

Resolution of Stepwise Cooperativities of Copper Binding by the Homotetrameric Copper-Sensitive Operon Repressor (CsoR): Impact on Structure and Stability

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Abstract: The cooperativity of ligand binding is central to biological regulation and new approaches are needed to quantify these allosteric relationships. Herein, we exploit a suite of mass spectrometry (MS) experiments to provide novel insights into homotropic Cu-binding cooperativity, gas-phase stabilities and conformational ensembles of the D_2 -symmetric, homotetrameric copper-sensitive operon repressor (CsoR) as a function of Cu^I ligation state. Cu^I binding is overall positively cooperative, but is characterized by distinct ligation state-specific cooperativities. Structural transitions occur upon binding the first and fourth Cu^I , with the latter occurring with significantly higher cooperativity than previous steps; this results in the formation of a holo-tetramer that is markedly more resistant than apo-, and partially ligated CsoR tetramers toward surface-induced dissociation (SID).

The structural, dynamic and thermodynamic origins of cooperativity of ligand binding (allostery) is a subject of considerable interest, motivated by the central role this process plays in the regulation of biological activity.^[1] Cooperativity can be positive or negative and involve the binding of the same (homotropic) or different (heterotropic) ligands, often to an homooligomeric protein.^[2,3] Bacterial repressors that function to control transition metal bioavailability in cells are typically homodimeric or homotetrameric and minimally bind two “ligands”: a DNA operator found upstream of metal-regulated genes and a specific (cognate) transition metal ion(s).^[4] Although a robust thermodynamic framework capable of quantifying both homo- and heterotropic cooperativity in these systems is available,^[5] these ensemble-based methods suffer from the limitation that a specific, partially ligated state can not be studied independently of other states. Such step-wise insights are required to understand allosteric coupling beyond a generally phenomenological description.^[1]

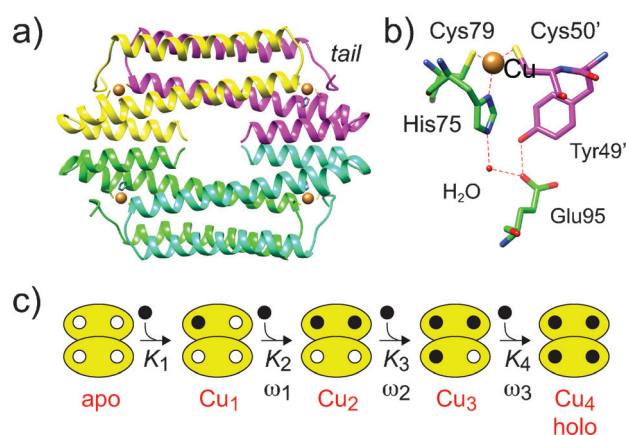


Figure 1. Ribbon representation a) of the structure of a thermophilic CsoR tetramer (PDB: 4M1P) closely related to *B. subtilis* CsoR studied here.^[16] Each protomer is shaded differently, with the Cu^I ions indicated by the brown spheres. The folded N-terminal tail (*tail*) is indicated. b) Close-up of the Cu^I binding pocket of CsoR. c) Schematic representation of the step-wise K_i and ω_i used in this analysis.

The Cu^I -specific homotetrameric metalloregulatory protein, copper-sensitive operon repressor (CsoR) (Figure 1a) binds four Cu^I ions to four identical subunit-bridging Cys₂-His sites (Figure 1b) with high affinity, $K_{\text{Cu}} \approx 10^{18} \text{ M}^{-1}$.^[6,7] Apo-CsoR forms a 2:1 CsoR tetramer:DNA “sandwich” complex, with bound DNA binding along one face of each of two tetramers.^[8–10] Cu^I binding leads to dissociation of CsoR from the DNA operator (negative heterotropic linkage) resulting in transcriptional derepression of Cu-resistance genes. CsoR regulates free Cu, which is buffered to vanishingly low bioavailability in the cytoplasm of cells.^[11] Copper toxicity^[12] is a well-established antimicrobial weapon employed by the host to combat bacterial infection.^[13,14] Previous studies reveal that Cu-mediated DNA dissociation is associated with a global quaternary structural compaction in the CsoR tetramer.^[15,16] However, there is no information on the structure or stability of partially Cu-ligated states of CsoR, nor is it known if the binding of Cu is cooperative.

Herein, we develop a soft-ionization-based mass spectrometry (MS) approach to unravel homotropic linkage relationships in CsoR by resolving partially Cu-ligated tetramers based on their unique m/z values. Although previous studies have determined the Hill coefficient for cooperativity of ligand binding to a homotetramer using mass spectrometry^[17] and prior work on monomeric metal binding

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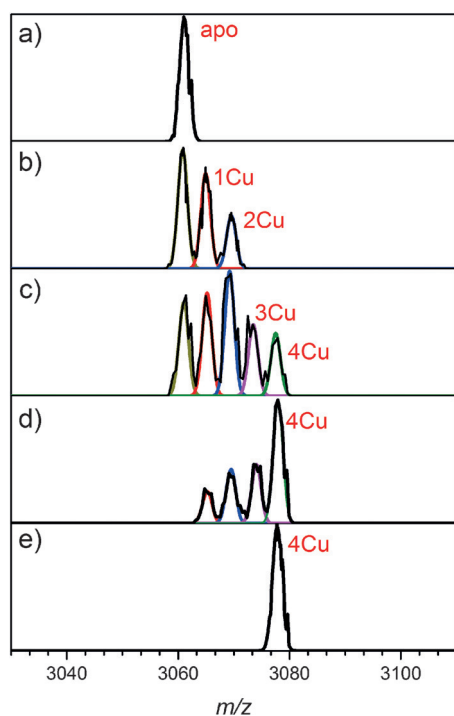


Figure 2. Representative normalized mass-to-charge spectra of the 15+ charge state of CsoR as a function of copper addition. Solutions were 500 μM in CsoR protomer (125 μM tetramer) with the addition of a) 0 μM CuCl, b) 125 μM CuCl, c) 250 μM CuCl, d) 375 μM CuCl, or e) 500 μM CuCl. Each spectrum was fit with Gaussians at the mass-to-charge ratio of each Cu-bound state to determine the intensity (I) of each metallated species (apo, yellow; Cu₁, red; Cu₂, blue; Cu₃, magenta; Cu₄, green; left to right). Mol species fractions are calculated from $I_i/\sum I_i$ for each addition of Cu (see Figure 3).

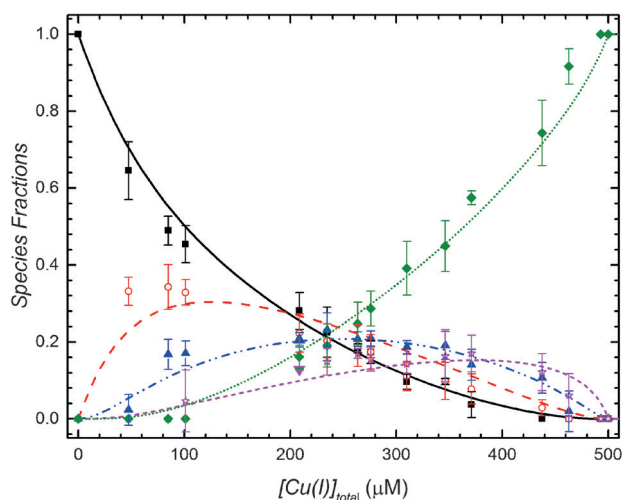


Figure 3. Mol species fractions of CsoR tetramers as a function of Cu^I concentration. Data represent triplicate experiments across all useable charge states, with standard error of the mean value defined by the error bar. The continuous lines are the results of a global fit of all species fractions as a function of added Cu^I to a step-wise cooperative binding model (see Supporting Information). Apo, Cu₁, Cu₂, Cu₃, Cu₄ CsoRs species are represented by black squares, red open circles, blue triangles, magenta open stars and green diamond symbols, respectively.

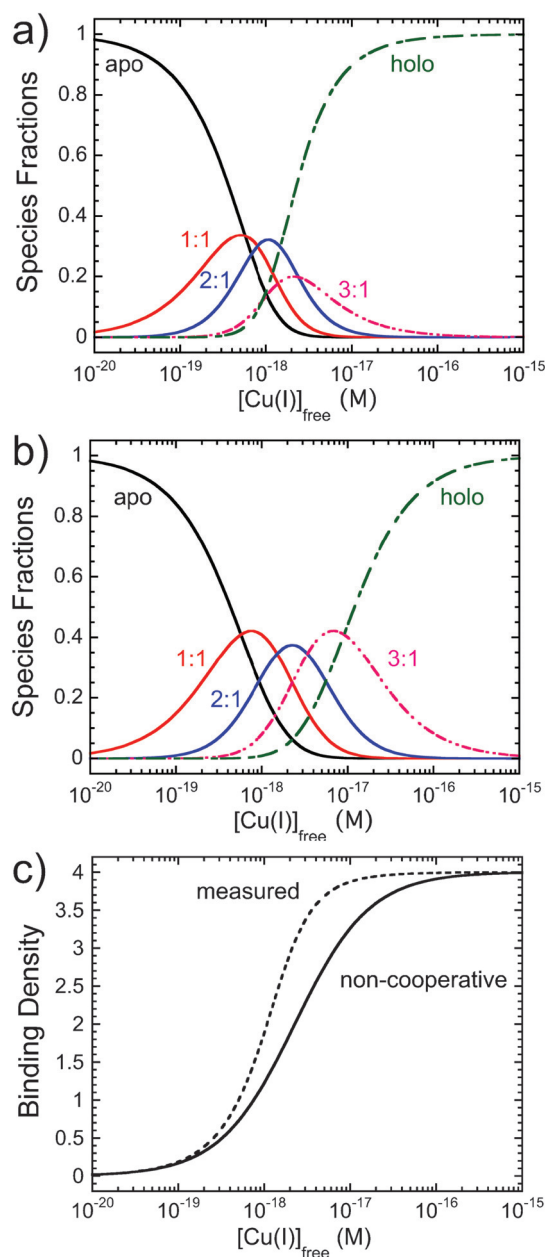


Figure 4. Simulated species fractions versus $[\text{Cu}]_{\text{free}}$ curves defined by the parameters resolved a) from the step-wise cooperative binding model and b) a non-cooperative binding model in which $k = 0.44 \times 10^{18} \text{ M}^{-1}$ (the fitted value from K_i (initial) of $1 \times 10^{18} \text{ M}^{-1}$; see Table S1). c) Cu-binding isotherms for the step-wise cooperative (dashed line, measured) and non-cooperative (solid line) binding models.

proteins shows that apo- and holoproteins obtained at substoichiometric metal can be monitored by mass spectrometry as a titration is carried out in solution,^[18–21] we develop here a generally applicable approach to rigorously determine the cooperativity of Cu^I binding at every Cu^I binding step, termed the step-wise cooperativities (Figure 1c).

To do this, we quantified the fractional concentrations of each i th ligated species of the CsoR tetramer (where $i = 0$ to 4) as a function of added CuCl up to stoichiometric equivalence in an anaerobic solution under conditions in

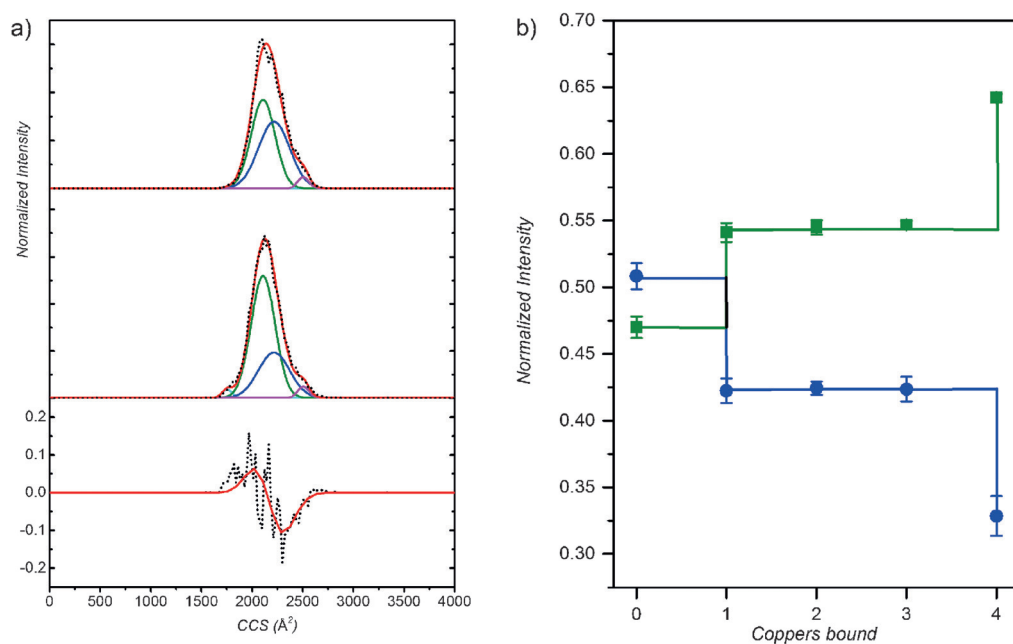


Figure 5. a) Cross-section mobility distributions for apo (top) and holo (middle) CsoRs, with the difference plots (holo-apo) of the raw data (black) and Gaussian-fitted model (red) shown (bottom panel). Major conformations are colored as green (2107 Å²) and blue (2216 Å²), while minor conformers are colored cyan (1755 Å²) and magenta (2505 Å²). Gaussian global fits superimposed on the data (black) are indicated by the red line. b) The normalized integrated intensities for the two major conformers as a function of Cu-ligation state.

which all added Cu^I is bound, i.e., [CsoR] $\gg$ 1/ K_{Cu} .^[22] The resulting mass spectra (see Figure S1 in the Supporting Information) of the 15+ charge state (Figure 2) show the change in abundance of each i th Cu^I-bound state ($i = 0-4$ in Figure 2a-e, respectively) as the total Cu^I concentration is increased. Even at a molar ratio of Cu^I to protomer of 2 (Figure 2c), the holo (Cu₄) species, characterized by an m/z of 3077, is present to a degree greater than would be expected for non-cooperative Cu^I binding, with further additions of Cu^I further increasing the abundance of the holo species (Figure 2d,e). This result holds for all charge states and can only be due to overall positively cooperative binding of Cu^I to the CsoR tetramer. The mol species fractions of each Cu _{i} state reveals a clear propensity to form holo-CsoR at substoichiometric concentrations of Cu (Figure 3).

A global fitting of the data in Figure 3 allows resolution of the step-wise equilibrium binding constants (K_1-K_4) and ligation state-specific Cu^I binding cooperativities, ω_1 , ω_2 , and ω_3 (see Supporting Information for details) (Figure 1c). The resolved step-wise cooperativities of $1.9(\pm 0.2)$, $2.3(\pm 0.2)$ and $5.2(\pm 0.3)$ for ω_1 , ω_2 and ω_3 along with K_4 ca. 2-fold larger than K_1 , reveal quantitatively that Cu^I binding to the CsoR tetramer is overall positively cooperative (Table S1; Figure S2). This cooperativity is also reflected in the species fractions versus [Cu]_{free} plots simulated from these K_i (Figure 4a), compared to a non-cooperative binding system with K_1 fixed at the value obtained for K_1 in the cooperative system (Figure 4b). This has the clear effect of moving the simulated Cu^I binding isotherm to the left of the non-cooperative binding isotherm, while making it appear more sigmoidal (Figure 4c); as a result, formation of holo-CsoR

occurs at ca. 5-fold lower [Cu]_{free} relative to the non-cooperative case.

The advantage of this approach for quantifying ligand binding cooperativities, subject of course to the proposal that gas-phase species fractions mirror those found in solution, is further illustrated by comparing simulated Cu^I binding curves obtained in a standard chelator competition assay using bathocuproine disulfonate (BCS) ($\log \beta_2^{Cu} = 19.8$ ^[23]) and described by the resolved cooperative vs. noncooperative binding parameters derived above (Figure 3). When these simulated curves are superimposed on published data for *B. subtilis* CsoR^[22] (Figure S3), binding parameters

derived from the noncooperative binding model ($K \geq 10^{19} M^{-1}$) can not be easily distinguished from those derived from the cooperative binding model developed above at all $K = K_1 \geq 10^{17} M^{-1}$ (Figure S3). Clearly, resolution of CsoR species fractions using the MS-based method outlined here is far superior to existing Cu^I-binding chelator competition methods,^[24] particularly in cases where distinct step-wise cooperativities (ω_i) are superimposed on large intrinsic ligand affinities (K_i) (Figure 1c).

We next investigated how these distinct cooperativities influence the structure and stability of the tetramer. Ion mobility spectrometry (IMS) allows for the assessment of gas phase conformational ensemble, which in many cases appears to be representative of the solution structural ensemble.^[16,25-31] Examination of the mobility distributions for the apo (Figure 5a, top) and holo (Figure 5a, middle) CsoRs show two similarly broad distributions, with clearly distinguishable peak centers of 2253 Å² for apo-tetramer and 2231 Å² for holo-tetramer (Figure 5a, bottom), given a peak center RMSD of 15.2 Å² (Figure S4), a finding inherent in the raw data and independent of the number of Gaussians used to fit these mobility distributions (Figure S5a). Although this trend in collisional cross-section is consistent with previous data revealing a hydrodynamically more compact holo species,^[16] the breadth of the IMS peaks suggests that multiple conformations are present. We therefore modeled these distributions for all five states (apo, Cu₁-Cu₄) with a set of four Gaussian functions (two major; two minor) (Figure 5a), since fitting to fewer Gaussians gives rise to statistically inferior fits (Figure S5b) (see Supporting Information). Although the two minor distributions ($\leq 3\%$ of the total)

do not change as a function of ligation state (Figure 5a), the relative abundance of the two major conformations at 2107 Å² and 2216 Å² change dramatically as a function of Cu ligation state (Figure 5b). These results show that the conformational ensemble becomes hydrodynamically smaller upon binding a single Cu^I, and does not vary greatly with the binding of the second and third Cu^I ions to the tetramer. However, binding of the fourth Cu^I induces a second shift to an even more structurally compact ensemble (Figure 5b). Previous studies on ubiquitin using both overtone mobility spectroscopy and IMS-IMS show that normally unresolved structures could be isolated and observed using this method.^[32,33] This is consistent with the idea that each ligation state of CsoR may represent an interconverting ensemble of apo- and holo-like “end” state structures, leading to broader-than-expected distributions. The origin of the Cu-dependent compaction is unknown, but previous solution studies of a related thermophilic CsoR suggests that this derives from kinking of the α2 helix, repacking of the tetramer interface, and folding of the N-terminal tail over the bound Cu (cf. Figure 1a).^[16] Remarkably, these changes in cross-section precisely mirror with the relative degrees of cooperativity upon filling the tetramer: the site-cooperativity is nearly independent of ligation state in transiting from the $i = 1$ to $i = 2$ and $i = 2$ to $i = 3$ states, but increases significantly in going from the $i = 3$ to $i = 4$ holo states.

Finally, we exploited our ability to resolve differentially i th ligated states of CsoR and determined their stabilities in the gas phase, as measured by the fragmentation of each tetramer through surface induced dissociation (SID) at 40 V acceleration voltage (Figure S6, 13+ charge state). As more Cu^I is bound, the precursor increases in relative abundance in the fragmentation spectrum, with the converse is true in the monomer fragmentation region. A scan from 10 to 100 V acceleration voltage reveals clearly that the fragmentation pathway differs markedly as a function of Cu^I loading (Figure 6). Apo-CsoR and all other substoichiometric Cu^I-CsoR tetramer complexes (Figure 6a–d) are completely fragmented by 70 V acceleration voltage, cleaving into predominately monomer as the acceleration voltage increases. In contrast, holo-tetramer begins to fragment at higher voltages but most striking is the differential dimer formation during fragmentation of the apo- versus Cu-loaded tetramer (Figure 6e). In fact, dimer fragments *increase* in intensity for the holo-tetramer with increased acceleration voltage, accounting for 27% of fragments at 100 V.

These data suggest that the dimer is an intermediate in the fragmentation of the Cu-loaded CsoR tetramer to monomer, a finding that likely derives from the full complement of Cu^I coordination bonds within each dimer of the Cu₄ tetramer (cf. Figure 1b), thus biasing the SID-induced fragmentation pathway by inducing dissociating at the dimer-dimer interface. Further, the persistence of the dimer with increasing acceleration voltages suggests that positively cooperative binding of Cu^I, most notable for the fourth Cu-binding step, strongly stabilizes both the $n = 2$ and holo forms of the tetramer. This finding is unanticipated from fragmentation patterns of D_2 -symmetric tetramers predicted by Proteins, Interfaces, Structures and Assemblies (PISA) analysis and

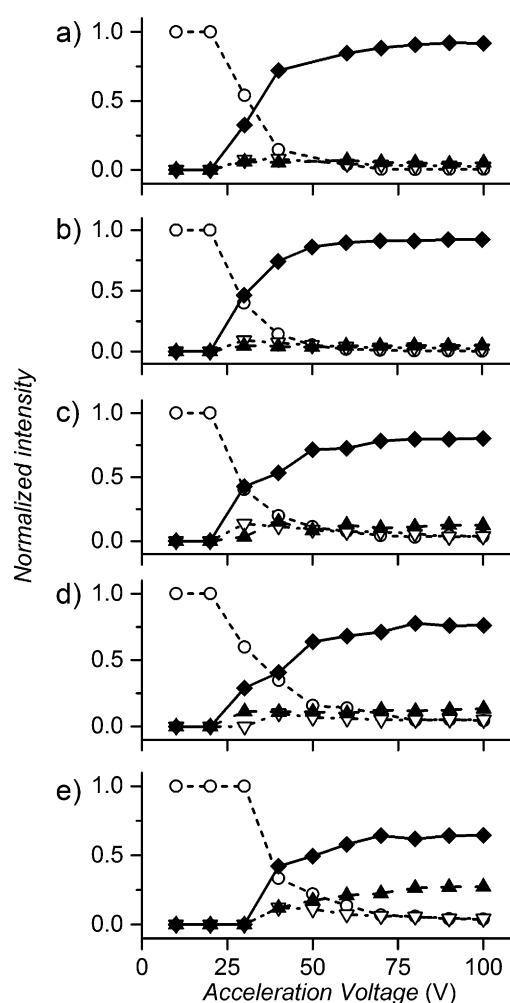


Figure 6. Fractional abundance of precursor and fragment peaks produced by SID as a function of acceleration voltage for a) apo, b) Cu₁, c) Cu₂, d) Cu₃ and e) holo Cu₄ CsoR species. The normalized fractional intensities of each fragment species are shown at each acceleration voltage. Tetramer, open circle trace; trimer fragment, open upside down triangle trace; dimer fragment, triangle trace; monomer fragment, diamond trace.

SID,^[34] but is likely a consequence of very strong, subunit bridging, coordinate covalent Cu^I–S bonds.

In summary, we have developed a general MS-based framework to extract and quantify ligation state-resolved homotropic cooperativities in an oligomeric metal-binding protein, and use IMS and SID fragmentation to examine the step-wise impact of metal binding on the structure and stability of the partially ligated states. This method is dependent on transition metal (Cu, Zn, Cd) complexes that are not disrupted by the electrospray ionization source.^[21,35,36] We find that the binding Cu^I to CsoR is characterized by a distribution of ligation state-dependent cooperativities, with the final Cu-binding event biasing the conformational ensemble toward a more highly compact and more stable structure. This degree of cooperativity has the effect of moving the biological response to lower free Cu^I and making the response more “all-or none”, as is often desired for a molecular switch. Current efforts are directed toward extending this method to

other metallosensors and exploring the role of the DNA operator in these coupled equilibria.

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