A single sensor controls large variations in zinc quotas in a marine cyanobacterium

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Marine cyanobacteria are critical players in global nutrient cycles that crucially depend on trace metals in metalloenzymes, including zinc for CO₂ fixation and phosphorus acquisition. How strains proliferating in the vast oligotrophic ocean gyres thrive at ultra-low zinc concentrations is currently unknown. Using Synechococcus sp. WH8102 as a model we show that its zinc-sensor protein Zur differs from all other known bacterial Zur proteins in overall structure and the location of its sensory zinc site. Uniquely, Synechococcus Zur activates metallothionein gene expression, which supports cellular zinc quotas spanning two orders of magnitude. Thus, a single zinc sensor facilitates growth across pico- to micromolar zinc concentrations with the bonus of banking this precious resource. The resultant ability to grow well at both ultra-low and excess zinc, together with overall lower zinc requirements, likely contribute to the broad ecological distribution of Synechococcus across the global oceans.

All major biogeochemical cycles, including those for carbon, nitrogen and phosphorus, are catalyzed by multiple enzymes, many of which require metal ions for activity. Therefore, all organisms involved in these cycles must ensure that they acquire appropriate amounts of the entire panel of essential metals. This also holds true for microorganisms that inhabit the most micronutrient-depleted regions of the open ocean, including photosynthetically active cyanobacteria of the genera Synechococcus and Prochlorococcus. Together, these smallest but most abundant photautotrophs contribute an estimated one-quarter of marine net primary production and hence are major drivers of the global carbon cycle. Much remains to be elucidated regarding their metal ion requirements, uptake and utilization strategies.

One element that has received comparatively little attention in this context is zinc. Typically, oceanic zinc concentrations follow a nutrient-like distribution, with pico- to nanomolar concentrations in surface waters. Although the importance of zinc for eukaryotic phytoplankton is undisputed, evidence for the zinc limitation of photosynthetically active cyanobacteria in the largest but most abundant photautotrophs contribute an estimated one-quarter of marine net primary production and hence are major drivers of the global carbon cycle. Much remains to be elucidated regarding their metal ion requirements, uptake and utilization strategies.

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by reduced zinc tolerance and altered zinc accumulation. Structural characterization of the recombinantly expressed SYNW2401 protein reveals unique features including a new zinc-sensing site. RNA-sequencing (RNA-seq) establishes a small set of genes regulated by Zn\(^{2+}\) and Zur, including a Zur-repressed znuABC uptake system and, unprecedentedly, a Zur-activated metallothionein that enables safe accumulation of intracellular Zn\(^{2+}\) and expands the range of zinc concentrations at which this strain can thrive.

**Results**

**Disruption of synw2401 alters zinc quota and tolerance.** Previous genome-mining identified several potential players in zinc homeostasis in (marine) cyanobacteria\(^{30,39}\). According to our analyses, which included analysis of multiple genome neighborhoods, WH8102 harbors two putative ABC (ATP-binding-cassette)-type zinc-uptake transporter systems (ZnuABC), encoded by the genes synw2479–synw2481 and synw0969–synw0971. A Zur-binding site (Zur box; Fig. 1a,b) is predicted in the intergenic region between synw0970 (znuC) and synw0971 (znuA)\(^{36}\), but none for synw2479–synw2481. However, two further Zur boxes are predicted in the promoter region for a gene encoding a bacterial metallothionein (synw0359, bmtA). Because inference of metal specificity by bioinformatics is not straightforward\(^{18}\), it is critical to experimentally establish whether the predicted Zur protein (SYNW2401) is indeed involved in zinc homeostasis and a true zinc sensor.

We therefore generated a single crossover interposon mutant (Extended Data Fig. 1) in which the synw2401 gene was disrupted. The metal-specific phenotype of this mutant strain was investigated first, focusing on growth and metal accumulation when cultured in chelexed and metal-supplemented artificial seawater (ASW) medium\(^{27}\) (Fig. 1c–e). At low zinc (0 or 77.2 nM added Zn\(^{2+}\)), wild-type (WT) and mutant strains grew equally well, whereas at the 'standard' ASW zinc concentration (772 nM\(^{27}\)), the mutant began to show growth impairment (Fig. 1c and Supplementary Fig. 1). At 2.5 μM zinc, the mutant was unable to grow, in contrast to the WT which only showed relatively mild growth impairment. Further increase in [Zn] exceeded the zinc tolerance of the WT as well. Cellular metal quotas, expressed as mmol metal per mol phosphorus, were determined at two (zur mutant) or three (WT) zinc concentrations (Fig. 1d,e and Extended Data Fig. 1c). At both 0 and 772 nM zinc added to the culture medium, the most severely altered metal quotas were those of zinc. Notably, although media had been treated with Chelex resin, the medium with 0 Zn added evidently still supplied sufficient zinc to be accumulated in the WT and the mutant, with no indication of zinc limitation, as previously observed\(^{26}\). In all cases, the determined zinc quotas were broadly within the ranges reported for both field samples (0.5–52 mmol Zn: mol P\(^{13}\)) and laboratory cultures (0.6–8.3 mmol Zn: mol P\(^{29,30}\)) of marine *Synechococcus* and other marine cyanobacteria. Quotas for other metals were also in ranges comparable with literature data, although trending toward the high end, owing to ASW being a comparatively rich medium.

At 0 added zinc, the mutant accumulated 3.6 times more zinc than the WT, whereas the quotas of all other metals inspected either decreased (Mn, Fe, Co) or remained unchanged (Ni, Cu). These observations are consistent with SYNW2401, like other Zur proteins, repressing transcription of (at least) znuA (synw0971), encoding a periplasmic binding protein. The absence of SYNW2401 in the mutant then leads to complete de-repression of znuA and hence maximal zinc import through the associated ZnuABC system. The drop in the quotas of other metals may indicate the operation of compensatory processes aiming to reduce metal influx nonspecifically, or could be related to mis-metallation of sensors for other metals.

As expected, the zinc quotas of both WT and mutant increased upon addition of zinc (772 nM) to the culture medium. Zinc quotas increased by factors of 6.5 (mutant) and 43 (WT), with the WT
accumulating more zinc overall than the mutant (Fig. 1c). Despite this higher cellular quota, the WT showed no growth impairment, although the mutant did. Moreover, although mild growth impairment was evident at 2.5 μM Zn, the WT was able to sustain a 129-fold increased cellular zinc quota compared with growth at 0 added Zn. This suggests that SYNW2401 also regulates a process that supports zinc accumulation without eliciting toxicity. The molecular basis for this remarkable ability to sustain zinc quotas that vary over two orders of magnitude is discussed later.

Thus, the WH8102 synw2401 mutant is characterized by altered accumulation of zinc and reduced tolerance to excess zinc. Together with previous bioinformatics analyses33,34, the results from these phenotyping studies demonstrate that SYNW2401 indeed corresponds to the zinc sensor Zur. Because no structural information for any cyanobacterial Fur-family protein is available, and the zinc-binding residues for sensory sites known from other Zur proteins are not conserved in cyanobacterial Zurs (Extended Data Fig. 2), we determined the structure of SYNW2401 (referred to as SynZur henceforth) by X-ray crystallography.

**Cyanobacterial Zurs differ from other Zur proteins.** SynZur was recombinantly overexpressed with a tobacco-etch-virus-protease-cleavable His-tag in *Escherichia coli*, using standard culture medium without additional metal supplementation, and the protein was purified using an approach that avoids denaturation and metal loss (Extended Data Fig. 3). The only metal ion that was present in substantial abundance was Zn2+ (2 molar equivalents per subunit; Fig. 2a and Supplementary Table 1).

The molecular mass derived from size-exclusion chromatography (SEC; Extended Data Fig. 3c) did not allow conclusive derivation of the oligomeric state, but nondenaturing SDS–PAGE (Extended Data Fig. 3d) and dynamic light scattering (Supplementary Fig. 2) results are both consistent with the protein being predominantly present as a dimer. Treatment with EDTA led to the loss of one of the two bound zinc ions (Fig. 2b). This process likely corresponds to zinc sensing. Indeed, although Zn2SynZur as isolated binds to the znuA promoter (as a dimer; Supplementary Fig. 3), the presence of EDTA abolished binding (Fig. 2c). This process is reversible, because addition of Zn2+ to EDTA-treated SynZur re-established DNA-binding ability (Fig. 2c). The remaining zinc ion in Zn2SynZur likely corresponds to a ‘structural’ site; the corresponding sites in other Fur-family proteins have repeatedly been found to be refractory to removal by EDTA33,34. The Zn2+-binding affinity of the EDTA-responsive site, that is the sensory site, was measured by spectrophotometric titration in competition with 2-[[8-[[bis(carboxylatomethyl)amino]-6-methoxyquinolin-2-yl]methoxy]-N-(carboxylatomethyl)-4-methylanilino]acetate (Quin-2) (Fig. 2d), giving a dissociation constant (Kd) of 8.3×10⁻¹⁴ M, similar to those measured for other Zur proteins (6.4×10⁻¹³ to 5.5×10⁻¹⁴ M)35.

Single crystals suitable for X-ray analysis were obtained in Mg(OAc)/MES buffer, pH 6, from protein purified by SEC, without further addition of zinc. The structure was solved to a resolution of 2.1 Å (Supplementary Table 2) employing single-wavelength anomalous diffraction with fluorescence detected at the zinc K absorption edge (9,666 eV). This approach was necessary because molecular replacement using a range of bacterial Fur-family proteins failed—indicating that SynZur adopts a structure that substantially differs from previously determined structures. The asymmetric unit of the crystal with the space group P63 contains four protein molecules. Interface analysis by PISA36 is consistent with SynZur forming a homodimer (Fig. 3a), with two dimers present in the asymmetric unit (Extended Data Fig. 4a). Like other Fur-family proteins33, each SynZur monomer consists of two domains, an N-terminal ‘winged helix’ domain that mediates interactions with DNA (DNA-binding domain (DBD); residues P6–A72) and a C-terminal domain that provides the dimerization interface (dimerization domain (DD); residues R76–P128) (Fig. 3b). The two domains are connected by a short ‘hinge’ (residues P73–D75).

Each monomer has two zinc ions bound with bond lengths that are within the expected ranges (Supplementary Table 3); one is bound tetrahedrally by four Cys residues (83, 86, 123 and 126) and corresponds to the structural site mentioned previously (Fig. 3b; site 1). The residues forming this site are (with a single exception) 100% conserved in cyanobacterial Zur sequences (Supplementary Fig. 4). Site 1 is located in the DD and tethers the C terminus to a region close to the second zinc site, which is formed by D77, H79, C95 and H115 (site 2, Fig. 3b). To confirm that this tetrahedral site is involved in zinc sensing, we generated a Cys95Ala mutant protein (Extended Data Fig. 5). Electrospray ionization mass spectrometry (ESI-MS) analysis of this mutant showed that the purified protein retained only one zinc ion. The mutant also displayed a similar elution volume in SEC, with no indication of dissociation of the dominant dimer, indicating that loss of sensory zinc does not lead to the dissociation of the dimer at concentrations accessible to SEC. However, electrophoretic mobility shift assay (EMSA) experiments demonstrate that the mutant is unable to interact with Zur boxes (Extended Data Fig. 5d). This strongly supports the notion that site 2 is involved in zinc sensing.

The ligand sphere (N2O5) of this new sensory zinc site in SynZur is very similar to sites found in other Zur proteins, including the single sensing site in *E. coli*38 and *Xanthomonas campestris*39 Zur, and the primary sensing sites in *Mycobacterium tuberculosis*35 and *Streptomyces coelicolor* Zur40. However, most remarkably, the SynZur sensory site is in a location that differs from all other confirmed sensory sites in Fur-family proteins37. (Extended Data Figs. 2 and 6). These invariably lie between the DBD and DD involving one or two residues from the DBD, one or two from the hinge region, and one or two from the DD. This inter-domain location previously provided a straightforward understanding of the canonical sensing mechanism in Fur-family proteins: the mutual orientation between DD and DBD is not fixed in the absence of the sensory metal, whereas the presence of this metal stabilizes a conformation of the dimer in which the two DBDs are optimally oriented to match the binding sites on the cognate DNA33,35,39,41. By contrast, three of four of the corresponding metal-binding residues are absent in SynZur (Extended Data Fig. 2), and its sensory site does not involve any residues from the DBD (Fig. 3b). Although some Fur-family proteins harbor additional metal-binding sites in an analogous location35,41, SynZur is the first Fur-family protein in which this site is the sole sensory site. The absence of the canonical sensory site and the presence of this new alternative sensory site are essentially conserved in Zur proteins from both marine and freshwater cyanobacteria (Supplementary Fig. 4), with H115 being 100% conserved, H79 and C95 being fully conserved with a single exception, and D77 being present in 86.8% of sequences, occasionally (11.0%) replaced by a histidine residue, or in rare cases separated by two instead of one residue from H79. The absence of an inter-domain zinc-binding site appears to be partially compensated by a network of hydrogen bonds and salt bridges that support this conformation and may also communicate the presence of Zn2+ in the sensory site to the DBD (Fig. 3c).

The two DBDs in either dimer can be superimposed with the two DBDs in either *Streptomyces Zur* (1.60 Å root mean squared deviation (r.m.s.d.) over 484 backbone atoms for dimer 2 (chains C+D); Extended Data Fig. 7a) or *E. coli Zur* (1.87 Å r.m.s.d. over 408 backbone atoms; Extended Data Fig. 7b) dimers. Separately, the DDs also match well with those for other Fur-family proteins (r.m.s.d. 1.40–1.80 Å for two DDs, over 212–264 backbone atoms; Extended Data Fig. 7). However, in all cases, it is impossible to simultaneously align both DBDs and DDs in either monomer or dimer. This is due to the mutual orientation of these two domains being ‘rotated’
with respect to these other proteins (Extended Data Fig. 7a). Thus, SynZur not only harbors a new zinc-sensing site, but also displays a unique orientation of DD and DBD.

SEC and CD spectroscopy of SynZur before and after treatment with EDTA (Supplementary Fig. 5) revealed no changes in shape, oligomerization state or secondary structure. The latter observation is not unexpected; the X-ray structures of apo- and holo-Zur from Xanthomonas campestris display the same secondary structure composition\(^\text{29}\). It is therefore likely that Zn\(^{2+}\) binding exerts more subtle effects on SynZur structural dynamics. Indeed, small differences in the conformations of the two dimers (Extended Data Fig. 8) point to a degree of conformational flexibility—even in the presence of zinc and in the crystal.

With SynZur now firmly established as a zinc sensor, we next explored its regulon in WH8102 by transcriptomic analysis.

**Zinc and SynZur regulate genes for zinc uptake and storage.** To study SynZur-dependent transcription, mutant and WT cells were grown in chelexed ASW medium\(^\text{27}\), to which 0 or 772 nM Zn\(^{2+}\) had been added. Cells were harvested in mid-exponential phase (optical density at 750 nm (OD\(_{750}\)) of \(\approx 0.3–0.4\)) and subjected to RNA-seq.

The most substantial changes in SynZur-dependent transcription occurred when comparing the WT and mutant at abundant zinc (772 nM; Fig. 4a and Extended Data Fig. 9a). Here, synw0971 (putative znuA) was the most upregulated gene in the mutant. The fact that removal of SynZur increases expression of znuA is consistent with the canonical mode of action of Zur sensors, namely repression of transcription when intracellular zinc is abundant enough to bind to Zur, which in turn enhances its DNA-binding affinity\(^\text{29}\). In fact, the entire gene cluster synw0968–synw0973, including znuB (synw0969; encoding the permease component of the ABC transporter) and znuC (synw0970; encoding the ATPase component of the ABC transporter), was upregulated, suggesting that all six of these genes are repressed by SynZur. synw0972 encodes an uncharacterized protein and is likely co-transcribed with synw0971. synw0973 and synw0968 are also annotated as uncharacterized proteins; how these are regulated by Zur is unclear. Surprisingly, synw0359, the bacterial metallothionein (bmtA), and its neighboring gene synw0360 (‘weak similarity to phage integrase’) were both downregulated in the mutant. This suggests that these two genes are not repressed but activated by SynZur; whether this activation requires zinc-bound SynZur is explored later.

The analogous comparison at 0 nM added zinc (Extended Data Fig. 9b,c) highlighted the same eight genes, although the...
fold-changes were smaller in each case. The apparent upregulation of zur (synw2401) itself in the mutant is likely a consequence of the single crossover nature of the mutant constructed. RNA-seq transcripts map only to the first 315 bp of the synw2401 gene; thus translation of this incomplete transcript would not result in a functional protein. Applying the criteria of log₂(fold change) > 2 and P < 0.05, no other genes were differentially regulated at both 0 and 772 nM Zn between the WT and zur mutant.

Comparison of the datasets at 0 and 772 nM added Zn for the WT informs about which genes are regulated by zinc availability (Extended Data Fig. 9d,e). A relatively small number of genes were differentially regulated by more than fourfold (log₂(fold change) > 2) between these two conditions. The most upregulated gene at 0 nM added Zn was again synw0971, with the adjacent synw0972 also upregulated. The upregulation of synw0971 in response to zinc availability further confirms that this periplasmic binding protein and its associated ABC-system components (Extended Data Fig. 10) correspond to ZnuABC, and that this system deals with zinc uptake when zinc is scarce. In turn, the two most downregulated genes at low zinc were synw0359 (bmtA) and its neighbor synw0360, the same two genes found to be most downregulated in the zur mutant. This means that bmtA transcript levels increase at higher [Zn], suggesting that activation of transcription requires zinc-loaded SynZur. It also implies that the BmtA protein sequesters excess zinc at higher concentrations.

By contrast, synw2401 transcript levels were not significantly altered at different zinc concentrations in the WT (P > 0.80), so synw2401 transcription is not zinc-dependent, a common observation for other Zur sensors. In accordance with neither Zur- nor zinc-regulation, no binding of SynZur to the synw2401 promoter region was apparent either (Supplementary Fig. 6). This supports the suggestion that the apparent partial overexpression of synw2401 in the mutant is a consequence of its single crossover nature.

The modulus of log₂(fold change) for differentially expressed genes decreases in the order mutant/WT at 772 nM zinc > mutant/WT at 0 nM zinc > WT at 0/WT at 772 nM zinc. For example, for znuA, log₂(fold change) values were 9.00, 5.76 and 3.99, respectively. To capture any genes that might be regulated simultaneously by zinc and SynZur, but in a less-pronounced way than specified by the log₂(fold change) > 2 criterion, we considered all transcript level changes that fulfilled the P < 0.05 criterion for the three comparisons discussed so far (Extended Data Fig. 9f,g). The only two genes that were downregulated in both the absence of SynZur (irrespective of zinc supply) and at low zinc in the WT are bmtA and synw0360, and the only four upregulated genes are synw0970–synw0973, that is zurC, znuA and two genes encoding proteins of unknown function. ZnuB (synw0969) was upregulated 1.4-fold in the WT at 0 zinc compared with 772 nM zinc, but with very low significance (P = 0.83). This is also the case for the adjacent gene synw0968. It is likely that divergent transcription of zurC and znuA is regulated by a single Zur box (Figs. 1a and 4b). Potential RNA polymerase-binding sites identified are shown in Fig. 4b, confirming that for both znuA and zurC, the Zur box overlaps the −10 promoter elements, consistent with repression occurring through blocking RNA polymerase binding. ZnuA expression appears to be more sensitive than that of znuB or zurC (Extended Data Fig. 10).

The corresponding analysis for the bmtA promoter (Fig. 4c) indicates that the first Zur box partially overlaps both −10 and −35 elements, whereas the second box lies beyond the −35 element. Ferguson analysis of SynZur binding to the PbmtA promoter (Supplementary Fig. 7) indicated that at low [SynZur], only a...
single dimer bound, whereas at higher [SynZur], a maximum of two dimers were bound. It is unclear whether an equilibrium involving the binding of one or two dimers relates to the activation mechanism. Analysis of promoter regions of Zur-regulated genes in a range of bacteria shows that there is no discernible correlation between the presence of two Zur boxes and activation (Supplementary Table 4). Neither do all bmtA promoters from marine cyanobacteria contain two Zur boxes (Supplementary Tables 5 and 6). Our Ferguson analysis also provided no evidence for oligomerization; the latter has been observed for Zur-activated genes in *S. coelicolor*[^42] and *Xanthomonas campestris*[^43]. Other possibilities for activation described for iron-responsive Fur proteins include regulation via small RNAs[^18] and via reversing H-NS silencing, as seen for ferritin expression in *E. coli*[^44]. However, we were unable to find evidence for Zur/zinc regulated sRNAs or H-NS binding sites within the *PbmtA* promoter. Therefore, the mechanism of activation of *bmtA* expression by SynZur does not appear to follow any precedents. The implications of Zur-activated *bmtA* expression in response to elevated zinc availability are explored in the following section.

**Zur activation of bmtA enables safe accumulation of zinc.**

Expression patterns for *znuA* and *bmtA* were further studied by quantitative polymerase chain reaction with reverse transcription (RT–qPCR; Fig. 4d,e). These data are broadly in line with the trends observed in the RNA-seq data; maximal expression of *znuA* was observed in the mutant, irrespective of zinc concentration, followed by lower expression in the WT at 0 Zn, and very low expression at 772 nM or 2.5 μM Zn. The latter two expression levels are indistinguishable, indicating that repression is already maximal at 772 nM Zn.

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[^18]: Reference to small RNAs
[^42]: Reference to *S. coelicolor*
[^43]: Reference to *Xanthomonas campestris*
[^1]: Reference to *E. coli*
The pattern for \( \text{bmtA} \) is essentially a mirror image of that for \( \text{znuA} \), but basal expression (at low \([\text{Zn}]\) or in the mutant) was higher than \( \text{znuA} \) expression at high \([\text{Zn}]\). At 772 nM Zn, \( \text{bmtA} \) transcripts were 125 times more abundant in the WT compared with the mutant (Fig. 4e). Even at 0 added Zn, the WT expressed seven times more \( \text{bmtA} \) than the mutant. In the WT, transcript levels at elevated Zn (772 nM or 2.5 \( \mu \)M) were higher by a factor of 16–17 compared with no added Zn. These data confirm that although some basal expression occurs in the mutant, Zur is required to activate \( \text{bmtA} \) transcription in the presence of zinc.

EMSA experiments in dependence of Zn\(^{2+}\) availability confirm that for both \( \text{znuA} \) and \( \text{bmtA} \), Zn\(^{2+}\) is required for DNA-binding (Fig. 4f,g). The two promoters respond at slightly different free Zn\(^{2+}\) concentrations. This means that the downregulation of \( \text{znuABC} \) occurs at lower \([\text{Zn}]_{\text{free}}\) than the upregulation of \( \text{bmtA} \). Similar observations have been made for other Zur proteins\(^{15,16}\). It can also be suggested that the narrow range defined by the two \( K_D \) values (1.8–7.0 femtomolar) corresponds to the optimal intracellular \([\text{Zn}]\) for \( \text{Synechococcus} \) sp. WH8102.

Crucially, \( \text{bmtA} \) upregulation at higher \([\text{Zn}]\) offers an obvious explanation (Fig. 5) as to why the WT was able to accumulate much more zinc than the mutant at 772 nM while suffering no growth impairment: it can be expected that each additional BmtA protein molecule will be able to sequester up to four zinc ions\(^{45}\). Overall, this keeps the concentration of intracellular free Zn\(^{2+}\) in a safe range and allows for a 43-fold increase (Figs. 1e and 5) in the total cellular zinc quota between the 0 and 772 nM added zinc conditions in the WT.

Although these data indirectly confirm that BmtA in WH8102 has a role in dealing with zinc ‘luxury’ and excess, the observation of appreciable basal transcription of \( \text{bmtA} \) in both the mutant and the
WT at 0 zinc (Fig. 4c) may indicate a more fundamental role for the BmtA protein, which may include redox buffering or zinc donation to other proteins. Indeed, previous proteomic work investigating the response of WH8102 to phosphorus and zinc scarcity showed that the abundance of BmtA followed similar trends to those of a putative alkaline phosphatase (SYNW2391), leading to the suggestion that BmtA might supply zinc to this enzyme. An analysis of the distribution of bmtA genes in cyanobacteria (Supplementary Table 5) may lend support to this hypothesis: bmtA genes are widespread in marine Synechococcus strains, with the majority of strains from clade III containing bmtA genes with two Zur boxes. The latter strains are dominant in warm oligotrophic waters that are permanently depleted in phosphorus.

Discussion

SynZur (SYNW2401) is a metallo sensor of the Fur family that responds to zinc, and hence a confirmed Zur protein. This is evidenced by: (1) impaired zinc tolerance and altered zinc accumulation in the zur mutant, (2) strong overlap between genes regulated by zinc and SYNW2401, (3) zinc-dependent DNA binding of the recombinantly expressed SynZur protein and (4) the presence of a sensory metal-binding site with a tetrahedral NOS coordination sphere that is typical for Zn2+. The SynZur crystal structure is distinct from previously characterized homologs in terms of domain orientation and location of the sensory zinc site. Given the high degree of conservation between SynZur and its predicted orthologs from both marine and freshwater cyanobacteria, this structure may also support further studies on any cyanobacterial Zur protein.

The Zur regulon of the marine cyanobacterium Synechococcus sp. WH8102 is small, comprising eight genes, six of which are repressed and two of which are activated by SynZur. Among the repressed genes are the three components of a znuABC Zn2+ uptake system (synw0969–synw0971). In contrast to repression of znuABC transcriptional activation by Zur proteins is rare, and Zur regulation of a bacterial metallothionein is unprecedented, as previously only transcriptional repression by SmtB-type zinc-sensor proteins has been reported. Thus, Zur in WH8102 regulates both zinc uptake (via znuABC) and storage (via bmtA) (Fig. 5).

Taken together, our data provide evidence for zinc being an essential element for a marine cyanobacterium. The low zinc quota for the WT at 0 added zinc, together with no evidence for the cultures being zinc-limited, suggests that the minimal zinc requirements of Synechococcus sp. WH8102 are very low, as may be expected for an oligotrophic strain. Yet, by expressing a bacterial metallothionein, WH8102 can deploy a considerable capacity for storage of surplus zinc — up to more than two orders of magnitude above these minimal levels (Fig. 5a). Similar ranges (24 to 1,138 zeptomoles per cell) have been found for zinc quotas in marine Synechococcus sampled from different types of mesoscale eddies in the Sargasso Sea, the original habitat of Synechococcus sp. WH8102. No other metal showed such a wide range. Indeed, such variations in cellular metal quotas are far from common: for example, metal quotas in E. coli cultured in different media including minimal and excess (0.1 mM) Zn2+ vary only two- to fourfold with respect to replete media.

A second putative znuABC system in this strain (synw2479–synw2481) was neither zinc- nor Zur-regulated. However, the periplasmic binding protein SYNW2481 was previously identified in the proteome of WH8102 cultured at 80 nM Zn2+, and our transcriptomic data indicate that all three components are expressed at appreciable levels in all conditions. This suggests that this system is constitutively expressed and could contribute to zinc uptake even when synw0969–synw0971 is completely repressed. The remarkable zinc accumulation at higher [Zn] may be facilitated either by this system and/or nonspecific transport through other metal transporters. It is also noteworthy that synw2479–synw2481 are upregulated under phosphorus depletion. Together with the finding of zinc-dependent abundance of an alkaline phosphatase at low [P] and the widespread distribution of Zur-regulated bmtA genes in clade III strains, this lends further support to the idea that zinc may be utilized for phosphorus acquisition from dissolved organic phosphates. Scavenging phosphorus from organic phosphates is a critical strategy for WH8102 and related strains being able to thrive in oligotrophic waters that are extremely scarce in phosphorus. Thus, the ability to avidly accumulate zinc when it becomes available may expand the ability of WH8102 and other oligotrophic strains that harbor bmtA genes to proliferate in these ‘ocean deserts.’

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41589-022-01051-1.

Received: 5 March 2021; Accepted: 5 May 2022;
Published online: 9 June 2022

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Methods

Bacterial strains and growth conditions. *Escherichia coli* cells were grown in liquid LB medium or on solid LB agar at 37 °C with kanamycin (Km, 50 µg/ml), ampicillin (Amp, 100 µg/ml) or chloramphenicol (Cm, 30 µg/ml) added where appropriate. Strains used are shown in Supplementary Table 7.

*Synechococcus* sp. WH8102 cells were cultured in 100 ml of ASW medium without added Zn (ASW−−Zn−). Supplementary Table 8) in 250 ml glass conical flasks. Cultures were maintained at 23 °C with continuous illumination (10 µE m−2 s−1 white light) and subcultured once a month by tenfold dilution into fresh ASW−−Zn− medium accompanied by checking for contamination. The mutant was maintained with 50 µg/ml Km.

Construction of a single crossover *Synechococcus* sp. WH8102 zur mutant. Genomic DNA extracted using a phenol–chloroform protocol61 was used as a PCR template. Vectors used in this study are shown in Supplementary Table 9. The zur−−ZurFABC−−zur gene was inserted into *ZurF* and *ZurABC* using *ZurF* and *ZurABC* (Supplementary Table 10) and MyTaq Read DNA Polymerase kit (Bioline). The insertion was ligated into pPG704CmKm vector at SalI and XbaI cloning sites and the mixture was transformed into *E. coli* strain S17-1 λPri. Conjugation was performed as described previously in the presence of sucrose-intolerant *Ruegeria pomeroyi* DSS-3 (pBBR MCSI Km r pKNG101)62. When colonies appeared, they were transferred into 1 ml of ASW−−Zn− containing 25 µg/ml Km and upon growth were gradually transferred into larger volumes of the medium with increasing concentrations of Km reaching 50 µg/ml. Single colony single crossover was assessed by colony PCR with primers *A_F* and *B_Re* or *C_Re* and *D_F* (Supplementary Table 10). Complete segregation was demonstrated using PCR with primers *A_F* and *C_Re*. Completely segregated mutant cultures were incubated overnight in ASW−−Zn− with 50 µg/ml Km, 100 µg/ml Amp and 10% (w/v) sucrose to remove *R. pomeroyi*. The mixture was then pour-plated using serial dilution to 0.22% (w/v) agarose overnight at 65 °C. Digests were diluted with 5.7 ml of MilliQ water to prepare the mid-log stage with 0.5 mM IPTG (ThermoFisher Scientific). Cells were lysed by sonication in Buffer 1 (50 mM NaH2PO4, 300 mM NaCl, 20 mM imidazole, pH 8.0). SynZur was purified using a Ni–Sepharose His-Trap column (GE Healthcare, 5 ml) using an AKTA purification system (GE Healthcare) with gradient elution with 200 mM NaH2PO4, 250 mM NaCl, 50 mM NiCl2, 100 mM imidazole, pH 6.8). The His-tag was cleaved by tobacco-etch-virus protease following buffer exchange into cleavage buffer (50 mM Tris–HCl pH 8.0, 1 mM DTT). Cleaved Zur was purified using the same His-Trap column. The purity of the protein was checked by SDS–PAGE in 14% Tris–glycine gel (Novex).

Growth rate comparison. Before adding the trace metal stock (Supplementary Table 8), the ASW macroeutrients solution was treated with Chelex 100 resin (Bio-Rad). *Synechococcus* sp. WH8102 WT and zur knockout mutant cultures were grown in chelched ASW−−Zn−, in triplicate until the late-log phase (OD750 >1). OD750 measurements were taken every 48–72 h until cultures reached the stationary phase. The initial OD750 and the experiments with different added zinc concentrations. Cultures were checked for contamination at each time point.

Specific growth rates were derived from the gradients of the linear portion of growth curves. Data are the mean of three biological replicates.

Trace metal analysis. The zur mutant and WT cells were grown in triplicate in chelced ASW−−Zn− with 0.772 mM and 2.5 µM of zinc added. For the zur knockout mutant, only the 0 and 772 µM zinc conditions were used. At the mid-log stage of growth (OD750 = 0.4–0.5), 50 ml of cells were harvested by centrifugation at 4000 r.p.m. for 30 min. Cells were then resuspended in 10 ml of MilliQ water to prepare the mid-log phase (OD750 >1). OD750 measurements were taken every 48–72 h until cultures reached the stationary phase. The growth rate of the experiments with different added zinc concentrations. Cultures were checked for contamination at each time point.

For RNA-seq analysis, HISAT2 (ref. 56) software was used to map FASTQ reads onto the genome. Resulting SAM files were converted to BAM and sorted BAM using Samtools63. FeatureCounts64 software was used to identify mapped genes. DESeq2 (ref. 65) as an R-package in R-studio software was used to normalize raw reads and calculate statistics.

For RT–qPCR analysis, reverse transcription was performed using the GoScript Reverse Transcription System (Promega). The RT–qPCR mixtures were prepared in 96-well Microamp microplates (Applied Biosystems) and covered with MicroAMP adhesive film (Applied Biosystems). PowerUp SYBR Green Master Mix (Applied Biosystems) was used to quantify amplification. All reactions had three technical replicates for each of the two biological replicates on a 7500 Fast Real-Time PCR System (Applied Biosystems). The presence of a single product was inspected by analysis of melting curves. Data were analyzed using 7500 software, v.2.3 (Applied Biosystems) and Microsoft Excel.

Primers for qPCR were designed using PrimerQuest Tool 150 from IDT60 and are given in Supplementary Table 10. The expression level of *synw2047* (synZur, phosphoenolpyruvate carboxylase) was used to normalize transcript abundance.

SynZur overexpression and purification. The sequence for *Synechococcus* sp. WH8102 Zur was codon-optimized for expression in *E. coli* and synthesized by GeneArt (Invitrogen) before cloning into a PET155-D-TOPO vector with an N-terminal His-tag (Invitrogen). SynZur was expressed in *E. coli* BL21(DE3)LYS8 (Invitrogen) grown in LB at 25 °C overnight following induction at the mid-log phase with 0.5 mM IPTG (ThermoFisher Scientific). Cells were lysed by sonication in Buffer 1 (50 mM NaH2PO4, 300 mM NaCl, 20 mM imidazole, pH 8.0). SynZur was purified using a Ni–Sepharose His-Trap column (GE Healthcare, 5 ml) using an AKTA purification system (GE Healthcare) with gradient elution with 200 mM NaH2PO4, 250 mM NaCl, 50 mM NiCl2, 100 mM imidazole, pH 6.8). The His-tag was cleaved by tobacco-etch-virus protease following buffer exchange into cleavage buffer (50 mM Tris–HCl pH 8.0, 1 mM DTT). Cleaved Zur was purified using the same His-Trap column. The purity of the protein was checked by SDS–PAGE in 14% Tris–glycine gel (Novex).

Protein characterization. Analytical SEC was carried out using an AKTA purifier equipped with a Superdex 200, 10/300 column (GE Healthcare) running a flow rate of 0.5 ml/min (50 mM Tris pH 8.0, 100 mM NaCl). The column was calibrated with Blue Dextran (2,000 kDa; to determine the void volume), BSA (66.4 kDa), carbonic anhydrase (29.2 kDa) and cytochrome c (12.2 kDa).

Dynamic light scattering was used to determine the hydrodynamic diameter of the protein. The protein was diluted to 20 µM with 50 mM Tris and filtered using a 0.2-µm pore-size filter (Sartorius Minisart RC4 syringe filter). The hydrodynamic diameter was measured at 25 °C using a Malvern Zetasizer Nano which was equilibrated for 300 s before each measurement. A total of six measurements were taken for each sample. Theoretical hydrodynamic diameters for nonmonomers, dimers and different tetrameric assemblies were calculated from the three-dimensional structure in this work, with the size of an ‘interwound’ tetramer based on the published structure for *Francisella tularensis* Fur (pdb 5nhk)67. For these calculations, radii of gyration (Rg) were calculated using WinHydroPro68 and converted to hydrodynamic radii (Rh) by employing the simple relationship Rg = Rh/0.774 (ref. 69). Correct calibration of the instrument and the validity of the approach to estimate the hydrodynamic sizes were checked using carbonic anhydrase (29.2 kDa) and cytochrome c (12.2 kDa) measured under the same conditions.

Nondenaturing SDS–PAGE was carried out as described in ref. 1. Protein samples were mixed with 4x sample buffer (100 mM Tris–HCl, 150 mM Tris base, 10% v/v glycerol, 0.1815% w/v Coomassie G-250, 0.00825% w/v Phenol Red, pH 8.5). Samples were loaded onto precast 10% Tris–glycine gels, and electrophoresis was performed in non-denaturing SDS buffer (50 mM Tris pH 7.3, 50 mM MOPS, 0.0375% w/v SDS) at 4 °C and 100 V until the dye front reached the bottom of the gel. Gels were visualized using SimplyBlue SafeStain (Life Technologies) and scanned.

Spectrophotometric determination of zinc affinity. Zinc affinity was determined following a well-established methodology suitable for metal sensors and is based on competition between apo-protein and the metallochroic dye Quin-2 (ref. 2). For removal of the sensory site Zn, SynZur at a concentration of 32 µM in 100 mM Tris–HCl, 50 mM NaCl, 10% v/v glycerol, 0.1815% w/v Coomassie G-250, 0.00825% w/v Phenol Red, pH 8.5). Samples were loaded onto precast 10% Tris–glycine gels, and electrophoresis was performed in non-denaturing SDS buffer (50 mM Tris pH 7.3, 50 mM MOPS, 0.0375% w/v SDS) at 4 °C and 100 V until the dye front reached the bottom of the gel. Gels were visualized using SimplyBlue SafeStain (Life Technologies) and scanned.
coefficient of 37.500 cm⁻¹ M⁻¹ (ref. 34). Protein concentration was estimated by absorbance at 280 nm, using an extinction coefficient of 3,485 cm⁻¹ M⁻¹.

The latter was determined by accurately measuring protein concentration through sulfur quantitation by inductively coupled plasma optical emission spectroscopy (ICP-OES), and is close to the theoretical value (3,400 cm⁻¹ M⁻¹). The zinc concentration in stock solutions and final samples was also determined by ICP-OES. A UV–visual spectrum was measured after each addition of ZnSO₄ with 20% glycerol (pH 8), chelexed before use. Mixtures were equilibrated for 30 min, loaded onto pre-run Novex WedgeWell gel, and incubated at 4 °C. Small rod-shaped crystals appeared after 1 week, grown by ICP-OES. A UV–visual spectrum was measured after each addition of ZnSO₄ to the atomic model automatically using ARPA®, at the positions of large positive peaks in the difference electron density map, only at places where the resulting water molecule fell into an appropriate hydrogen-bonding environment. Restricted isotropic temperature factor refinements were carried out for each individual atom. The polypeptide chain was traced continuously through electron density maps (2Fo–Fo and Fo–Fo) from residues 6–104 and 108–128 for chains A, B and D, and residues 6–102 and 108–128 for chain C, respectively. Data collection and refinement statistics are given in Supplementary Table 2.

Promoter analyses. The 150 bp promoter regions of marine cyanobacterial metallothionein genes were extracted manually from CyanoBak®. Putative cyanobacterial Zur-binding box was inferred from RegPrecise® (NT/NTAGTA TA/TTAN). Scanning across cyanobacterial metallothionein promoters was performed using FIMO (part of the MEME suite) with default parameters®. Bacterial genes with predicted double Zur boxes were extracted from RegPrecise. The ~10–35 elements were identified by Softberry BPR®

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. E. coli strains (Supplementary Table 7), plasmids (Supplementary Table 9) and oligonucleotides (Supplementary Table 10) are provided in the Supplementary Information. The atomic coordinates and structure factors for Synecococcus sp. WH1012 Zur have been deposited in the Protein Data Bank under the accession number 7NE9. RNA-sequencing data have been deposited in the EMBL-EBI database (accession number F-MTAB-10194). RNA-seq FASTQ files are available at https://cris.liv.ac.uk/illum/LIMS16562_259a058713 a41db/. Source data are provided with this paper. Other data supporting our findings are available in extended data and supplementary information. Any raw biophysical and RT–qPCR and construct DNA sequencing data are available from the corresponding author c.blindauer@warwick.ac.uk.

Code availability. R-Studio scripts used for RNA-seq analysis are available at https://github.com/AlevtinaMikh/Syn1012.

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Acknowledgements
This work was supported by the University of Warwick through a Chancellor’s International scholarship to A.M. We would like to thank Diamond Light Source, UK for beamtime (proposal MX14692), and the staff scientist K. McAuley of beamline I03 for assistance with crystal testing and data collection. We also thank L. Song for excellent assistance in inorganic and molecular mass spectrometry and R. Puxty for useful discussions on our RNA-seq data. This work was supported by the Biotechnology and Biological Sciences Research Council (grant reference BB/M003523/1) and the Natural Environment Research Council (grant reference NE/I00985X/1). Some equipment used in this research was obtained through Birmingham Science City with support from Advantage West Midlands and the European Regional Development Fund.

Author contributions
C.A.B. and D.J.S. conceived the project. A.Z.K. participated in the experimental design, constructed the wild-type Zur overexpression plasmid and conducted initial Zur expression, characterization and EMSA trials. A.M. optimized Zur expression, characterization and EMSA, designed, oversaw, carried out and analyzed all other experiments except for X-ray structure analysis and ICP-MS data acquisition. R.C.W. and V.F. carried out all aspects of X-ray structure analysis, including SEC purification, crystal screening and optimisation of crystallization conditions (R.C.W.) and structure calculations (V.F. and R.C.W.). J.P.C.C. undertook ICP-MS data acquisition. E.M. and D.S. assisted in protein expression, EMSA, SEC and dynamic light scattering data acquisition and analysis. A.M. and C.A.B. wrote the manuscript with input as appropriate from all other authors.

Competing interests
The authors declare no competing interests.

Additional information
Extended data is available for this paper at https://doi.org/10.1038/s41589-022-01051-1.

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41589-022-01051-1.

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Peer review information *Nature Chemical Biology* thanks the anonymous reviewers for their contribution to the peer review of this work.

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Extended Data Fig. 1 | See next page for caption.
Extended Data Fig. 1 | Construction, confirmation and metal accumulation of a synw2401 disruption mutant. a, Construction of a single crossover Synechococcus sp. WH8102 synw2401 interposon mutant. Primers used to confirm the successful mutation are indicated with letters (also see Supplementary Table 10). b, Agarose gel electrophoresis of PCR products obtained with these primers. This analysis was repeated multiple times (n > 3), always successfully. c, Cellular metal quotas, expressed as mmol metal per mol phosphorus for mutant and WT grown at different zinc concentrations.
Extended Data Fig. 2 | Residues critical for zinc sensing identified in other Zur proteins are not conserved in cyanobacterial homologs. Multiple sequence alignment comparing three predicted Fur-family proteins from *Synechococcus* sp. WH8102 with selected homologs for which functional and/or structural information is available. Confirmed or predicted metal-binding residues are highlighted in grey (structural ZnCys4 site 1), red (canonical major sensory site 2; absent in cyanobacterial Zurs), and yellow (site 3). The residues highlighted in cyan form important inter-domain or inter-subunit hydrogen bonds or salt bridges (see Fig. 3c and Extended Data Fig. 8). The other two WH8102 Fur family proteins can be predicted to possess a canonical site 2 capable of binding Fe$^{2+}$ or Mn$^{2+}$. Pdb entries used for metal site analysis: 2o03 (*Mycobacterium tuberculosis* FurB (Zur))$^{34}$; 3mwm (*Streptomyces coelicolor* Zur)$^{39}$; 4mtd (*Escherichia coli* Zur)$^{33}$; 7dh8 (*Xanthomonas campestris* Zur; sequence lacks disordered N-terminus)$^{35}$; 2xig (*Helicobacter pylori* Fur)$^{41}$; 5nhk (*Francisella tularensis* Fur)$^{51}$; 4raz (*Magnetospirillum gryphiswaldense* Fur)$^{40}$; 3f8n (*Bacillus subtilis* PerR; Jacquamet, L. et al. Structural characterization of the active form of PerR: insights into the metal-induced activation of PerR and Fur proteins for DNA binding. *Mol Microbiol* 73, 20–31 (2009)).

| Syn. WH8102 Zur | ----- | MTGSPALNAR---- | QALLTNAGC---- | DQMSQQLRSLID-- | DEASMG |
| Anabaena Zur | ----- | MRAIR----- | TRGQERWLNLQTIK | QGISAQIDYVELRNRNQSMG |
| Synechocystis Zur | ----- | MLPLTPAVLRELSTLVNQRLVQALQRET | EPLSQAQLFAKLR- | ETRKKG |
| Mycobacterium Zur | ----- | MASAAG---- | VSTR----- | OORLASTLETLD- | DPRSAQELHDLERRGENI |
| Streptomyces Zur | ----- | VTTAGPVPKGRAT----- | QRAVSAALQEVE-E | EFRSAQELHDMLKHKGDAVG |
| E. coli Zur | ----- | MEKTQTELQAEKCIAQTNVLTPQELVRLIMSLQD- | GAIASYDLDDLIR-LAE-P | PerR |
| Xanthomonas Zur | ----- | DDDNFQVRAVERACSERGLRLTPIRANVLRLEDAGA----- | KQPVAYELLDDVR- | ESKGQV |
| Helicobacter Fur | MKRLLETSLERLMLRSIKKGLKNSKQREWEEVSVELWRSSG- | TLSPEIRTHSIRQDKNTS |
| Francisella Fur | ----- | MNSKNLKDFTGFQTRGLVPPVEILKLFEKNKD- | KPSLFDPVFSLKAGSTTG |
| Magneto spirillum Fur | ----- | GMVRSTQICDRICDDKMTDQRVIAQVLSDSA----- | DPDBVEVYRRTAKDDPRIS |
| Syn. WH8102 Fur | MSLPSFAPADGLQQGPHQGDRLTPQKRLELFERGCGLSLAEVHQIALQSLKQV |
| B. subtilis PerR | ----- | MAHAHELKELETKLKETGVTPQHRAILEYVNSM- | APTADDIYKALEGKFPNMS |
| Syn. WH8102 PerR | ----- | MFRCQRLGMRLSRQMRRMLDLWSEQ----- | SLDIADFEKLNARSGIG |

| Syn. WH8102 Zur | ----- | IATVYRNLRQLQRLVRCRLPTEALAPVD----- | ALHKLHLYGQTQVLDIHCPH |
| Anabaena Zur | ----- | IATVYRALDEALKGELEQTVLPNEALYSIAQ----- | ODYHLTHIQLQGVSIPIHCPVH |
| Synechocystis Zur | ----- | IATVYRALDKLFQI0HQAAMTGELYYLTE----- | ODYHLTHIQLQGVSIPIHCPVH |
| Mycobacterium Zur | ----- | IATVYRLGSMAAVLVQPIEHSVYRRT----- | EHHDLRSGTSCSTIEGDHEV |
| Streptomyces Zur | ----- | IATVYRLSLADAVEQSLVEASGYSR----- | STD-GEHELVRAGKAVEVEPAG |
| E. coli Zur | ----- | AKPPTVYRLDLFELQEQAVKESNYSVL1FLQDFNP-IATSMICIDRCQGAVKKEACAGV |
| Xanthomonas Zur | ----- | ADAPTPVYRLDLFELQEQAVKESNYSVL1FLQDFNP-IATSMICIDRCQGAVKKEACAGV |
| Helicobacter Fur | ----- | ISSYVRILNFEKLEPNISVLTSEKSSRRT----- | IAEE-EBHHMLGCHKIKT-FADPBE |
| Francisella Fur | ----- | IATVYRVLNQFSAIINRLKLDNEQVMY----- | LBG-EHDLIIYKCNMIOFYSFSP |
| Magneto spirillum Fur | ----- | IATVYRTVRLEFESILERLHDPCGDGRY----- | EAP-EHDLIDDNARSVFTSPTE |
| Syn. WH8102 Fur | ----- | IATVYRTLALLDMPFQFLQELLELSGRR---FLDAGHDR-HELHVRAGKCEFTEFSEPEVL |
| B. subtilis PerR | ----- | VATVVNLVRVFRELSVLKELTYGDASRRPVTS----- | D-YAIENICGKVIFVHPGDL |
| Syn. WH8102 PerR | ----- | HTSAQNYLIALQSAVIGECRDLANG-RLYGCR----- | DPSL-LDQELTVCEIDVELPE |

| Syn. WH8102 Zur | GIDVPADSRGDFELLF1LLEFFGEESSRPRQSSK----- |
| Anabaena Zur | NLEEQLQTAKHEKFIYRTLEFFGCLGKCOMHASEI- |
| Synechocystis Zur | SLEENLQANYFRIYYTLEFFGCLGKCAK- | GSD- |
| Mycobacterium Zur | AWAAYAVTKHGGDSVTVETIEFTGCDSDSR |
| Streptomyces Zur | KWAEEIAAEGYVNNVETEIFGTCDACAGASG |
| E. coli Zur | DIMHTLAKMFGALRNVIHELAAHVCEACRHPEQCQHDSVQKP |
| Xanthomonas Zur | SQILEARAKALGFQPAQTLVHELCAKAAAG |
| Helicobacter Fur | NQRNEVKKYQAKLISHDMKMFVWCEQSES |
| Francisella Fur | ALQKQIVESFAGMDLYSLNYKCSREKI |
| Magneto spirillum Fur | ALQREIKHGFRLVGRLELYGVLTSGGSDDD |
| Syn. WH8102 Fur | MAGADAAQQFGKLISSLNVALENCPNLQ |
| B. subtilis PerR | EVEQLAHTVGTKVSHIRLEIYVQGIRCSEKHEL |
| Syn. WH8102 PerR | DLDLDTQKRTFRIESTYLQNLNRGRPLER |

Extended Data Fig. 2 | Residues critical for zinc sensing identified in other Zur proteins are not conserved in cyanobacterial homologs. Multiple sequence alignment comparing three predicted Fur-family proteins from *Synechococcus* sp. WH8102 with selected homologs for which functional and/or structural information is available. Confirmed or predicted metal-binding residues are highlighted in grey (structural ZnCys4 site 1), red (canonical major sensory site 2; absent in cyanobacterial Zurs), and yellow (site 3). The residues highlighted in cyan form important inter-domain or inter-subunit hydrogen bonds or salt bridges (see Fig. 3c and Extended Data Fig. 8). The other two WH8102 Fur family proteins can be predicted to possess a canonical site 2 capable of binding Fe$^{2+}$ or Mn$^{2+}$. Pdb entries used for metal site analysis: 2o03 (*Mycobacterium tuberculosis* FurB (Zur))$^{34}$; 3mwm (*Streptomyces coelicolor* Zur)$^{33}$; 4mtd (*Escherichia coli* Zur)$^{33}$; 7dh8 (*Xanthomonas campestris* Zur; sequence lacks disordered N-terminus)$^{35}$; 2xig (*Helicobacter pylori* Fur)$^{40}$; Snhk (*Francisella tularensis* Fur)$^{35}$; 4raz (*Magneto spirillum gryphiswaldense* Fur)$^{39}$; 3f8n (*Bacillus subtilis* PerR; Jacquamet, L. et al. Structural characterization of the active form of PerR: insights into the metal-induced activation of PerR proteins for DNA binding. *Mol Microbiol* 73, 20–31 (2009)).
Extended Data Fig. 3 | See next page for caption.
Extended Data Fig. 3 | Purification and characterization of SynZur from Synechococcus sp. WH8102. a, Two-step IMAC purification of SynZur, including SDS-PAGE of fractions. Purification was repeated multiple times (n ≈ 50), always yielding the same product. b, The full ESI-MS spectrum of purified apo-Zur at pH 2. The deconvoluted spectrum is shown in the top right-hand corner. The pH of 5 μM SynZur in 20 mM NH₄HCO₃ was adjusted to 2.0 with formic acid. The theoretical neutral mass of apo-SynZur, including 6 additional residues from the cleaved tag, is 15466.43 Da. c, Analytical SEC of Zn₅SynZur (40 μM) in 20 mM Tris-HCl (pH 8) and 300 mM NaCl at 0.5 mL min⁻¹. Standards at concentration 1 mg mL⁻¹ used were: Blue Dextran, 2000 kDa; BSA, 66.4 kDa; Carbonic anhydrase, 29 kDa; Cytochrome c, 12.2 kDa. d, Native SDS protein gel electrophoresis shows four bands, with the strongest band likely to correspond to a dimer. The numbers 1-4 refer to likely oligomeric states. A repeat of the experiment showed identical results.
Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | Oligomeric states in the crystal. a, The two dimers in the asymmetric unit of SynZur. There are no physiologically relevant contacts between these two dimers. b, The symmetry-related zinc site formed by chains B (green) and D (blue) of different crystallographic units. In the zoomed-in inset, the electron density map (σ level 1.5) is displayed as a mesh. The origin of this ‘surplus’ zinc ion (0.5/dimer) is unclear, but it is most likely that its presence is related to crystal packing. This assessment is based on the observation that this region is followed by a short stretch (residues 105–107 in chains A, B and D, 103–107 in chain C) with unresolved electron density. Based on structural comparisons and secondary structure predictions, residues 97–106 are expected to form an α-helix, but this is absent in the SynZur X-ray structure. We suggest that the conformational changes imposed by binding the inter-dimer zinc have caused structural disorder in this region. The non-conservation of H94 in Zur proteins from marine or freshwater cyanobacteria (Supplementary Figure 4) also argues against this zinc site relating to a physiological process, although we cannot exclude that Zur regulation in WH8102 may differ from that in other marine cyanobacteria.
Extended Data Fig. 5 | A Cys95Ala mutant loses the ability to bind to the znuA promoter. a, Deconvoluted ESI-MS spectrum under denaturing conditions (pH 2), confirming the expected mass for the C95A mutant. b, Deconvoluted ESI-MS spectrum under near-native conditions. The mass is consistent with that expected for the mutant protein with one zinc ion bound. Dimers are still observed in the gas phase. c, Analytical size-exclusion chromatogram of Cys95Ala SynZur (30 µM in 20 mM Tris-HCl (pH 8), 300 mM NaCl; Superdex G200, 10/300, GE Healthcare; 0.5 mL min⁻¹). The observed mass is somewhat smaller than that for the wild-type (48.3 kDa), for which dominance of the dimer has been confirmed by dynamic light scattering analysis (Supplementary Fig. 2). We infer that the C95A mutant also still forms dimers in the solution state. d, The C95A mutant has no meaningful binding ability to the znuA promoter, similar to EDTA-treated wild-type SynZur (Fig. 2e). In contrast, wild-type SynZur as purified showed 100% binding at 100 nM.
Extended Data Fig. 6 | Comparison of locations of metal-binding sites on Zur and Fur proteins. The boxes in the top left comparison are coloured to match Extended Data Fig. 2. The canonical sensing sites in other Zur and Fur proteins (grey ribbons and metal ions) is labelled ‘Site 2’, and its inter-domain location is thought to be essential to stabilise a ‘closed’ conformation with high affinity for DNA. The absence of this site in SynZur (blue ribbons and purple zinc ions) is evident. The location of the new sensory site in SynZur is similar to that of site 3 in S. coelicolor or M. tuberculosis Zur. The ligand sphere of site 3 in these proteins consists of 3 His and 1 Glu residue. In contrast, the composition of the single sensory site in SynZur (AspHis2Cys) resembles that of the primary zinc-sensing sites in all other characterised Zur proteins (2 His, 1 Cys and one carboxylate). All proteins shown also harbour the structural Cys₄ site 1. For the superpositions shown, SynZur chain C (blue) was superposed with other Fur-family proteins (chain A in each case) by first matching the four Cys residues forming the structural site 1, and then improving the fits further using the respective tool in the Swiss pdb viewer v. 4.1. As can be seen, this gives a good match for the β strands in the dimerisation domain (typically around 1.2 Å RMSD for backbone heavy atoms).
a

Optimized for alignment of DNA-binding domains

b

| Pdb code | organism          | Protein name (condition) | RMSD (Å) (number of bb atoms aligned) |
|----------|-------------------|--------------------------|-------------------------------------|
|          |                   |                          | Full-length dimer | 2 DDs in dimer | 2 DDs in dimer | Single DBD |
| 3MWWM    | Streptomyces coelicolor | Zur (holo)               | 7.08 (924) | 1.40 (224) | 1.60 (484) |
| 4MTD     | E. coli           | Zur (holo+DNA)           | 7.15 (476) | 1.49 (264) | 1.87 (408) |
| 2003     | Mycobacterium tuberculosis | Zur (partially loaded) | 17.25 (936) | 1.47 (212) | n.a.         | 1.31 (260) |
| 7DH8     | Xanthomonas campestris | Zur (holo)               | 12.90 (268) | 1.65 (292) | n.a.         | 1.41 (276) |
| 1M2B     | Pseudomonas aeruginosa | Fur (holo-Zn)           | 6.53 (880) | 1.61 (240) | n.a.         | 1.19 (284) |
| 2XIG     | Helicobacter pylori  | Fur (holo-Zn)           | 13.17 (728) | 1.80 (308) | n.a.         | 1.25 (300) |
| 4RAY     | Magnetospirillum graftiswaldense | Fur (apo)       | 11.84 (916) | 1.77 (232) | n.a.         | 1.29 (244) |
| 4RAZ     | Magnetospirillum graftiswaldense | Fur (holo-Mn) | 7.99 (916) | 1.63 (272) | n.a.         | 1.53 (296) |
| 4RB2     | Magnetospirillum graftiswaldense | Fur (holo-Mn-DNA) | 6.47 (900) | 1.14 (140) | n.a.         | 1.38 (288) |
| 5NHK     | Francisella tularensis | Fur (holo-Fe)           | 7.32 (868) | 1.51 (220) | n.a.         | 1.29 (288) |
| 3F8N     | Bacillus subtilis | PerR (holo-Mn)          | 7.47 (936) | 1.34 (300) | n.a.         | 1.52 (284) |

Extended Data Fig. 7 | See next page for caption.
Extended Data Fig. 7 | The overall structure of SynZur differs from other Fur-family proteins. a, Superposition of the SynZur C-D dimer (blue) with the S. coelicolor dimer (grey; the dimer was generated from pdb 3mwm using the PISA server). The top superpositions demonstrate that the two DBDs are in similar positions in the two dimers (121 residues across two DBDs aligned with RMSD = 1.50 Å), but that the dimerization domains are in rather different orientations. The bottom superposition shows that the dimerization domains can be aligned well (RMSD = 1.40 Å over 56 residues), but that the DBDs are in different locations relative to the dimerization domains. Superpositions were performed using the ‘explore domain alternate fit’ tool in Swiss pdb viewer v. 4.1. b, Structural alignment/superpositions of SynZur (dimer 2, chains C and D) with other Fur family proteins. Single domain comparisons refer to chain C for SynZur and chain A for each of the other proteins in each case. ‘Apo’ and ‘holo’ refer to occupation of sensory sites, irrespective of the presence of the ZnCys₄ site. RMSDs for full-length dimers refer to values obtained using simple ‘magic fit’, whilst those for individual domains were obtained using the ‘explore domain alternate fit’ tool, both in Swiss pdb viewer v. 4.1. Footnotes: a) Functional dimers were generated from the original pdb files using the PISA server at https://www.ebi.ac.uk/pdbe/pisa/. Where more than two chains were present in the original pdb file, the chain identifiers are given. b) ‘n.a.’ = not applicable; this refers to cases where no superpositions were found that encompassed both DBDs.
Extended Data Fig. 8 | Subtle differences in structure and weak interactions between the two dimers in the crystal. **a**, Superposition of dimer 1 (chains A and B) with dimer 2 (chains C and D), where alignment of chains B and D has been optimized (carried out in Swiss pdb viewer v. 4.1). In dimer 2 (chains C and D), the two DBDs are ca. 3 Å closer together than in dimer 1 (chains A and B). This is accompanied by changes to the dimer interface and subtle variations in inter-protomer hydrogen-bonds as shown in (b). For dimer 1, the interface area is with 1668.3 Å² slightly smaller than that for dimer 2 (1703.1 Å²). **b**, These differences seem to be driven by conformational dissimilarities between monomers, as comparisons between individual monomers using the SuperPose webserv (http://superpose.wishartlab.com/) shows. Chain A differs the most from other chains. (RMSDs 1.08–1.23 Å over 120 Cα carbons; c, Residues engaged in inter-protomer hydrogen bonds and salt bridges (green lines), as derived from analysis using PISA (https://www.ebi.ac.uk/pdbe/pisa/). The residues highlighted in grey are involved in salt bridges. d, Particularly noteworthy are inter-subunit H-bonds between Gln55 and Asp85. These H-bonds between the DBD of one subunit and the DD of the other may confer enhanced stability to the SynZur dimer. Again, some variability is observed in inter-subunit H-bonding between Gln55 and Asp85. For chains B, C and D, the NH4 of Gln55 forms an H-bond with the backbone carbonyl oxygen of Asp85, whilst for chain C, an additional interaction with the carboxylate is also likely. For chain A, Gln55 does not undergo either of these interactions.
Extended Data Fig. 9 | See next page for caption.
Extended Data Fig. 9 | Transcriptomic analysis of mutant and wild-type *Synechococcus* sp. WH8102. Genes upregulated in the mutant (parts (a)-(c)) or at low [Zn] (parts (d) and (e)) are highlighted in red; genes downregulated in the mutant or at high [Zn] are highlighted in blue. a, Identity and annotations of genes that are differentially expressed in the mutant at 772 nM Zn (see Fig. 4a for Volcano plot). b, Identity and annotations of genes that are differentially expressed in the mutant at 0 Zn. c, Volcano plot depicting differentially expressed genes at 0 Zn. d, Identity and annotations of genes that are differentially expressed in the wild-type in dependence on [Zn]. e, Volcano plot depicting differentially expressed genes in the wild-type in dependence on [Zn]. In parts (a)-(e), only genes with log₂(fold change) > 2 and p-values < 0.05 are highlighted and listed. f, g, Venn diagrams including all up- and down regulated genes where P < 0.05, for the three comparisons referred to in Fig. 4a and parts (a)-(e) here. We observe that synw0969-0971 had been originally annotated with the addition ‘possibly Mn transport’. Noting that inference of metal specificity is non-trivial, we have therefore removed this incorrect specification in these Tables. Statistical analysis for parts (a), (b) and (d) involved default parameters for DeSeq2: Wald test for significance testing based on 3 biological replicates with two-sided P-values. Bonferroni cut-off was used for multiple comparisons adjustment.
Extended Data Fig. 10 | Expression levels of znuA, znuB and znuC from RNAseq data. The conditions (0, 772 nM, and 2.5 μM Zn²⁺ added to the culture medium) are the same as reported in Fig. 4d, e, and Extended Data Fig. 9. Transcripts (RPKM, reads per kilobase of transcript per million mapped reads) are normalized with respect to those of the pepC housekeeping gene. ZnuA and znuBC, being divergently transcribed, have different promoters but share a Zur box. Overall, trends are similar, but expression of znuA appears to be more sensitive to either Zn²⁺ or SynZur compared to those of znuB or znuC. Data are presented as mean ± standard deviation over 3 independent biological replicates for each condition.
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  *Give P values as exact values whenever suitable.*
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Software and code

Policy information about availability of computer code

**Data collection**

No custom code was used.

- X-ray data were collected at the zinc absorption edge (9666 eV) using beamline I03 and a Pilatus 6M detector at the Diamond Light Source, Didcot, Oxfordshire, UK.
- SDS-PAGE gels were scanned using a Ricoh Aficio MFD.
- UV-Vis data were acquired on a Cary 50 UV-Vis spectrophotometer.
- DLS data were acquired on a Malvern Zetasizer Nano instrument.
- CD spectroscopic data were acquired on a Jasco J-815 spectropolarimeter.
- The ICP-MS measurements were performed using an Agilent 7900 ICP-MS instrument. Data were acquired using Mass Hunter 4.3.
- RNA-Seq library preparation and sequencing were performed by the Centre for Genomic Research, Institute of Integrative Biology, at the University of Liverpool.
- RT-qPCR data were acquired on a 7500 Fast Real-Time PCR System (Applied Biosystems).
- DNA for EMSA experiments was visualized with SYBR-green (SIGMA-ALDRICH) using a luminescent image analyser ImageQuant LAS 4000 (GE Healthcare Bio-Sciences AB).

**Data analysis**

- Growth rate data were analyzed and plotted using R-studio Version 1.4.1717.
- The ICP-MS data were processed using Mass Hunter 4.3 and analyzed and plotted using R-studio Version 1.4.1717.
- Quin-2 competition experiments were analyzed using DynaFit 4.0.
- DLS data were analyzed using Malvern Instruments Zetasizer software and plotted in Excel. Theoretical radii of gyration (RG) were calculated using WinHydroPro and converted to hydrodynamic radii (RG) by employing the simple relationship RH = RG/0.774.
- CD spectroscopic data were plotted in Excel, and secondary structure was analysed using SELCON, CONTIN (accessed via DiChroWeb server) and RAUSSENS (webserver) algorithms.
- RT-qPCR data were analysed using 7500 Software, v2.3 (Applied Biosystems) and Microsoft Excel (v. 2202).
- EMSA gels for ferguson analysis were processed and analysed in Gimp (GNU Image Manipulation Program, v. 2.8.10).
- Bands from EMSA gels for assessing SynZur binding to PmuA and PbmIA were quantified using ImageJ, and data were fitted in Dynafit v. 4.0.
For RNA-Seq analysis Hisa2 software was used to map FASTQ reads onto the genome. Resulting SAM files were converted to BAM and sorted BAM using Samtools. FeatureCounts software was used to identify mapped genes. DESeq2 as a R-package in R-studio software was used to normalize raw reads and calculate statistics. R codes are available at https://github.com/AlevtinaMikh/Syn8102. X-ray data was processed using the XDS package and further handling took placing within the CCP4 software package. The crystal structure was solved SHELX. Refinement of the structure was carried out Coot and Refmac. Water molecules were added to the atomic model automatically using ARP.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

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E. coli strains (Supplementary Table 7), plasmids (Supplementary Table 9) and oligonucleotides (Supplementary Table 10) are provided in the Supplementary Information. The atomic coordinates and structure factors for Synechococcus sp. WH8102 Zur have been deposited in the Protein Data Bank under the accession number 7NE9. RNA-sequencing data have been deposited in the EMBL-EBI ArrayExpress database under the accession number E-MTAB-30194. RNA-seq FASTQ files are available at http://cgp.liv.ac.uk/illum/JM516056_259a058713a14129b/. Source data are provided with this paper. Other data supporting our findings are available in extended data and supplementary information. Any raw biophysical and RT-qPCR and construct DNA sequencing data are available from the corresponding author c.blindauer@warwick.ac.uk. All biological materials are available from the authors on request.

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Life sciences study design

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| Sample size | Experiments requiring statistical analysis were repeated as three independent biological replicates (RNA-seq, ICP-MS), as is common practice in the field. RT-qPCR data for each biological replicate were repeated at least 3 times, again as is common in the field. Electrophoretic Mobility Shift Assays were repeated at least twice, in keeping with common practice. The results of the Quin-2 competition experiments are from 3 independent titrations. Microbial growth experiments were performed with sample sizes of three replicates as is standard in microbiology. |
| Data exclusions | No data were excluded from the analysis. |
| Replication | All experiments requiring statistical analysis were reproducible and were repeated at least twice or more as repeats or biological replicates; details are provided in the figure captions. As is common in the field, and due to the nature of the work, X-ray crystallography and mutant construction, neither of which are subject to statistical analysis, were not repeated. However, the zur knockout mutant was repeatedly used in multiple experiments with three replicates (e.g. growth experiments, metal quotas, RNAseq) demonstrating reproducibility of their phenotypes. |
| Randomization | Randomization is not relevant to the current study, as for the experiments requiring statistical analysis, the order of analysis does not affect results. Thus, for example, randomization is not applicable to microbial growth experiments which involve clonal populations and tight control of abiotic factors other than those under investigation. As such, randomization, the purpose of which is to control for genetic and environmental variability, is unnecessary. Consequently, randomization is not part of the current standard of practice for defined microbial growth assays. |
| Blinding | Blinding is not relevant to the current study, as data evaluation for all experiments requiring statistical analysis is rule-based. |

Behavioural & social sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Study description | Briefly describe the study type including whether data are quantitative, qualitative, or mixed-methods (e.g. qualitative cross-sectional, quantitative experimental, mixed-methods case study). |
| Research sample | State the research sample (e.g. Harvard university undergraduates, villagers in rural India) and provide relevant demographic information (e.g. age, sex) and indicate whether the sample is representative. Provide a rationale for the study sample chosen. For studies involving existing datasets, please describe the dataset and source. |
Ecological, evolutionary & environmental sciences study design

All studies must disclose on these points even when the disclosure is negative.

**Study description**
Briefly describe the study. For quantitative data include treatment factors and interactions, design structure (e.g. factorial, nested, hierarchical), nature and number of experimental units and replicates.

**Research sample**
Describe the research sample (e.g. a group of tagged *Passer domesticus*, all *Stenocerus thurberi* within Organ Pipe Cactus National Monument), and provide a rationale for the sample choice. When relevant, describe the organism taxa, source, sex, age range and any manipulations. State what population the sample is meant to represent when applicable. For studies involving existing datasets, describe the data and its source.

**Sampling strategy**
Note the sampling procedure. Describe the statistical methods that were used to determine sample size OR if no sample size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient.

**Data collection**
Describe the data collection procedure, including who recorded the data and how.

**Timing and spatial scale**
Indicate the start and stop dates of data collection, noting the frequency and periodicity of sampling and providing a rationale for these choices. If there is a gap between collection periods, state the dates for each sample cohort. Specify the spatial scale from which the data are taken.

**Data exclusions**
If no data were excluded from the analyses, state so OR if data were excluded, describe the exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.

**Reproducibility**
Describe the measures taken to verify the reproducibility of experimental findings. For each experiment, note whether any attempts to repeat the experiment failed OR state that all attempts to repeat the experiment were successful.

**Randomization**
Describe how samples/organisms/participants were allocated into groups. If allocation was not random, describe how covariates were controlled. If this is not relevant to your study, explain why.

**Blinding**
Describe the extent of blinding used during data acquisition and analysis. If blinding was not possible, describe why OR explain why blinding was not relevant to your study.

**Field work, collection and transport**

**Field conditions**
Describe the study conditions for field work, providing relevant parameters (e.g. temperature, rainfall).

**Location**
State the location of the sampling or experiment, providing relevant parameters (e.g. latitude and longitude, elevation, water depth).

**Access & import/export**
Describe the efforts you have made to access habitats and to collect and import/export your samples in a responsible manner and in compliance with local, national and international laws, noting any permits that were obtained (give the name of the issuing authority, the date of issue, and any identifying information).

**Disturbance**
Describe any disturbance caused by the study and how it was minimized.
Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☒   | Antibodies            |
| ☒   | Eukaryotic cell lines |
| ☒   | Palaeontology and archaeology |
| ☒   | Animals and other organisms |
| ☒   | Human research participants |
| ☒   | Clinical data |
| ☒   | Dual use research of concern |

Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☒   | ChIP-seq |
| ☒   | Flow cytometry |
| ☒   | MRI-based neuroimaging |

Antibodies

- **Antibodies used**: Describe all antibodies used in the study; as applicable, provide supplier name, catalog number, clone name, and lot number.

- **Validation**: Describe the validation of each primary antibody for the species and application, noting any validation statements on the manufacturer’s website, relevant citations, antibody profiles in online databases, or data provided in the manuscript.

Eukaryotic cell lines

- **Policy information about cell lines**
  - **Cell line source(s)**: State the source of each cell line used.
  - **Authentication**: Describe the authentication procedures for each cell line used OR declare that none of the cell lines used were authenticated.
  - **Mycoplasma contamination**: Confirm that all cell lines tested negative for mycoplasma contamination OR describe the results of the testing for mycoplasma contamination OR declare that the cell lines were not tested for mycoplasma contamination.
  - **Commonly misidentified lines (See ICLAC register)**: Name any commonly misidentified cell lines used in the study and provide a rationale for their use.

Palaeontology and Archaeology

- **Specimen provenance**: Provide provenance information for specimens and describe permits that were obtained for the work (including the name of the issuing authority, the date of issue, and any identifying information).

- **Specimen deposition**: Indicate where the specimens have been deposited to permit free access by other researchers.

- **Dating methods**: If new dates are provided, describe how they were obtained (e.g., collection, storage, sample pretreatment and measurement), where they were obtained (i.e. lab name), the calibration program and the protocol for quality assurance OR state that no new dates are provided.

- **Tick this box to confirm that the raw and calibrated dates are available in the paper or in Supplementary Information.**

- **Ethics oversight**: Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Animals and other organisms

- **Policy information about studies involving animals**: ARRIVE guidelines recommended for reporting animal research

- **Laboratory animals**: For laboratory animals, report species, strain, sex and age OR state that the study did not involve laboratory animals.

- **Wild animals**: Provide details on animals observed in or captured in the field, report species, sex and age where possible. Describe how animals were caught and transported and what happened to captive animals after the study (if killed, explain why and describe method; if released, say where and when) OR state that the study did not involve wild animals.

- **Field-collected samples**: For laboratory work with field-collected samples, describe all relevant parameters such as housing, maintenance, temperature, photoperiod and end-of-experiment protocol OR state that the study did not involve samples collected from the field.
Ethics oversight (identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Human research participants

Policy information about studies involving human research participants

Population characteristics Describe the covariate-relevant population characteristics of the human research participants (e.g. age, gender, genotypic information, past and current diagnosis and treatment categories). If you filled out the behavioural & social sciences study design questions and have nothing to add here, write "See above."

Recruitment Describe how participants were recruited. Outline any potential self-selection bias or other biases that may be present and how these are likely to impact results.

Ethics oversight (identify the organization(s) that approved the study protocol.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Clinical data

Policy information about clinical studies

All manuscripts should comply with the ICMJE guidelines for publication of clinical research and a completed CONSORT checklist must be included with all submissions.

Clinical trial registration Provide the trial registration number from ClinicalTrials.gov or an equivalent agency.

Study protocol Note where the full trial protocol can be accessed OR if not available, explain why.

Data collection Describe the settings and locales of data collection, noting the time periods of recruitment and data collection.

Outcomes Describe how you pre-defined primary and secondary outcome measures and how you assessed these measures.

Dual use research of concern

Policy information about dual use research of concern

Hazards

Could the accidental, deliberate or reckless misuse of agents or technologies generated in the work, or the application of information presented in the manuscript, pose a threat to:

- No
  - □ Public health
  - □ National security
  - □ Crops and/or livestock
  - □ Ecosystems
  - □ Any other significant area

Experiments of concern

Does the work involve any of these experiments of concern:

- No
  - □ Demonstrate how to render a vaccine ineffective
  - □ Confer resistance to therapeutically useful antibiotics or antiviral agents
  - □ Enhance the virulence of a pathogen or render a nonpathogen virulent
  - □ Increase transmissibility of a pathogen
  - □ Alter the host range of a pathogen
  - □ Enable evasion of diagnostic/detection modalities
  - □ Enable the weaponization of a biological agent or toxin
  - □ Any other potentially harmful combination of experiments and agents
ChIP-seq

Data deposition
- Confirm that both raw and final processed data have been deposited in a public database such as GEO.
- Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

Data access links
May remain private prior to publication.
For "initial submission" or "revised version" documents, provide reviewer access links. For your "final submission" document, provide a link to the deposited data.

Files in database submission
Provide a list of all files available in the database submission.

Genome browser session
Provide a link to an anonymized genome browser session for "initial submission" and "revised version" documents only, to enable peer review. Write "No longer applicable" for "final submission" documents.

Methodology

Replicates
Describe the experimental replicates, specifying number, type and replicate agreement.

Sequencing depth
Describe the sequencing depth for each experiment, providing the total number of reads, uniquely mapped reads, length of reads and whether they were paired- or single-end.

Antibodies
Describe the antibodies used for the ChIP-seq experiments; as applicable, provide supplier name, catalog number, clone name, and lot number.

Peak calling parameters
Specify the command line program and parameters used for read mapping and peak calling, including the ChIP, control and index files used.

Data quality
Describe the methods used to ensure data quality in full detail, including how many peaks are at FDR 5% and above 5-fold enrichment.

Software
Describe the software used to collect and analyze the ChIP-seq data. For custom code that has been deposited into a community repository, provide accession details.

Flow Cytometry

Plots
- Confirm that:
  - The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
  - The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a ‘group’ is an analysis of identical markers).
  - All plots are contour plots with outliers or pseudocolor plots.
  - A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation
Describe the sample preparation, detailing the biological source of the cells and any tissue processing steps used.

Instrument
Identify the instrument used for data collection, specifying make and model number.

Software
Describe the software used to collect and analyze the flow cytometry data. For custom code that has been deposited into a community repository, provide accession details.

Cell population abundance
Describe the abundance of the relevant cell populations within post-sort fractions, providing details on the purity of the samples and how it was determined.

Gating strategy
Describe the gating strategy used for all relevant experiments, specifying the preliminary FSC/SSC gates of the starting cell population, indicating where boundaries between "positive" and "negative" staining cell populations are defined.

Magnetic resonance imaging

Experimental design

Design type
Indicate task or resting state; event-related or block design.
**Design specifications**
Specify the number of blocks, trials or experimental units per session and/or subject, and specify the length of each trial or block (if trials are blocked) and interval between trials.

**Behavioral performance measures**
State number and/or type of variables recorded (e.g. correct button press, response time) and what statistics were used to establish that the subjects were performing the task as expected (e.g. mean, range, and/or standard deviation across subjects).

### Acquisition

**Imaging type(s)**
Specify: functional, structural, diffusion, perfusion.

**Field strength**
Specify in Tesla

**Sequence & imaging parameters**
Specify the pulse sequence type (gradient echo, spin echo, etc.), imaging type (EPI, spiral, etc.), field of view, matrix size, slice thickness, orientation and TE/TR/flip angle.

**Area of acquisition**
State whether a whole brain scan was used OR define the area of acquisition, describing how the region was determined.

**Diffusion MRI**
- [ ] Used
- [x] Not used

### Preprocessing

**Preprocessing software**
Provide detail on software version and revision number and on specific parameters (model/functions, brain extraction, segmentation, smoothing kernel size, etc.).

**Normalization**
If data were normalized/standardized, describe the approach(es); specify linear or non-linear and define image types used for transformation OR indicate that data were not normalized and explain rationale for lack of normalization.

**Normalization template**
Describe the template used for normalization/standardization, specifying subject space or group standardized space (e.g. original Talairach, MINI305, ICBM152) OR indicate that the data were not normalized.

**Noise and artifact removal**
Describe your procedure(s) for artifact and structured noise removal, specifying motion parameters, tissue signals and physiological signals (heart rate, respiration).

**Volume censoring**
Define your software and/or method and criteria for volume censoring, and state the extent of such censoring.

### Statistical modeling & inference

**Model type and settings**
Specify type (mass univariate, multivariate, RSA, predictive, etc.) and describe essential details of the model at the first and second levels (e.g. fixed, random or mixed effects; drift or auto-correlation).

**Effect(s) tested**
Define precise effect in terms of the task or stimulus conditions instead of psychological concepts and indicate whether ANOVA or factorial designs were used.

**Specify type of analysis**
- [ ] Whole brain
- [ ] ROI-based
- [x] Both

**Statistical type for inference**
(See Eboud et al. 2016)
Specify voxel-wise or cluster-wise and report all relevant parameters for cluster-wise methods.

**Correction**
Describe the type of correction and how it is obtained for multiple comparisons (e.g. FWE, FDR, permutation or Monte Carlo).

### Models & analysis

**n/a Involved in the study**
- [ ] Functional and/or effective connectivity
- [ ] Graph analysis
- [ ] Multivariate modeling or predictive analysis

**Functional and/or effective connectivity**
Report the measures of dependence used and the model details (e.g. Pearson correlation, partial correlation, mutual information).

**Graph analysis**
Report the dependent variable and connectivity measure, specifying weighted graph or binarized graph, subject- or group level, and the global and/or node summaries used (e.g. clustering coefficient, efficiency, etc.).

**Multivariate modeling and predictive analysis**
Specify independent variables, features extraction and dimension reduction, model, training and evaluation metrics.