 222 nm divided by  $-15,410 \text{ deg cm}^2 \cdot \text{dmol}^{-1}$  gave their secondary structure content.

Thermal denaturation curves of BABZ proteins were also obtained on the MOS 450 AF/CD using the Peltier accessory with 10 mm quartz cuvettes. The CD signal at 222 nm was recorded. The protein concentration was 0.02 mg/mL in the same buffers as in far-UV CD, and heating was performed at 1°C/min speed from 25°C to 90°C.

### Gel filtration chromatography

Protein samples were dissolved in 20 mmol/L Tris-HCl buffers (pH 7.3) containing 2 mmol/L ZnSO<sub>4</sub> or 1 mmol/L EDTA, and 100 mmol/L NaCl to final concentrations of 50  $\mu$ mol/L, and then loaded onto a Superdex<sup>TM</sup> Peptide 10/300 GL column (GE Healthcare). The column was eluted using the same buffer with a flow rate of 0.4 mL/min and UV absorption at 280 nm was monitored.

The molecular markers are cytochrome C (12.4 kDa), aprotinin (6512 Da), and vitamin B12 (1355 Da). The lg(MW)–V plot (MW: molecular weight, V: elution volume) was obtained for the column and used to calculate the apparent molecular weight of BABZ proteins. The apparent MW divided by the calculated MW gave their aggregation states.

### Isothermal titration calorimetry

An iTC200 Microcalorimeter (MicroCal, USA) was used to measure the binding affinities ( $K_d$ ) between Zn<sup>2+</sup> and BABZ proteins: 400  $\mu$ mol/L BABZ protein dissolved in 100 mmol/L HEPES (pH 7.3), 50 mmol/L NaCl and 100  $\mu$ mol/L TCEP was loaded into the cell, and 4 mmol/L ZnSO<sub>4</sub> in the same buffer was added automatically by syringe. The titration was performed at 25°C with an initial 0.4  $\mu$ L injection in 0.8 s, followed by nineteen 2  $\mu$ L injections in 4 s. The spacing between injections was 150 s, and the stirring speed during the titration was 1000 rpm. Data were collected every 5 s, and then analyzed using MicroCal Origin software by fitting to the single-site binding model. Correction for ligand dilution and other nonspecific interactions was carried out by performing a control experiment, using the same ZnSO<sub>4</sub> and HEPES buffers, and the heat effect was subtracted in the original data.

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