Co(II)-detection does not follow $K_{\text{Co(II)}}$ gradient: channelling in Co(II)-sensing†

Carl J. Patterson, Rafael Pernil, Samantha J. Dainty, Buddhapriya Chakrabarti, Clare E. Henry, Victoria A. Money, Andrew W. Foster and Nigel J. Robinson*

The MerR-like transcriptional activator CoaR detects surplus Co(II) to regulate Co(II) efflux in a cyanobacterium. This organism also has cytosolic metal-sensors from three further families represented by Zn(II)-sensors ZiaR and Zur plus Ni(II)-sensor InrS. Here we discover by competition with Fura-2 that CoaR has $K_{\text{Co(II)}}$ weaker than $7 \times 10^{-8}$ M, which is weaker than ZiaR, Zur and InrS ($K_{\text{Co(II)}} = 6.94 \pm 1.3 \times 10^{-10}$ M; $4.56 \pm 0.16 \times 10^{-10}$ M; and $7.69 \pm 1.1 \times 10^{-9}$ M respectively). $K_{\text{Co(II)}}$ for CoaR is also weak in the CoaR-DNA adduct. Further, Co(s) promotes DNA-dissociation by ZiaR and DNA-association by Zur in vitro in a manner analogous to Zn(II), as monitored by fluorescence anisotropy. After 48 h exposure to maximum non-inhibitory [Co(s)], CoaR responds in vivo yet the two Zn(II)-sensors do not, despite their tighter $K_{\text{Co(II)}}$ and despite Co(s) triggering allostery in ZiaR and Zur in vitro. These data imply that the two Zn(II) sensors fail to respond because they fail to gain access to Co(s) under these conditions in vivo. Several lines of evidence suggest that CoaR is membrane associated via a domain with sequence similarity to precorrin isomerase, an enzyme of vitamin B$_{12}$ biosynthesis. Moreover, site directed mutagenesis reveals that transcriptional activation requires CoaR residues that are predicted to form hydrogen bonds to a tetrapyrrole. The Co(II)-requiring vitamin B$_{12}$ biosynthetic pathway is also membrane associated suggesting putative mechanisms by which Co(II)-containing tetrapyrroles and/or Co(II) ions are channelled to CoaR.

Introduction

Cells control the buffered concentration of each metal in the cytosol through the combined actions of proteins of metal homeostasis including metal-importers, -exporters, -storage proteins, -delivery proteins and -sensors. In bacterial cells the metal-sensors are commonly, although not exclusively, DNA-binding, metal-binding transcriptional regulators. These sensors are categorised into different families and function as metal-dependent activators, de-repressors or co-repressors. The ability of the sensors to discern and respond to the correct metal(s) as opposed to all other ions is paramount, since this influences the expression of the homeostatic proteins which in turn influences which metals are available to occupy other metallo-proteins. Factors which dictate metal selectivity in metal-sensors have been summarised under the headings affinity, allostery and access.

The cyanobacterium Synechocystis PCC 6803 contains a Co(u)-responsive transcriptional activator of the MerR-family, CoaR. In elevated concentrations of cobalt CoaR activates transcription of a gene encoding a cobalt-exporting P$_1$-type ATPase, CoaT, from a promoter with sub-optimal spacing between canonical $-10$ and $-35$ RNA polymerase binding sites. Variants in which this spacing is shortened confer constitutive transcriptional activity consistent with CoaR residues that are predicted to form hydrogen bonds to a tetrapyrrole. The Co(II)-requiring vitamin B$_{12}$ biosynthetic pathway is also membrane associated suggesting putative mechanisms by which Co(u)-containing tetrapyrroles and/or Co(u) ions are channelled to CoaR.

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co-repressor of the Fur family, Zur.8,14–19 We recently explored the hypothesis that selectivity in favour of Ni(II)-sensing by InrS is partly a function of the relative affinities of the complement of sensors in a cell.17 The Ni(II) sensor InrS was shown to have the tightest $K_{Ni(II)}$ of the set enabling InrS to de-repress Ni(II)-efflux at a cytosolic concentration below $K_{Ni(II)}$ of the other sensors, providing a mechanism by which the actions of InrS sustain a buffered [Ni(II)] sufficiently low to prevent the other sensors from gaining access to Ni(II).17 Here we unexpectedly discover that an analogous model cannot explain the Co(II) specificity of CoaR, prompting a search for additional mechanisms to account for this apparent anomaly.

**Results**

DNA and Co(II)-binding CoaR can be purified using n-dodecyl $\beta$-D-maltoside

Inability to purify sufficient CoaR for *in vitro* biochemical studies stalled analyses beyond an initial characterisation over a decade ago.12 Three independent algorithms now predict surface exposed, hydrophobic, potentially membrane associated regions within a domain of CoaR which can be modelled on the structure of precorrin isomerase from *Pseudomonas denitrificans* (Fig. S1, ESI†). Inclusion of a non-ionic detergent in buffers enabled purification of CoaR from recombinant *E. coli* cell extracts.17 Titration of a hexachlorofluorescein-labelled 36 bp coa operator-promoter fragment containing a 13-6-13 hyphe-nated inverted repeat corresponding to the CoaR binding site in *P. denitrificans* (coa-O/P) showed binding of the recombinant protein to its deduced DNA target with $K_{DNA} \sim 28$ nM (Fig. 1A). Moreover, addition of Co(II) to an anaerobic solution of CoaR generated Co(II)-dependent spectral features consistent with ligand to metal charge transfer (LMCT) due to formation of Co(II)-thiolate bonds ($\epsilon_{310 \text{ nm}} \approx 2,500 \text{ M}^{-1} \text{ cm}^{-1}$), which saturate at ~ one molar equivalent of Co(II) suggestive of two sites per CoaR dimer of $K_{Co(II)} \leq 10^{-6}$ M (Fig. 1B). Co(II)-dependent features around 650 nm indicate d-d transitions of sufficient intensity to report upon tetrahedral coordination environments, which moreover contain at least two Cys-thiols per site based upon the intensity of the LMCTs (Fig. 1B).20

$K_{Co(II)}$ is weaker for CoaR than for ZiaR, Zur or InrS

To explore why CoaR is the cellular sensor for Co(II), by analogy to recent investigations into the specificity of Ni(II)-detection in the same cells,15 $K_{Co(II)}$ was estimated for Zn$_2$Zur (one atom of Zn(II) is kinetically trapped in a structural site),21 ZiaR, InrS and CoaR; representing one member from each family of cytosolic metal-sensors encoded within the *Synechocystis* PCC 6803 genome.17 Co(II)-dependent spectral features were described previously for InrS, and are presented here for ZiaR and Zn$_2$Zur (Fig. S2, ESI†).17 Co(II)-titrations of ZiaR reveal LMCT and d-d-transitions requiring at least two molar equivalents of metal to reach saturation, consistent with four tetrahedral sites per dimer. Unlike the d-d feature at 585 nm, the LMCT feature at 310 nm does not show a linear [Co(II)]-dependent increase in intensity (Fig. S2A, ESI†). ZiaR contains two conserved metal binding sites; a thiol-free z5 site with predicted ligands (two His, Asp, Glu) derived from antiparallel z5 helices and an z3N site with predicted ligands (three Cys, His) derived from z3 helices and an amino terminal extension.7,15 These data (Fig. S2A, ESI†) imply that thiol-free z5 sites have the tightest $K_{Co(II)}$, with two to three Cys-thiols present in z3N sites of weaker $K_{Co(II)}$. Co(II)-titration of Zn$_2$Zur also reveals d-d transitions at 585 nm of intensity consistent with tetrahedral coordination and LMCT features at 310 nm of intensity consistent with a single Cys-thiol, and both features saturate at ~ one molar equivalent of Co(II) (Fig. S2B, ESI†). Co(II)-titration of InrS previously showed saturation at ~ two molar equivalents of Co(II), a maximal stoichiometry which is consistent with the resolution of Co(II)-bound InrS via gel filtration chromatography (Fig. S3, ESI†).17 The ratiometric fluorescent metal chelator Fura-2 has been used previously to probe protein Co(II) affinities in the nM range.22 Incubation of Fura-2 with ZiaR, Zn$_2$Zur or InrS in the presence of a sub-stoichiometric
concentration of Co(II), sufficient to almost completely fill a single tightest metal binding site on each protein (Tables S2–S4, ESI†), resulted in partial Co(II)-occupancies of sensor-proteins and chelator as reported by an intermediate magnitude of quenching of Fura-2 fluorescence emission (Fig. 2). Using a determined $K_{Co(II)}$ for Fura-2 of 7.03 nM (Fig. S4, ESI†), $K_{Co(II)}$ of the tightest site per dimer for ZiaR and Zn1Zur is calculated to be 6.94 ($\pm$1.3) x 10^{-10} M and 4.56 ($\pm$0.16) x 10^{-10} M respectively (Fig. 2E; Fig. S2–S4, ESI†). In contrast to the other three sensors, CoaR failed to compete with Fura-2 implying $K_{Co(II)}$ at least one order of magnitude weaker than Fura-2, but $K_{Co(II)}$ must be tighter than the half-saturation [Co(II)] (~10^{-6} M) for linear Co(II)-dependent spectral features (Fig. 1B inset). Thus, unexpectedly, CoaR binds Co(II) more weakly than any of the other tested cytosolic metal sensors (Fig. 2F), raising questions about how CoaR gains access to Co(II) in vivo, in a cytosolic environment shared by the Zn(n)-sensors ZiaR and Zur.

**Zur outcompetes CoaR for Co(II)**

To test the order of affinity in Fig. 2F, the sensor with the tightest estimated $K_{Co(II)}$ (Zn1Zur) was competed with the weakest (CoaR). Zn1Zur does not contain tryptophan residues resulting in weak fluorescence emission when excited at 295 nm (Fig. 3A), unlike CoaR (Fig. 3B), enabling discrimination between metal-binding to the two proteins. In the presence of equimolar Zn1Zur quenching of CoaR fluorescence by Co(II) is lost, consistent with CoaR binding to the tighter but invisible site of Zn1Zur, using $\lambda_{em} = 295$ nm (Fig. 3C). In an analogous experiment, addition of ~1 molar equivalent of Zn1Zur also gave no quenching of CoaR fluorescence, but quenching occurred upon subsequent addition of Co(II), consistent with Zn1Zur preferentially binding and saturating Zn1Zur allowing Co(II) to subsequently partition to CoaR (Fig. 3D).

**$K_{Co(II)}$ is relatively weak for CoaR–DNA complexes**

As a MerR-like transcriptional regulator, activated Co(II)–CoaR remains associated with DNA. Therefore we explored the possibility that $K_{Co(II)}$ might be tighter for the DNA-adduct. Based on the estimated $K_{DNA}$ (Fig. 1), CoaR at 1.0 μM in the

![Fig. 2](image-url) Determination of Co(II) binding constants for each sensor by competition with the Co(II) binding chelator Fura-2. Co(II)-dependent quenching of Fura-2 fluorescence emission at 510 nm ($\lambda_{em} = 360$ nm) was measured in the absence of protein (closed symbols). Open symbols show the fluorescence emission, at equilibrium, following addition of InrS (A), Zn1Zur (B), ZiaR (C) and CoaR (D) to a mixture of Fura-2 and Co(II) under anaerobic conditions. Expected fluorescence values if Fura-2 fully outcompetes the protein for Co(II) are shown (arrows). Mean $K_{Co(II)}$ values for ZiaR, Zn1Zur and InrS, and a $K_{Co(II)}$ range for CoaR, are tabulated (with standard deviations) (E) and shown in graphical form (F). Numerical parameters used to calculate mean $K_{Co(II)}$ values are shown in Tables S2–S4 (ESI†).

![Fig. 3](image-url) (A) Fluorescence emission spectra for Zn1Zur (5 μM) in the presence (dotted line) and absence (solid line) of one molar equivalent of Co(II). (B) Emission spectra of CoaR (5 μM) in the absence (solid line) and presence (dotted line) of 0.9 molar equivalents of Co(II). (C) Emission spectra from a solution of equimolar (5 μM) apo-CoaR and Zn1Zur (solid line) and after addition of 0.9 molar equivalents of Co(II) (dotted line). (D) Emission spectra from an experiment analogous to (C) but with 0.9 molar equivalents Zn(II) (dashed line, under solid line) followed by 1.1 molar equivalents of Co(II) (dotted line). $\lambda_{em} = 295$ nm for all experiments.
presence of surplus (1.2 mM), coa-O/P DNA (without fluorescent label), should be fully DNA-bound. Fura-2 was titrated with Co(II) in the presence or absence of this CoaR–DNA complex and in common with an analogous experiment using CoaR alone, negligible Co(II) was withheld from Fura-2 (Fig. 4A and B). In contrast, control titrations with ZiaR and Zn1Zur do show evidence of Co(II) being withheld from Fura-2 (Fig. 4C and D). Thus, DNA-binding does not enhance the $K_{\text{Co(II)}}$ of CoaR and its weak affinity relative to the other sensors remains anomalous.

**Co(n) inhibits DNA-binding by ZiaR and promotes DNA-binding by Zur**

Although Co(n) binds tightly to the cytosolic cyanobacterial sensors for other metals, it might be ineffective at triggering the allosteric mechanism to alter DNA-binding and hence to modulate gene expression by these other proteins. This was tested in vitro by fluorescence anisotropy (Fig. 5). Addition of Co(n) to a pre-formed complex of ZiaR and hexachlorofluorescein-labelled zia operator–promoter DNA (zia-O/P) caused a decrease in anisotropy consistent with Co(n)-dependent DNA-dissociation (Fig. 5A), analogous to previous observations with Zn(n). Consistent with this observation, Co(n) also inhibited the association of ZiaR with zia-O/P DNA [Fig. S5, ESI†]. Addition of Co(n) to Zn1Zur caused an increase in anisotropy consistent with Co(n)-dependent DNA association by this metal-dependent co-repressor (Fig. 5B), which is similar to previous observations with Zn(n).21 Co(n) is also effective at dissociating InrS–DNA complexes (Fig. 5C). Thus, each of the other three sensors is competent to respond to Co(n) in vitro.

**Either z3N or z5 metal-sites of ZiaR are sufficient for allostery**

Only 0.5 molar equivalents of Co(n) were required to dissociate ZiaR–DNA complexes (Fig. 5A). This was unexpected because previously it was shown that mutation of putative metal ligands at either z5 or z3N sites impaired Zn(n) sensing in vivo. This implied that both z5 and z3N sites were needed for the protein to respond to Zn(n) whereas the stoichiometry in Fig. 5A implies that, at least for Co(n), binding to only one of four sites (per dimer) is needed to dissociate the protein from DNA. Variants of ZiaR were generated in which metal-binding residues at each of the sites were replaced individually or in combination with non-binding alternates. Titration of these variants with Co(n) confirmed binding of ~ one molar equivalent of metal solely to a thiol-free, and hence LMCT-free (at 310 nm), z5 site in the z3N mutants, and binding of ~ one molar equivalent of metal solely to a thiol-containing, and hence LMCT-generating, z3N site in the z5 mutants (Fig. 6). Double mutants missing both pairs of sites failed to dissociate from DNA upon addition of up to ten molar equivalents of Zn(n), as expected. However, mutants missing either individual pair of sites (at z5 or at z3N) remained inducer responsive (Fig. 6). Thus either site is sufficient to trigger the allosteric response and it is reasonable to consider the affinity of the tightest Co(n) site of ZiaR in comparisons with the other sensors.

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**Fig. 4** Titration of Fura-2 (1 μM) with Co(n) in the absence (closed symbols) or presence (open symbols) of CoaR (1 μM), either in the absence (A) or presence (B) of unlabelled coa-O/P DNA (1.2 μM). At these DNA and protein concentrations all of the CoaR should be DNA-bound. Titrations analogous to (A) but for ZiaR (C) and Zn1Zur (D) (1.5 μM Fura-2 and protein). All titrations in the presence of protein were performed anaerobically.

**Fig. 5** (A) Anaerobic titration of a complex of ZiaR (1 μM) and zia-O/P DNA (10 nM) with Co(n), monitoring fluorescence anisotropy. (B) Titration (in the presence of 1 mM TCEP) of znu-O/P DNA (10 nM) and Zn1Zur (100 nM) with Co(n), monitoring fluorescence anisotropy. (C) Anaerobic titration of a complex of InrS (1 μM) and nrsD-O/P DNA (10 nM) with Co(n), monitoring fluorescence anisotropy.
CoaR, but not ZiaR or Zur, responds after prolonged exposure to Co(II) \textit{in vivo}

ZiaR is reported to respond to Zn(II) and not Co(II), but CoaR the opposite \textit{in vivo}.\textsuperscript{12,15} In view of the observation that the two Zn(II) sensors ZiaR and Zur bind Co(II) at least two orders of magnitude more tightly than CoaR (Fig. 2) and in both cases Co(II) alters DNA-binding in a manner analogous to Zn(II) (Fig. 5), we coincidently tested the abundance of \textit{ziaA}, \textit{znuA} and \textit{coaT} transcripts, regulated by ZiaR, Zur and CoaR respectively, in common populations of nucleic acids isolated from cells exposed to elevated Zn(II) (16 μM) or Co(II) (2 μM) (maximum non-inhibitory concentrations) for 48 h (Fig. 7A). Transcripts encoded by the gene activated by CoaR, namely \textit{coaT}, accumulated in response to Co(II) not Zn(II), transcripts de-repressed by ZiaR, \textit{ziaA}, accumulated in response to Zn(II) not Co(II), while the abundance of transcripts repressed by Zn, Zur, \textit{znuA}, declined in response to Zn(II) but not Co(II). Thus, the observed selectivity precisely replicates predictions based upon the literature.\textsuperscript{12,15,19} \textit{InrS} is known to respond to Co(II) as well as Ni(II) and was not included in these analyses.\textsuperscript{17}

\textbf{Co(II) does initially trigger ZiaR \textit{in vivo}}

Failure to de-repress expression of \textit{ziaA} in response to 48 h exposure to Co(II) implies that by the end of this incubation ZiaR does not have access to excess Co(II). We tested whether shorter (1 h) exposure to Co(II) might elicit a response. Fig. 7B shows accumulation of \textit{ziaA} and \textit{ziaR} transcripts under these conditions. Both these genes are regulated by ZiaR. Thus ZiaR is responsive to Co(II) \textit{in vivo} as well as \textit{in vitro} but after longer exposure (48 h) Co(II) is somehow sensed by CoaR but not by ZiaR despite the >100 fold tighter \textit{K}_{Co(II)} of the latter.

\textbf{A deduced tetrapyrrole binding-site in CoaR is required for cobalt-sensing}

CoaR contains a domain with homology to precorrin isomerase, an enzyme which catalyses a methyl isomerisation reaction required during the biosynthesis of vitamin B\textsubscript{12} from precursor tetrapyrrole molecules.\textsuperscript{12} Homologous proteins catalysing this reaction in organisms utilising the so-called aerobic B\textsubscript{12} synthesis pathway are termed CobH and those from anaerobic B\textsubscript{12} synthesis pathways are termed CbiC.\textsuperscript{24} Sequence alignment of CoaR and \textit{Pseudomonas denitrificans} CobH reveals that only two out of the eleven residues (Ser17 and Ala44), that in CobH form hydrogen bonds to its reaction product hydrogenobyrinic acid (HBA), are absolutely conserved in CoaR (Ser120 and Ala44) (Fig. S7, ESI†). Using the structure of \textit{P. denitrificans} CobH in which the tight-binding hydrogenobyrinic acid molecule was co-visualised,\textsuperscript{25} a dimeric model of the precorrin isomerase-like domain of CoaR was produced that included hydrogenobyrinic acid associated with modelled CoaR tetrapyrrole binding sites (Fig. S8, ESI†). By making a comparison with the analogous structure of CobH from \textit{P. denitrificans}, it was possible to identify residues in CoaR in equivalent spatial locations to the residues known to form hydrogen bonds to the tetrapyrrole in CobH (Fig. 8; Fig. S8 and Table S5, ESI†). Three of these CoaR residues (Trp265, Leu191 and Gly342) are incapable of forming side chain hydrogen bonds while Ala192 overlays a residue in CobH (Ala44) which forms a hydrogen bond from a backbone nitrogen atom. Site directed mutagenesis was performed on the remainder to test for effects on Co(II)-dependent transcriptional activation. Conspicuously, Tyr287 protrudes into the predicted tetrapyrrole binding site at the opposite face to a ligand in CobH, Tyr14, and this has also been mutated. Loss and gain of Co(II)-responsive expression of a reporter gene (\textit{lacZ}) driven from the \textit{coaT} operator–promoter correlated with Tyr287 substitutions which lose (Phe) and restore (Glu) hydrogen bonding, respectively (Fig. 8). Co(II)-responsive expression was lost or impaired by Ala substitutions at five of the remaining six deduced ligand positions (Fig. 8). The exception (Ser290) overlays with a residue (Thr140) which forms one of two hydrogen bonds to a functional group in the tetrapyrrole and hence its loss may be compensated by an alternate bond. Three of the substitutions (Ser167Ala, His267Ala and Tyr287Phe) gave no Co(II)-response (Fig. 8) and two of these variants were over-expressed and purified from \textit{E. coli}, and both confirmed to form complexes with the \textit{coa}-O/P DNA with similar \textit{K}_{DNA} to wild type CoaR (Fig. 9), consistent with loss of effector recognition rather than global mis-folding. In \textit{Synechocystis} PCC 6803 Δ\textit{cbiE}, unable to

![Fig. 6](https://example.com/fig6.png)  
**Fig. 6** Anaerobic titration of a complex of either Δα3N Δα5 ZiaR (circles), Δα5 ZiaR (squares) or Δα3NΔα5 ZiaR (triangles) (each at 1 μM) and zia-O/P DNA (10 nM) with Zn(II), monitoring fluorescence anisotropy.

![Fig. 7](https://example.com/fig7.png)  
**Fig. 7** (A) Abundance of \textit{ziaA}, \textit{znuA} and \textit{coaT} transcripts in common populations of RNA analysed by RT-PCR following 48 h exposure to Zn(II) (16 μM) or Co(II) (2 μM). (B) Abundance of \textit{ziaA} and \textit{ziaR} transcripts in response to the same concentrations of Zn(II) and Co(II) following 1 h exposure. \textit{rps1} is included as a control in both cases.
synthesise the substrate for precorrin isomerase, expression of a reporter gene from the coaT operator–promoter was enhanced and this was taken to imply that intermediates in the vitamin B12 biosynthetic pathway are inhibitory to transcriptional activation by CoaR. In contrast, the observations reported here suggest that a deduced binding site for such molecules in CoaR is required for Co(II)-dependent transcriptional activation.

A carboxyl-terminal Co(I)-site in CoaR is required for cobalt-sensing

If CoaR sensed a tetrapyrrole rather than directly sensing Co(I) this could account for its response to Co(I) in vivo despite its relatively weak $K_{\text{Co(II)}}$. However, previously a triple mutant within a carboxyl-terminal CysHisCys motif in CoaR was shown to lose Co(I)-dependent transcriptional activation and it was suggested that this was a likely Co(I) binding site. Another residue (Cys121) is conserved in CoaR and acts as a metal ligand in the homologous E. coli MerR-like sensors ZntR (Zn(II)) and the CueR (Cu(I)).

We have examined the effects of mutation of Cys121 and the first Cys residue from the carboxyl-terminal CysHisCys motif (Cys363) on the Co(I)-dependent spectral features of purified CoaR (Fig. 10 inset). Conversion of either Cys to Gly reduced the intensity of the LMCT features at 310 nm by $10^3$ M$^{-1}$ cm$^{-1}$ consistent with loss of a single Co(I)–S bond in each mutant. Both variants showed either no, or negligible, Co(I)-responsive expression of a reporter gene from the coaT operator–promoter when examined in E. coli (Fig. 10).

Discussion

In this research we observe binding of the cobalt responsive transcriptional activator CoaR to Co(I) and to DNA (Fig. 1), but unexpectedly estimate its $K_{\text{Co(II)}}$ to be the weakest among a sub-set of metal-sensors from Synechocystis PCC 6803: a sub set which includes one sensor from each family encoded by this genome (Fig. 2). This relatively weak $K_{\text{Co(II)}}$ is supported by

Fig. 8 A simplified model of the predicted tetrapyrrole binding site of CoaR is shown (central panel) in complex with hydrogenobyrinic acid. Residues spatially analogous to tetrapyrrole ligands in CobH are annotated (see Fig. S8, ESI†). Outer panels show expression from the coaT promoter in E. coli cells containing pET3a coa and variants of coaR, in the absence (open bars) or presence (closed bars) of 100 μM CoCl$_2$. In each dataset the left hand bars represent a wild-type CoaR control performed in the same assay. All data are mean activities from assays performed in triplicate shown with standard deviations.
analyses of metal-dependent quenching of intrinsic CoaR fluorescence in the presence of zinc-sensing Zn$_1$Zur (estimated to have the tightest $K_{	ext{Co(II)}}$ of the set of sensors) (Fig. 3). Quenching of CoaR fluorescence upon addition of one molar equivalent of Co(II) is lost in the presence of equimolar Zn$_1$Zur, consistent with Co(II) binding to the tighter sites of the Zn(II)-sensor, but quenching is regained if the sensory sites of Zn$_1$Zur are first saturated with a molar equivalent of Zn(II) (Fig. 3). These data indicate that $K_{	ext{Zn(II)}}$ Zn$_1$Zur is tighter than $K_{	ext{Co(II)}}$ CoaR, consistent with the former but not the latter detecting Zn(II) in vivo, and in contrast with the inverse correlation between $K_{	ext{Co(II)}}$ and CoaR-sensing. The results for Co(II) also contrast with the order of affinity for $K_{	ext{Ni(II)}}$ of the same set of proteins in which the cytosolic Ni(II)-sensor, InrS, has the tightest $K_{	ext{Ni(II)}}$.17 Ni(II) and potentially Zn(II), are thus expected to partition to the sensor of tightest affinity, triggering expression of efflux, preventing the buffered cytosolic concentration of Ni(II) or Zn(II) from rising to a sufficiently high level to bind to other sensors of weaker $K_{	ext{Ni(II)}}$ and $K_{	ext{Zn(II)}}$. In contrast, the specific detection of Co(II) by the weakest Co(II)-binding sensor CoaR is enigmatic.

The possibility that the $K_{	ext{Co(II)}}$ of CoaR may be enhanced by conformational changes in the CoaR–DNA adduct was explored, since this adduct is the active species. Co(II)–CoaR is inferred to induce transcription from the coa operator–promoter by under winding DNA in a mechanism analogous to Hg(II)–MerR.12,13,27 In the presence of excess coa operator–promoter at a concentration at least an order of magnitude greater than $K_{	ext{DNA CoaR}}$ (Fig. 1A), CoaR failed to compete with Fura-2 for Co(II) (Fig. 4). Both ZiaR and Zn$_1$Zur do compete with Fura-2 for Co(II) (Fig. 2 and 4). The possibility that selectivity may not relate to Co(II) binding but may instead be due to Co(II) failing to trigger the allosteric mechanisms of ZiaR and Zur was investigated next. However, Co(II) does drive DNA-dissociation of ZiaR and encourages DNA-association by Zn$_1$Zur (Fig. 5).

CoaR, ZiaR and Zur show perfect discrimination between Co(II) and Zn(II) after prolonged exposure (48 h) of cells to maximum non-inhibitory metal concentrations but crucially this selectivity is not detected for ZiaR after short term (1 h) exposure (Fig. 7). While the latter observation is consistent with in vitro data for ZiaR (Fig. 5), the former is not. The relative abundance of CoaR and ZiaR in the cell has not been determined but the mode of action of CoaR implies that mass action could not provide a bias in favour of CoaR detecting cobalt. The precedent is that metal-bound forms of MerR-like activators have slightly weaker $K_{	ext{DNA}}$ than apo-forms and hence a larger number of CoaR molecules would further dis-favour activation of the coaT operator–promoter in response to Co(II).27 Failure to respond to cobalt at 48 h is inferred to reflect an in vivo regime under which CoaR somehow gains access to surplus Co(II) in preference to ZiaR. This kinetic contribution must be sufficient to overcome a dis-favourable thermodynamic gradient of at least two orders of magnitude (Fig. 2F).

A requirement for 0.1% w/v $n$-dodecyl $\beta$-D-maltoside to purify CoaR suggests association with membranes via its precorrin isomerise-like domain. Enzymes of vitamin B$_{12}$ biosynthesis are
known to be membrane associated, and one of the homologues of precurrin isomerase in Synechocystis PCC 6803 (Srl1467) has been shown by proteomics to be membrane associated. Srl1467 shares sequence similarity (45%) with the precurrin isomerase-like domain of CoaR and the two proteins have analogous predicted hydrophobic carboxyl-terminal regions (Fig. S1, ESI†). Negative charge associated with the inner leaflet of plasma-membranes would be attractive to cations such as Co(II), raising the possibility that Co(n) and CoaR encounter at the membrane whereas Co(n) and ZiaR or Zur or Zur encounter in the cytosol. Ignoring any contribution from thylakoid membranes, if the radius of the cell is ~ three orders of magnitude greater than that of CoaR (Fig. S9, ESI†), a three dimensional search between Co(n) and ZiaR or Zur would take approximately two orders of magnitude longer than a two dimensional search for CoaR (associated with one of the multiple chromosome copies) at the membrane (Fig. S9, ESI†).

The pathway for vitamin B12 biosynthesis is also known to be membrane associated in other bacteria with intermediates channelled between enzymes. The precurrin isomerase-like domain of CoaR might be expected to associate with this machinery, in common with the precurrin isomerase enzyme that acts in the pathway. Previously, bacterial two-hybrid interactions were not detected between the cobalt-exporter CoaT and the cobalt-chelatase for B12 biosynthesis CbiX, but interaction with the cobalt importer HupE or with CoaR is formally possible, and might enable Co(n) to be channelled to CoaR.

Mutation of a candidate tetrapyrrole binding site in CoaR suggests that B12, one of its precursors or a related tetrapyrrole, is required for Co(n)-dependent transcriptional activation (Fig. 8). Notably, this requirement was observed in E. coli which does not possess a complete B12 biosynthetic pathway and imports B12. The siroheme ferrochelatase (CysG) of E. coli can however introduce Co(n) into tetrapyrroles, but CoaR remains responsive to Co(n) in AcysG E. coli (Fig. S10, ESI†), implying that CoaR does not detect a Co(n)-containing product of this enzyme. A hyper-responsive phenotype was previously observed in Synechocystis PCC 6803 ΔchbiE and it was inferred that this mutant is missing a tetrapyrrole that represses CoaR. However, cobalt is inserted into the corrin ring sooner than previously anticipated in this organism via the so-called anaerobic pathway, and an explanation consistent with current data (Fig. 8), is that there is hyper-accumulation of a Co(n)-containing activator of CoaR in ΔchbiE. In this alternative scenario the tetrapyrrole may either be the sole activator, potentially channelled to CoaR via the B12 assembly pathway (albeit it is unclear how a Co(n)-cofactor is generated in E. coli), or a co-activator with additional channelling of cobalt to a site involving Cys121 associated with the MerR-like domain, and carboxyl-terminal Cys363 (Fig. 10). Potentially, K_{Co(n)} may be enhanced on binding tetrapyrroles.

In conclusion, these studies highlight the merit in considering selectivity in metal sensing as an integrated system: a combined function of a cells set of sensors (a metallomic approach), rather than solely a property of individual sensors acting in isolation. For Ni(n), and provisionally Zn(n), specificity matches the thermodynamic gradient among the set of sensors. However for Co(n) this is not the case. This uncovers a requirement for additional studies to explain the mechanism of action of CoaR and its interplay with vitamin B12 biosynthesis.

Methods

General reagents, bacterial cultures and cloning procedures

All reagents and chemicals were sourced from standard suppliers. E. coli strains DH5α, JM101 and BL21(DE3) were grown in standard Luria Bertini (LB) medium. Constructs (in pET29a) for overexpression of wild-type CoaR, ZiaR, Zur and InrS were produced previously, followed by ligation to Bsal and Sali sites to form pET3aacoa (the derivative of pET3a also contained cbiX controlled by an IPTG-responsive promoter which was not activated in these studies). Mutated variants of coaR were produced by QuikChange mutagenesis (primers in Table S1, ESI†). Due to the large size of pET3aacoa, nicks were first sealed using T4 DNA ligase (2 μl 10× T4 DNA ligase buffer (Promega), 1 μl T4 DNA ligase (Promega), 15 μl H2O, 2 μl DNA, 15 °C for 4 h) followed by enzyme inactivation at 65 °C for 20 min. All gene variants were confirmed by sequencing.

Protein expression and purification

Expression and purification of ZiaR, Zn2Zur, CoaR InrS have been described previously. Mutated variants of ZiaR and Δz3N ZiaR (C71S/C73S), Δx5 ZiaR (H116R), and Δz3NΔx5 ZiaR variants, were overexpressed in E. coli BL21(DE3) and purified using the same procedures as for wild-type ZiaR, with the exception that HiTrap heparin affinity chromatography (GE Healthcare) replaced Ni(n)-affinity chromatography for Δz3NΔx5 ZiaR (which binds weakly to Ni(n)-affinity matrices). C121G and C363G CoaR variants were expressed and purified as described previously for wild-type ZiaR. S167A and Y287F CoaR variants were expressed as described previously, but purified by refolding protein recovered from inclusion bodies, re-suspended in 4 ml of 500 mM NaCl, 10 mM Hepes (pH 7.8), 5 mM DTT, 8 M urea, 1% w/v SDS. Post incubation (37 °C, 1 h), supernatant (13 400 × g, 10 min) was diluted 10-fold into 500 mM NaCl, 5 mM DTT, 10 mM Hepes (pH 7.8), 0.5% w/v n-dodecyl β-D-maltoside, further incubated (1 h) and supernatant (13 400 × g, 10 min) diluted 50-fold into 100 mM NaCl, 1 mM DTT, 10 mM Hepes (pH 7.8), 100 mM imidazole, 1 mM EDTA, 0.05% w/v n-dodecyl β-D-maltoside. This solution was applied to a 1 ml HiTrap heparin affinity column (GE Healthcare) pre-equilibrated in the same buffer and subsequently eluted in buffer containing 1M NaCl and fractions analysed by SDS-PAGE.

ZiaR, Zur, CoaR and InrS (plus variants of ZiaR and CoaR) were quantified as described previously, with the exception of Y287F CoaR which has an altered theoretical extinction
coefficient ($\varepsilon = 2.5 \times 10^{-4} \text{M}^{-1} \text{cm}^{-1}$). Proteins were confirmed to be >95% apo by ICP-MS analysis. Anaerobic stocks of protein were confirmed to be >95% reduced by reaction with 5,5'-dithiobis-(2-nitrobenzoic acid) (DTNB) as described previously. CoaR showed inconsistent thiol-reactivity although DNA-binding and metal-binding remained consistent.

**UV-visible absorption spectroscopy**

Proteins were diluted into chelex-treated, N$_2$-purged buffer (containing 400 mM KCl, 100 mM NaCl, 10 mM Hepes (pH 7.8) and, in the case of CoaR, 0.1% w/v n-dodecyl β-D-maltoside) in an anaerobic chamber. UV-visible spectra were collected under anaerobic conditions, using sealed, gas tight quartz cuvettes, using Cary 4E UV-visible (Varian, UK) or Perkin Elmer λ35 spectrophotometers. Concentrations of ZnCl$_2$ and CoCl$_2$ stocks used in titrations were verified by ICP-MS.

**Fluorescence anisotropy**

Fluorescent zuu-O/P, nrsD-O/P and zia-O/P probes have been described previously. Oligonucleotides (primers 25 and 26, Table S1, ESI†) containing a region of the coa operator–promoter including the 13-6-13 CoaR binding site were annealed as described previously to produce 36 bp hexachlorofluorescein-labelled coa-O/P. Wild-type and mutant forms of apo-CoaR protein were titrated against 10 nM coa-O/P in buffer containing 150 mM NaCl, 5 mM DTT, 1 mM EDTA, 200 mM imidazole, 10 mM Hepes (pH 7.8). For anaerobic dissociation experiments with ZiaR and ImRs, protein–DNA complexes were pre-formed by adding protein to a final concentration of 1 µM in a sealed quartz cuvette containing 10 nM DNA in chelex-treated, N$_2$-purged buffer containing either 120 mM KCl, 30 mM NaCl, 10 mM Hepes (pH 7.8) (for ZiaR) or 240 mM KCl, 60 mM NaCl, 10 mM Hepes (pH 7.8) (for ImRs). Protein–DNA complexes were titrated with Co(II) under anaerobic conditions. A solution of Zn$_1$Zur and Zn$_1$Zur (1.5 µM protein, 1.5 µM Fura-2) (no DNA was present). The affinity of Fura-2 under the buffer conditions used in these analyses (400 mM KCl, 100 mM NaCl, 10 mM Hepes pH 7.8) was determined directly by competition with the nitriultricatic acid (NTA) which binds Co(II) with $K_{\text{Co(II)}} = 3.59 \times 10^{-9} \text{M}$ at pH 7.8 (Fig. S4, ESI†).

**CoaR-Zur Co(i) competition**

Recombinant, wild-type CoaR and Zn$_1$Zur were each diluted separately to 5 µM in chelex-treated, N$_2$-purged buffer containing 400 mM KCl, 100 mM NaCl, 10 mM Hepes (pH 7.8) and 0.1% w/v n-dodecyl β-D-maltoside in an anaerobic chamber. Fluorescence emission spectra ($\lambda_{ex} = 295 \text{ nm}$ for all experiments) were collected in sealed quartz cuvettes for apo-proteins and following addition of Co(i). CoaR and Zn$_1$Zur were competed for Co(i) by diluting both to 5 µM in a quartz cuvette. Fluorescence emission spectra were collected for apo-proteins, one molar equivalent of CoaR was then added in an anaerobic chamber and the emission spectrum re-recorded. In an analogous experiment 0.9 molar equivalents Zn(i) was added followed by 1.1 molar equivalents of Co(a).

**RNA-isolation and reverse transcriptase PCR**

Cells were inoculated to an OD$_{595}$ of 0.1 and cultured under standard conditions with maximum non-inhibitory concentrations of ZnSO$_4$ (16 µM) or CoCl$_2$, [2 µM] (concentrations determined in growth experiments to give <10% growth inhibition), for either 1 h or 48 h. Nucleic acid was isolated from logarithmically growing Synechocystis PCC 6803 using an established method. Nucleic acids were treated with DNase I (Sigma) and 1 µg used to produce cDNA using an Im-PromII reverse transcription kit with random hexameric primers (Promega). Reverse transcriptase was omitted from negative control reactions, which in all cases confirmed the absence of contaminating DNA. PCR primers for ziaA, rps1, znaA (described previously) and coaT (primers 23 and 24, Table S1, ESI†) were used to amplify ~300 bp of each gene. Cycling conditions included denaturation (95 °C for 2 min) then 25–30 cycles of 95 °C 1 min, 60 °C 1 min, 72 °C 20 s, followed by a final extension step at 72 °C for 5 min. Products were analysed on 1% w/v agarose gels.

**Structural modelling**

A dimeric model of the precorrin isomerase-like domain of CoaR bound to hydrogenobyrinic acid (HBA) was produced by threading residues 162–358 onto the co-crystal structure from P. denitrificans (PDB code: 1I1H) using SwissModel (http://swissmodel.expasy.org/). The resulting monomer was converted to a dimer and aligned with CobH in WinCoot, and the dimer analysed in PyMol and Swiss PDB Viewer.

**β-Galactosidase assays**

JM101 cells containing pET3acoa (and variants), grown at 37 °C overnight were diluted 100-fold into fresh LB medium and
cultured in the absence or presence of 100 μM CoCl₂ (maximum non-inhibitory concentrations used previously) at 37 °C until an OD₅₉₅ nm of 0.2–0.3 was reached. Cells were assayed in triplicate for β-galactosidase activity as described previously.  

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References


