Two zinc-binding domains in the transporter AdcA from *Streptococcus pyogenes* facilitate high-affinity binding and fast transport of zinc

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Two AdcA zinc-binding domains facilitate zinc transportation

In this study, we applied molecular modeling and molecular dynamics simulation to assess the dynamic structure and functional features of the two domains of Streptococcus AdcA. Based on the computational results, we propose a sequential structural stabilization model for the two fused domains that can conduct an interdomain conformational change when bound to zinc, an exceptional feature that enhances zinc uptake efficiency in zinc-deficient environments as compared with single-domains proteins. We further experimentally validated the predicted features and the double-domain organization that endows Streptococcus with unique survival advantages in zinc-depleted environments.

Results

Double-domain AdcA homologs are conserved and overrepresented in Streptococcus

We first constructed the S. pyogenes AdcA structure using homology modeling, taking as templates the most homologous crystal structures of B. subtilis Bsu-YcdH (43.9% sequence identity to the N-terminal domain; Protein Data Bank code 2O1E) and E. coli San-YodA (47.3% sequence identity to the C-terminal domain; Protein Data Bank code 1TXL) (Fig. S1B). The constructed AdcA structure contains the amino acid residues 31–515 (Fig. 1A), which is almost full-length except for the predicted signal peptide (amino acids 1–30; Fig. S1A). The modeled structure gained a discrete optimized potential energy score of −49,727.9, indicating that this structure was approximatively at the optimal low-energy state (20). A Ramachandran plot indicated that the modeled three-dimensional structure was reasonable (Fig. S1C). Virtual docking analysis suggested a zinc-binding center in the N-terminal domain (His-36, His-122, His-186, and Glu-261) and a zinc-binding center in the C-terminal domain (His-436, His-445, and His-447) (Fig. 1B). The two predicted zinc-binding sites, located in the highly conserved area, were in conformity with the principle of ConSurf evolutionary conservation patterns (21–23).

To determine the universality of the double-domain organization of AdcA homologs in bacteria, we compared S. pyogenes AdcA against 3,449 completely sequenced bacterial genome sequences in the NCBI database using the Blastx tool. We found 649 proteins with more than 30% homology to AdcA. Among these, 31 proteins were single-domain zinc-binding proteins homologous to the C-terminal domain of AdcA; blue dots, single-domain proteins homologous to the N-terminal domain of AdcA; gray dots, double-domain proteins homologous to the entire AdcA. D, phylogenetic tree of the bacterial species with double-domain AdcA homologs.
tances were maintained below 0.22, indicating that these double-domain zinc-binding proteins were highly conserved in *Streptococcus* during evolution from the common ancestor as compared with the other species (Fig. 1D). These structural results implied that *Streptococcus* AdcA possesses a unique ability for zinc uptake.

**The N-terminal domain stabilizes the protein structure more strongly than does the C-terminal domain upon zinc binding**

To predict the properties of *S. pyogenes* AdcA, we performed molecular dynamics (MD) simulations on the homology model. Independent replications of the MD simulation using random initial atomic velocities resulted in similar trajectories. All trajectories reached equilibrated conformation within 5 ns. These timescales are 1–2 orders of magnitude shorter than the timescale for forming any secondary structure. To exclude the formation of new secondary structures, we further performed 300-ns MD simulations for apo-AdcA and Zn$_2$-AdcA, a timescale that is sufficient to represent the conformational change of complex proteins (24, 25). Compared with the initial state, there was no obvious change in conformation or hydrogen bonds in the equilibrium state (Fig. S5, A–C). Moreover, the trend of change in secondary structures, caused by binding of the two zinc ions, in MD simulations was consistent with CD spectrum experimental results (Fig. 2C and Table S2). The merged results from the initial structure and the average structure of the equilibrium state are shown in Fig. S5D. These show that the simulated structure is close to its native state in physiological solutions.

Next, we simulated AdcA structures without zinc atoms (apo-AdcA), with zinc in the N-terminal binding center (Zn-N-AdcA) alone, with zinc in the C-terminal binding center (Zn-C-AdcA) alone, and with zinc in both binding centers (Zn$_2$-AdcA) (Fig. 2A). Trajectories of 40-ns MD simulations revealed the highest root mean square displacement (RMSD) for apo-AdcA, indicating that zinc binding stabilizes the protein structure (Fig. 2B). Indeed, Zn$_2$-AdcA had the lowest RMSD, indicating that it has the highest rigidity among all four forms. This was confirmed by CD spectra (Fig. 2C): the unordered fraction of AdcA was reduced from 19.1 ± 0.14 (apo-AdcA) to 11.2 ± 0.56% (Zn$_2$-AdcA) when saturated by zinc, and the helix fraction slightly increased from 53.2 ± 0.62 to 58.8 ± 0.14% (Table S2).

However, the structural rigidity of the two zinc-binding centers deviated remarkably. We calculated the root mean square fluctuation (RMSF) of the zinc-binding residues; a lower RMSF indicates higher rigidity. The four residues of the N-terminal domain (His-36, His-122, His-186, and Glu-261) showed reduced RMSF values after binding zinc, whereas two of the three residues in the C-terminal domain (His-436, His-445, and His-447) showed increased RMSF values (Fig. 2D) (24). These results indicated that zinc stabilizes only the N-terminal domain and that flexibility of the C-terminal domain is not affected by zinc. This was echoed by the RMSF of all C-α atoms of the two domains (Fig. 2E): 132 of 291 residues (45.5%) in the N-terminal domain showed reduced RMSF after binding zinc, whereas 45 of 194 residues (23.9%) in the C-terminal domain showed reduced RMSF after binding zinc.

Moreover, the radius of gyration ($R_g$) within the period of equilibrium also suggested a more flexible structure of Zn-C-AdcA, Zn-N-AdcA, and Zn$_2$-AdcA compared with apo-AdcA (Fig. 2, F and G). To further analyze the $R_g$ values between these three curves, we used symbolic aggregate approximation to evaluate the statistical differences in detail (Fig. S6). The major goal of the symbolic aggregate approximation algorithm is to convert time-series data to a symbolic representation, e.g. a $<$ b $<$ c $<$ d. Then, the mean value of each section is calculated (26). This method can be used to accurately distinguish differences in the data (27, 28). The strings, converted from the $R_g$ values of Zn-C-AdcA, Zn-N-AdcA, and Zn$_2$-AdcA, were shown as dcbbcbcb, cbacbacb, and cbbaaabc, respectively (Fig. S6). Thus, the $R_g$ of Zn-C-AdcA was considerably greater than that of Zn-N-AdcA, and Zn-N-AdcA had a similar $R_g$ compared with Zn$_2$-AdcA (Fig. 2, F and G, and Fig. S6). Therefore, we posit that the N-terminal domain stabilizes the protein structure more than does the C-terminal domain and possesses a higher affinity for zinc. Interestingly, zinc binding in one domain influenced rigidity in the other domain. Zinc binding to the N-terminal domain (Zn-N-AdcA) reduced the RMSF of 18.7% of the residues in the C-terminal domain, and Zn-C-AdcA reduced the RMSF of 12.4% of the residues in the N-terminal domain (Fig. 2E). This suggested that the two domains, each with distinct zinc-binding properties, may synergize upon zinc binding, creating new conformational features that do not exist in the single domains.

**Affinity and speed: New features emerge by synergy of the two domains**

To validate the above mentioned postulations, we mutated key binding residues to alanine to abolish the N-terminal (H36A/H122A/H186A/E261A) or C-terminal (H436A/H445A/H447A) zinc-binding sites while maintaining the full lengths of N-AdcA and C-AdcA. To completely remove the interactions between the two domains, we created an N-lobe (residues 31–321) and a C-lobe (residues 322–515) to mimic the single-domain AdcA that exists in most bacterial species (Fig. 3A). These mutants and the WT AdcA were expressed and purified (Fig. S3, A and B). The inductively coupled plasma MS measurement showed that all the purified proteins did not contain any metal ions.

To detect the strength differences of the two binding domains of AdcA, a 4-(2-pyridylazo)resorcinol (PAR) competition test with Zn$_2$-AdcA was performed. As shown in Fig. 3B, PAR could only capture one zinc from Zn$_2$-AdcA under normal conditions, whereas it could capture other zinc atoms only under harsh denaturation conditions, e.g. 6 M guanidine hydrochloride. These results of the competition test showed that the differences in zinc binding strength of the N- and C-terminal domains are significant.

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5 The abbreviations used are: MD, molecular dynamics; PAR, 4-(2-pyridylazo)resorcinol; TPEN, $NN'NN''N$-tetrakis(2-pyridylmethyl)ethylenediamine; RMSF, root mean square fluctuation; RMSD, root mean square displacement; $R_g$, radius of gyration; ITC, isothermal titration calorimetry; SASA, solvent-accessible surface area.
Two AdcA zinc-binding domains facilitate zinc transportation

We further experimentally determined the zinc-binding affinities of WT and mutant AdcA using isothermal titration calorimetry (ITC), and the calculated values of $N$, $K_D$, $\Delta H$, and $\Delta S$ are shown in Fig. 3, C–G. The $K_D$ of the WT AdcA for Zn$^{2+}$ was measured as $27 \pm 7.3$ nM, much lower than that of any mutant (Fig. 3C). The independently expressed N-lobe and C-lobe had $K_D$ values of $93 \pm 8.2$ nM and $2.6 \pm 0.3$ µM, respectively (Fig. 3, F and G), confirming that the N-lobe has higher...
Two AdcA zinc-binding domains facilitate zinc transportation

zinc-binding affinity than does the C-lobe. Moreover, the full-length expressed mutants N-AdcA and C-AdcA had dissociation constants of 390 ± 10.5 nM and 1.1 ± 0.2 μM (Fig. 3, D and E). Notably, the independently expressed N-lobe had a stronger binding affinity for Zn^{2+} than did full-length N-AdcA, implying that the two domains interact with each other, influence each other’s conformation, and thus regulate zinc affinity. Thermodynamic data on enthalpy changes (ΔH) and entropy changes (TΔS) of AdcAs were detected by ITC. For zinc binding, WT AdcA, C-AdcA, N-AdcA, the N-lobe, and the C-lobe were, respectively, found to have ΔH values of −17.6 ± 5.1, −10.2 ± 1.04, −11.4 ± 4.4, −9.9 ± 0.5, and −12.2 ± 1.1 kcal/mol and

Figure 3. Zinc-binding affinity of AdcAs and PAR competition for zinc ions. A, the five protein constructs used to determine the biophysical properties. The constructs were overexpressed in E. coli BL21 (DE3) strain and purified. B, UV-visible spectra of PAR, which competes for zinc, for Zn-C-AdcA, Zn-N-AdcA, and Zn_{2}-AdcA in the presence or absence of guanidine hydrochloride (Gdn.HCl). C–G, isothermal titration calorimetry binding curves of WT AdcA (C), C-AdcA (D), N-AdcA (E), the N-lobe (F), and the C-lobe (G) at 25 °C. The parameters N, K_{D}, ΔH, and ΔS are shown in the diagrams.
Two AdcA zinc-binding domains facilitate zinc transportation

Figure 4. Biochemical, thermodynamic, and kinetic characterization to verify protein stability. A, kinetics of AdcA binding to zinc. Time-dependent reactions were created using 10 μM apo-form proteins mixed with 40 μM Zn(PAR)₂. B, proteinase K sensitivity of WT AdcA. Proteins (15 μg) were subjected to proteinase K (30 μg/liter) for 0, 1, 3, 5, 7, 10, and 30 min. C, proteinase K sensitivity of C-AdcA. D, thermal stabilities of WT AdcA, Zn₂-AdcA, and Zn-C-AdcA. Thermal unfolding transitions were monitored using far-UV CD spectra at 223 nm.

The ΔS values of -7.3 ± 0.7, -2.0 ± 0.3, -2.7 ± 0.4, -0.3 ± 0.07, and -4.5 ± 1.5 kcal/mol. The values of ΔH and ΔS indicated that zinc binding to the five models of AdcA is both enthalpically and entropically favorable with ΔH being the main driving factor.

Next, we measured the zinc binding kinetics of AdcA. As expected, N-AdcA, with higher affinity-bound zinc, had very fast kinetics and a two-component reaction (k₁ = 45.45 ± 0.07 s⁻¹ and k₂ = 0.93 ± 0.02 s⁻¹, A₁ = 0.099 ± 0.000 and A₂ = 0.161 ± 0.002). In contrast, C-AdcA, with a less rigid structure, bound zinc at a much lower rate and had a second-order reaction. The fast reaction was a minor reaction with an amplitude of A₁ = 0.154 ± 0.026 and k₁ = 5.02 ± 0.03 s⁻¹, whereas the major reaction had the kinetic parameters A₂ = 0.271 ± 0.036 and k₂ = 0.15 ± 0.01 s⁻¹ (Fig. 4A). WT AdcA, with two binding centers, revealed a similarly fast process (k₁ = 41.05 ± 0.01 s⁻¹) as compared with N-AdcA and a much-accelerated slow process (k₂ = 0.75 ± 0.02 s⁻¹) as compared with C-AdcA (Fig. 4A). This indicated that, in the presence of a low concentration of zinc, the N-terminal domain rapidly bound a zinc atom and stabilized the entire protein structure. The C-terminal domain was then stabilized to obtain a faster binding speed. Both the zinc binding velocity and affinity constant of the C-terminal domain are 1 order of magnitude slower and weaker, respectively, than that of the N-terminal domain.

To validate the structural stability of the zinc-bound N-terminal domain, we performed proteinase K digestion assays. Zinc-saturated AdcA was almost intact after 30 min of proteinase K digestion, whereas the apo form of AdcA was mostly digested (Fig. 4B). In contrast, C-AdcA was vulnerable to proteinase K attack in both the apo and zinc-bound forms (Fig. 4C) due to the lack of stabilization by the zinc-bound N-terminal domain. This was echoed by the melting temperature of these two proteins. The melting temperature of Zn-C-AdcA was similar to that of WT AdcA and much lower than that of Zn₂-AdcA (Fig. 4D). These results suggested that the zinc in the N-terminal domain was crucial for enhancing the zinc binding rate in the C-terminal domain. As the C-terminal domain binds zinc with less affinity, it can pass zinc downstream to other zinc-binding proteins that finally transport zinc into the cell.

Structural alteration upon zinc binding

To investigate in detail the structural alterations after zinc binding, we compared the MD simulation trajectories of apo-AdcA, Zn-N-AdcA, and Zn₂-AdcA. The mean distance matrix showed remarkably decreased interdomain distances when the N-terminal domain bound a zinc atom (29) (Fig. 5A), which can be visualized in the steady-state three-dimensional conformations (Fig. 5B and Fig. 5C). This can be explained by N-terminal domain stabilization of the C-terminal domain via interatomic interactions of the peptide chains. The mean distance matrix of Zn-N-AdcA showed adjacency of the residue pairs Asn-237 and Lys-460, Glu-235 and Lys-457, Ser-258 and Asp-453, Lys-280 and Asp-453, Lys-280 and Tyr-430, and Asp-273 and Lys-421. All these interactions are interdomain interactions (30, 31) spanning the entire interaction surface between the two domains (Fig. 5C).

Moreover, the interdomain surface is linked by at least 10 hydrogen bonds (Fig. 5C, red dashed lines). This multianchored stabilization mechanism explains the stability of the induced conformational change. The induced conformation dramatically decreased the distance among the residues around the zinc-binding pocket of the C-terminal domain (32), specifically Lys-300 and Glu-481, Glu-299 and Tyr-360, and Lys-354 and Lys-409 (Fig. 5, D and E), facilitating the binding to zinc. This explained the increased affinity of the C-terminal domain when the N-terminal domain was zinc-bound. In summary, these analyses demonstrated the structural basis of the synergy between the two domains for both metal-binding affinity and rate.
We showed that the interface interactions rely on 10 hydrogen bonds between the N-terminal and C-terminal domains, involving the 12 amino acid residues depicted in Fig. 5. To evaluate the contributions of the 12 residues to the interface interaction, we successively established 12 single-site mutants of Zn-N-AdcA. However, we found that none of the single-site mutants caused any reduction of the surface formation (Fig. S7). Subsequently, we established four Zn-N-AdcA mutants, mutating groups of the abovementioned residues according to their vicinity: D273A, T276A, and K421A; K280A, S258A, Y430A, D453A, and G452A; E235A, N237A, K460A, and K457A; and a variant mutated for all 12 residues. In the fully mutated protein, only one zinc ion was located in the N-terminal domain of AdcA.

The trajectory analyses of the 100-ns MD simulations showed that these mutant proteins took a much longer time (~20 ns; Fig. 6A) to reach the equilibrium state than did the WT Zn-N-AdcA (~5 ns; Fig. 2B). Especially the fully mutated pro-
Two AdcA zinc-binding domains facilitate zinc transportation

The result revealed that the 12 residues of the interaction surface are crucial to stabilizing the skeleton of the whole protein. Among the 12 residues, the six residues at positions 273, 276, 280, 430, 452, and 453 showed increased RMSF values after mutation to alanine (Fig. 6B).

We next assessed whether the hydrogen bonds stabilized the interaction surface, measuring the solvent-accessible surface area (SASA) where a smaller SASA value represents tighter structures. The four corresponding average separation distances for each of the above mentioned mutants were 15.8, 28.8, and 10.6 Å (Fig. 6C, D273A, T276A, and K421A); 12.5, 20.4, and 22.1 Å (Fig. 6D, K280A, S258A, Y430A, D453A, and G452A); 8.7, 20.9, and 10.9 Å (Fig. 6E, E235A, N237A, K460A, and K457A); 21.5 Å, 22.3 Å, and 24.6 Å (Fig. 6F, fully mutated variant). Significantly, conformational changes in the C-terminal domain induced by the N-terminal domain of these four mutant proteins were reduced to varying extents when compared with WT Zn-N-AdcA (11.0, 13.1, and 8.1 Å; Fig. 5E). Among these, the fully mutated variant had the greatest impact on the conformation of the C-terminal domain. Based on these results, we propose a possible mechanism: the N-terminal domain firmly and rapidly binds a zinc ion and draws the C-terminal domain closer, relying on hydrogen bonds. Subsequently, this stabilizes the C-terminal domain and tightens the zinc-binding pocket, which facilitates zinc binding, especially in zinc-depleted environments.

A synergistic double-domain AdcA endows growth advantages at low zinc concentrations

The double-domain AdcA increased both affinity and rate of zinc binding. Therefore, we postulate that bacteria use the synergistic organization of the double-domain AdcA to more efficiently take up zinc, even under conditions of extremely low environmental zinc concentrations, to maintain growth. Due to the lack of genetic manipulation tools in S. pyogenes, we performed experiments in S. pneumoniae. We used N,N,N”,N”-tetraakis(2-pyridylmethyl)ethylenediamine (TPEN) to chelate zinc ions in the media and to create a zinc-deficient environment. Based on these results, we propose a possible mechanism: the N-terminal domain firmly and rapidly binds a zinc ion and draws the C-terminal domain closer, relying on hydrogen bonds. Subsequently, this stabilizes the C-terminal domain and tightens the zinc-binding pocket, which facilitates zinc binding, especially in zinc-depleted environments.

Discussion

Generally, bacterial genomes tend to encode smaller proteins (267 amino acids long on average) than do eukaryotes (33). Major reasons include that mRNA is degraded by endonucleases in bacteria, making it difficult to translate full-length long proteins (19), and multidomain large proteins need proper translational pausing sites for correct folding, which may be disturbed by environmental and molecular factors (34–36). Indeed, in most bacterial species, the homologs of Streptococcus AdcA are single-domain small proteins. Therefore, the evolution of the double-domain AdcA in Streptococcus (515 amino acids) should be unlikely unless an emergent feature benefits the adaptive survival of Streptococcus in host environments, such as lung tissue (1–2 μM zinc) (7). A previous study suggested that N-terminal domains in two-domain proteins are biased to be shorter and are predicted to fold faster than their C-terminal counterparts (37). However, AdcA is a special case of an overall trend in two-domain proteins as its N-terminal domain is longer than its C-terminal domain.

In this study, we found that double-domain AdcA with two zinc-binding sites is significantly overrepresented in Streptococcus species by homology comparison between AdcA and the library of known genomes of bacteria. We have shown evidence that fusion of the two domains creates a new, emergent, structure-based functional advantage that is greater than the sum of the advantages provided by two independent domains. In the presence of zinc, the N-terminal domain firmly and quickly binds a zinc ion and changes its conformation. This conformation change in the N-terminal domain subsequently stabilizes the C-terminal domain and tightens its zinc-binding pocket, facilitating zinc binding, especially in zinc-depleted environments. The relatively less stable conformation and weak binding of the C-terminal domain ensure a rapid transfer of zinc to the downstream transporter, the transmembrane protein AdcB of Streptococcus species (Fig. 8). Otherwise, overly stable and strong binding would limit the cyclic utilization of zinc transporters on the cell surface and the flux of zinc into the cell as shown in Fig. 7B. This interdomain synergy via interatomic interaction of the peptide chains endows Streptococcus with exceptional zinc uptake efficiency in zinc-depleted media, thus benefiting its survival in harsh conditions, e.g. in lung alveolar lavages with low zinc concentration (7). Therefore, the emergent features of the fused domains could be a simple evolutionary response to specific metal-deficient conditions (Fig. 8).

As organisms need multiple trace elements for physiology, other metal ion transporters are also found to exist in double-domain formats with two binding sites for the same ion. Examples are not restricted to bacteria and include the human transferrin receptor that binds two samarium ions (Protein Data Bank code 1CX8), human copper-lactoferrin that binds two copper ions (Protein Data Bank code 1LFI), R. thomisiana hemocyanin that binds two copper ions (Protein Data Bank code 1LN.L.), lactoferrin that binds two Fe2+ ions (Protein Data Bank code 2G2N).
Figure 6. Identification of the N- and C-terminal interaction surfaces of Fig. 5C. A, the time evolution of RMSD values in the four mutant Zn-N-AdcAs, D273A/T276A/K421A (blue), S258A/K280A/Y430A/G452A/D453A (gray), E235A/N237A/K457A/K460A (red), and the fully mutated variant (black). B, the RMSF of the C-α for all residues of the four mutant proteins calculated over the 100-ns trajectory. The RMSF values of the mutated amino acids became greater. Hydrogen bonds (red dashes), missing hydrogen bonds (black dashes with “/”), interaction surfaces (dotted line frame), and conformation changes of the C-terminal domain are shown in the four mutated forms of Zn-N-AdcA, D273A/T276A/K421A (C), S258A/K280A/Y430A/G452A/D453A (D), E235A/N237A/K457A/K460A (E), and the fully mutated variant (F). G, the calculated SASA values of the interface area, respectively, and error bars represent S.D.
Bank code 1B7Z), and others. Our model for emergent interdomain synergy may provide new insights to better understand such cases.

Our finding also emphasizes the interdomain conformation change after metal binding, which may implicate a novel target for antimicrobial drugs against pathogenic Streptococcus species. No AdcA homolog was found in the human genome using the HMMER tool (Fig. S8), suggesting that such a conformation may not exist in the human proteome. Therefore, a rationally designed blocker molecule that binds the interdomain surfaces may abolish the synergy of the two domains, sufficiently reduce the influx of zinc, and inhibit the growth of the pathogen. The action mechanism of the designed blocker molecule would be totally different from that of existing antibiotics. To be noted, the interdomain interaction is stabilized by as many as 10 hydrogen bonds (38, 39), suggesting that mutation of a few amino acids will not abolish the interaction, thus minimizing the probability of the bacteria to evolve resistance by simple point mutations. This finding may help during the treatment of bacterial infections caused by Streptococcus.

In conclusion, we suggest a special zinc transportation mechanism mediated by AdcA in Streptococcus species. An N-terminal binding site preferentially binds a zinc ion and induces stabilization of the overall conformation of the protein via interdomain interaction. This allows the C-terminal binding domain to acquire a zinc ion while also allowing zinc bound by the C-terminal domain to be more likely and preferentially released (Fig. 8). This model elucidates the significance of the evolution of the domain fusion, provides new insights on double-domain transporter proteins with two binding sites for the same ion, and implicates a novel target for antimicrobial drugs against pathogenic Streptococcus species.

**Experimental procedures**

**MD simulation**

The preprocessed structures of apo-AdcA, Zn-N-AdcA (mutation of the three zinc-binding ligands in the C-terminal domain), Zn-C-AdcA (mutation of the four binding ligands in the N-terminal domain), and Zn2-AdcA, obtained from homology modeling, were used as starting conformations. These conformations were solvated in cubic periodic boxes containing 0.15 M Na\(^+\) and Cl\(^-\) ions to neutralize the system (40, 41). No zinc ions were added to the simulated solution, representing a zinc-deficient environment. Energy minimization in each AdcA system was performed for the first relaxed energy through 400 steps of the steepest descent energy method and then continued with 25,000 steps of the conjugate gradient method. All MD simulations were simulated at a temperature of 310.15 K (42) and pressure of 1 atm (43) by the Gromacs 4.6.6 package with the simple point charge (SPC) model for liquid water as described previously (44). The zinc ion force field of the Gromacs package has been developed in various aspects (45), including bonds, angles, impropers, metal center, dihedrals, and normal van der Waals, and it is suitable for the protein-zinc simulation (46–48). Therefore, we used the default
parameters of Gromos43a1 force field for the simulation of AdcAs with zinc. We set the value of the entol convergence criterion at 1,000 kJ/mol/nm, and the temperature in the box was determined by the v-rescale temperature coupling method. We used the particle mesh Ewald method to calculate electrostatic interactions within the system. The position restraint simulations for the systems were executed for 50 ps, and then we actualized the 40-ns unrestrained simulation.

We simulated four structures of the AdcA system as mentioned above, and each system was repetitively carried out three times with random initial velocities. To avoid equilibration artifacts, we calculated the structural features using trajectory data ranging from 10 to 40 ns, which represents the structures at their equilibrated states. To validate the equilibrium of each structure, we also performed 300-ns MD simulations for apo-AdcA and Zn$_2$-AdcA. We analyzed RMSD, RMSF, secondary structure, and $R_g$ using Gromacs tools g_rms, g_rmsf, do_dssp, and g_gyrate, respectively. The mean distances between residues were calculated by the Gromacs tool g_dist. Protein structures were visualized using PyMOL 1.7. The matrix of residue distance fluctuations was represented by methods described previously, and it can be used to describe the plasticity...
and elasticity of residues in structural fluctuations. To reduce the deviation, all MD trajectories data were derived from three replicates (Fig. S2).

**Homology modeling**

The NCBI BLAST search tool was used to find several proteins that have the highest homology to AdcA. Then a multiple sequence alignment and cluster analysis were performed using proteins with high scores through the software ClustalX 2.0. The tertiary structure of AdcA was modeled using MODELLER in Accelrys Discovery Studio Client 4.1 (52, 53). Proteins San-YodA (Protein Data Bank code 1TXL) and BsuYcdH (Protein Data Bank code 2O1E), with a high level of amino acid sequence homology to AdcA, were selected as the templates to model the initial stage of the three-dimensional structure of AdcA. Subsequently, flexible molecule docking between AdcA and zinc was processed via the LigandFit module of the software to find the lowest energy conformation combining the ligand and the receptor in the active site (Fig. S1B). The reliability of the model was evaluated by discrete optimized potential energy and Ramachandran plots (54).

**Construction, expression, and purification of WT AdcA**

Genomic DNA extracted from *S. pyogenes* MGAS5005 (ATCC BAA-947TM) was used as a template to amplify the adcA gene (1,488 bp without the N-terminal signal). PCR primers were designed to introduce the restriction enzyme sites of BamHI and Sall for the adcA gene (Table S1). Detailed methods were described previously (55). Purified AdcA protein was confirmed using 12% SDS-PAGE and identified using MS (ABI 4800 MALDI-TOF/TOF) according to a method described previously (56) (Fig. S3D).

**Protease K resistance experiments**

To test the protease K (Roche Diagnostics) sensitivity of AdcA, the same amounts of apo-, zinc-saturated WT, and mutant AdcA (15 μg) were incubated with 30 μg/μl protease K in 20 mM Tris-HCl buffer (pH 8.0) containing CaCl₂ (10 mM) at 25 °C for 0, 1, 3, 5, 7, 10, and 30 min. Proteolysis was stopped by addition of 0.1% phenylmethylsulfonyl fluoride and boiling for 10 min. Digestion fragments were analyzed by 12% SDS-PAGE. Gels were stained with Coomassie Blue R-250 and scanned with Image Scanner II (GE Healthcare).

**Circular dichroism spectroscopy**

Far-UV CD studies were performed with a CD spectrometer (Chirascan, Applied Photophysics Ltd., Leatherhead, UK) using a quartz cuvette with a 0.1-cm optical path length at a wavelength range of 260–190 nm at room temperature. CD data were collected for 5 μM apo-, zinc-saturated WT, and mutant AdcA in 20 mM Tris-HCl (pH 7.4) in a data pitch of 1 nm at a scanning rate of 100 nm/min. Each CD spectrum was repeated three times, and a blank containing the same buffer was subtracted as a reference. Analysis of the experimental data was performed using the software CDPro. CD spectra were also used to study the thermal stability of proteins in the absence or presence of zinc ions. Thermal unfolding curves were monitored from 20 to 90 °C using an increase rate of 2 °C/min by detecting the loss of secondary structures at 222 nm. Each data set was obtained three times using steps of 0.5 °C, and Tm values were calculated by the included Glob3 software.

**Growth media and growth curve assays**

Casein-based semisynthetic liquid culture medium (C+Y medium) was used to culture WT and mutant D39 strains (57). To establish Zn(II) starvation conditions, TPEN (Sigma-Aldrich) was added into C+Y medium at final concentrations of 10, 20, 30, and 35 μM (8, 58, 59). We determined the growth curves for WT and ΔadcA ΔadcAll double-mutant strains cultured in zinc starvation medium at 37 °C with 5% CO₂ for 12 h by measuring A600 values at different time points. For the double-mutant strain, 20 μM TPEN significantly inhibited bacterial growth at an A600 of ~0.15 after 12 h of culture. We selected 20 μM TPEN to add to the medium to create zinc deficiency. Different treated *S. pneumoniae* strains were inoculated into C+Y medium, and growth curves were determined three times.

**Construction, expression, and purification of mutant AdcAs**

Based on the predicted structure of AdcA with molecular docking, the N-terminal binding site is composed of His-36, His-122, His-186, and Glu-261, and the C-terminal binding site is composed of His-436, His-445, and His-447. The four amino acids in the N-terminal domain and the three amino acids in the C-terminal domain were simultaneously mutated to alanine to generate the four-residue mutant H36A/H122A/H186A/E261A (C-AdcA) and the three-residue mutant H436A/H445A/H447A (N-AdcA), respectively, using a QuickChange mutagenesis kit (Stratagene) with the original pGEX-4T-AdcA plasmid as template. The primers used for constructing the mutants are listed in Table S1. All the constructed plasmids were transformed into *E. coli* BL21 (DE3) for expression. Expression and purification of mutant AdcAs were conducted as done for WT AdcA.

**Construction of the ΔadcA/ΔadcAll double-mutant strain**

The primer sequences used to construct mutant strains are listed in Table S1. The double mutant strain was constructed as described previously (60, 61). Long flanking homology PCR products contained an antibiotic resistance cassette (erythromycin or spectinomycin) flanked by 600-bp-long fragments homologous to the end of each target gene, adcA or adcAll. Then the long flanking homology PCR fragments were transformed into *S. pneumoniae* D39 competent cells. Transformants were selected with antibiotic-containing Columbia sheep blood agar plates after overnight culture at 37 °C with 5% CO₂ and confirmed by DNA sequence analysis and PCR (Fig. S3C). The mutant strain was stable after six sequential passages in Todd-Hewitt broth with 0.5% yeast extract (THY) medium in the absence of antibiotics.

**Construction of the three types of overexpression strains**

To construct overexpression strains of *S. pneumoniae* D39 for recovery of expression of different types of AdcA, the plas-
mid pLB169 (p169) was used in this study (62). We constructed three recombinant plasmids, p169-N-lobe (expressing the N-lobe alone), p169-C-lobe (expressing the C-lobe alone), and p169-adcA (expressing full-length adcA) (Fig. S3C). The constructed plasmids were transformed into the ΔadcA ΔadcAll double mutant strain, and the positive clones were screened using Columbia blood plates with 4 μg/ml chloramphenicol. The transformants with the recovered genes were verified by PCR. All primers are listed in Table S1.

Comparison of the zinc binding strength of the two domains of AdcA

To compare the binding strength of the N-terminal and C-terminal domains of AdcA, a special zinc metallochormic indicator, PAR, was used. A final concentration of 100 μM PAR was added to 5 μM Zn-N-AdcA, Zn-C-AdcA, and Zn2-AdcA in 20 mM Tris-HCl (pH 7.4) with or without 6 mM guanidine hydrochloride. UV/visible absorbance spectra were obtained from 300 to 600 nm after a 5-min equilibration at room temperature.

Zinc-binding affinity determination

To determine the binding affinities of WT and mutant AdcAs with zinc, ITC experiments were performed at 25 °C using a MicroCalorimeter Auto-ITC 200 (Malvern, UK). Prior to the experiments, the instrument was washed with deionized water, and the ZnCl2 and AdcAs were dissolved in the same solution buffer (20 mM Tris-HCl, 100 mM NaCl (pH 7.4)) (55). Typically, an experiment consisted of loading the syringe with zinc ions at a concentration at least 10-fold higher than the AdcAs samples, which were placed in the cell. The titration parameters were set as follows: 2 μl of ZnCl2 solution were injected into the 300-μl protein sample cell during each titration with 15–20 injections. The delay time between injections was set at 200 s to ensure thermal equilibrium before the next injection. The background heat effect was subtracted by addition of zinc alone to the buffer. All integrated heat data were analyzed using Origin 7.0 software for fitting calculations.

Stopped-flow absorbance kinetics

Stopped-flow spectroscopy was performed on a stopped-flow reaction analyzer (Chirascan SF.3, Applied Photophysics Ltd.) using the absorbance mode to monitor absorbance changes at 495 nm over time. The Zn(PAR)2 complex (200 μM PAR in 20 mM Tris-HCl buffer combined with 40 μM Zn2+) was loaded into the A drive syringe, the B drive syringe was filled with 10 μM apo-AdcAs, and transient mixing of the reaction was driven by bottled nitrogen. Experimental parameters were set as follows: 1-nm bandwidth, 10-mm optical path, 495 nm scanned wavelength with a 475-nm filter, and a 25 °C water bath temperature. Buffer incubated with Zn(PAR)2 was used as the reference. Tests were repeated until consistent results were obtained. The collected data were analyzed using exponential equations.

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