Identification of major zinc-binding proteins from a marine cyanobacterium: insight into metal uptake in oligotrophic environments†

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Marine cyanobacteria make a significant contribution to primary production whilst occupying some of the most nutrient-poor regions of the world’s oceans. The low bioavailability of trace metals can limit the growth of phytoplankton in ocean waters, but only scarce data are available on the requirements of marine microbes for zinc. Recent genome mining studies suggest that marine cyanobacteria have both uptake systems for zinc and proteins that utilize zinc as a cofactor. In this study, the oligotrophic strain Synechococcus sp. WH8102 was grown at different zinc concentrations. Using metalloproteomics approaches, we demonstrate that even though this organism’s growth was not affected by extremely low zinc levels, cells accumulated significant quantities of zinc, which was shown to be protein-associated by 2D liquid chromatography and ICP-MS. This indicates that the mechanisms for zinc uptake in Synechococcus sp. WH8102 are extremely efficient. Significantly, expression of SYNW2224, a putative porin, was up-regulated during growth in zinc-depleted conditions. Furthermore, along with 30 other proteins, SYNW2224 was captured by immobilised zinc affinity chromatography, indicating the presence of surface-exposed site(s) with metal-binding capacity. It is proposed that this porin plays a role in high-affinity zinc uptake in this and other cyanobacteria.
marine cyanobacteria for zinc are scarcely studied,\textsuperscript{23–25} whereas for some freshwater cyanobacteria effects of zinc toxicity and deprivation have been examined in some detail.\textsuperscript{26,27} The genomes of over 30 marine cyanobacterial strains have been sequenced, and our genome mining work has discovered not only genes for several enzymes predicted to require a zinc cofactor, including alkaline phosphatase and one or more carbonic anhydrases, but also for proteins with likely roles in zinc uptake and trafficking, including putative periplasmic zinc binding proteins (ZnuA).\textsuperscript{28–30} All strains inspected also harboured a gene for the zinc uptake regulator Zur, with recognition sites for this zinc sensor protein predicted to occur in the upstream regions of several relevant genes. Whilst these bioinformatic studies strongly point towards dedicated networks for zinc uptake and utilisation, an absolute requirement of marine cyanobacteria for zinc has not been demonstrated conclusively.

In the present study we have used the oligotrophic open-ocean strain \textit{Synechococcus} sp. WH8102\textsuperscript{24} as a model cyanobacterium to study its growth and cellular zinc quota under zinc-depletion conditions, and to probe the presence of major Zn\textsuperscript{2+}-binding proteins using solution-state metalloproteomics approaches. Metalloproteomics, a sub-discipline of metallomics, is dedicated to the provision of experimental evidence for metal–protein interactions.\textsuperscript{32–36} Combinations of inorganic and molecular mass-spectrometry are particularly powerful approaches, whilst separation techniques have posed the major experimental challenges.\textsuperscript{2}

Previously, we examined the applicability of different liquid-chromatography-based approaches to probe the Fe\textsuperscript{3+}, Ni\textsuperscript{2+}, and Co-related proteome of \textit{Synechococcus} sp. WH8102.\textsuperscript{25,27} In the present study, immobilised zinc affinity chromatography (Zn\textsuperscript{2+}-IMAC) has enabled the capture and detection of several proteins with potential functions in zinc metabolism. Besides the detection of a predicted periplasmic zinc-binding protein (ZnuA) along with several other periplasmic binding proteins, the most significant finding concerns a predicted porin, a novel candidate for mediating zinc uptake across the outer membrane of marine cyanobacteria.

**Materials and methods**

**Suppliers of chemicals and reagents**

All chemicals used were of the highest grade available and were purchased from either Fisher Scientific (UK) or Sigma-Aldrich (UK) unless otherwise indicated.

**Bacterial strains and growth conditions**

Axenic cultures of \textit{Synechococcus} sp. WH8102 were grown in an artificial seawater medium based on Aquil,\textsuperscript{38,39} prepared using ultrapure (MilliQ 18 MΩ cm\textsuperscript{-1}) water (Table S1, ESI\textsuperscript{t}). The base medium (macronutrients) was Chelex-treated and then autoclaved. Micronutrient components were filter-sterilised before adding to the base medium. Media bottles and culture flasks were washed with 5% trace metal clean HCl prepared in house by sub-boiling point distillation and rinsed with MilliQ water before use.

The theoretical free Zn\textsuperscript{2+} concentration in the growth medium at pH 8.0 was calculated from the total EDTA and Zn\textsuperscript{2+} concentrations using the “Species” module within the IUPAC stability constants database (Data version 4.56, L.D. Pettit, Academic Software, UK), taking also into account the concentrations of other metal ions. Stability constants and pH\textsubscript{0} values were also extracted from this database.

Stocks of \textit{Synechococcus} sp. WH8102 were maintained through sub-culturing in the different Zn media that were tested, over a long period of time. Cells were initially transferred from standard medium to zinc-depleted medium by collecting cells by centrifugation, gently washing and resuspending into the zinc-depleted medium. Before the growth measurements were carried out, cells were acclimated over several serial transfers across at least 12 weeks into fresh medium with either no zinc or 80 nM added zinc. Cultures were grown at 25 °C with continuous illumination at 10 μE m\textsuperscript{-2} s\textsuperscript{-1} and shaken at 150 rpm. This light level corresponds to levels found at the bottom of the surface mixed layer, a region where clade III \textit{Synechococcus} genotypes (of which \textit{Synechococcus} sp. WH8102 is a member) proliferate. Such a light level also assists in stable culture maintenance, without inducing photodamage. Growth was monitored by measuring the optical density at 750 nm or by flow cytometry using a FACScan flow cytometer (Becton Dickinson, NJ, USA). Cultures were checked at regular intervals for contamination with other microorganisms by plating onto Aquil-Agar plates containing 500 mg l\textsuperscript{-1} yeast extract. Cells were harvested by centrifugation at 6000 × g. All growth and subsequent separation experiments were carried out in duplicate.

**Preparation of whole cell lysates**

Cell pellets were re-suspended in 1–10 ml of 10 mM HEPES pH 7.2, 0.5% (w/v) octyl β-D-glucopyranoside, with a dissolved Complete\textsuperscript{TM} EDTA-free protease inhibitor cocktail tablet (Roche, UK). Cells were broken by sonication and debris was removed by centrifugation at 12 000 × g. The BCA method\textsuperscript{40} was used to determine the total protein content of cell lysates.

**Preparation of soluble and insoluble cell fractions**

Cell pellets were re-suspended in 8 ml of 20 mM HEPES pH 7.2 containing a dissolved Complete\textsuperscript{TM} EDTA-free protease inhibitor cocktail tablet (Roche, UK), and broken by sonication. Unbroken cells were removed by centrifugation at 6000 × g. The soluble and insoluble fractions were separated by ultracentrifugation at 200 000 × g for 30 minutes using a Beckman TLA100.3 rotor. The supernatant (soluble fraction) was collected, and the pellet (insoluble fraction) was re-suspended in 1 ml of 20 mM HEPES pH 7.2, 0.5% (w/v) octyl β-D-glucopyranoside to solubilise membrane proteins. The prepared fractions were further clarified by filtration using 0.2 μm pore-sized filters.

**Preparation of a carboxysome enriched fraction**

Carboxysomes were prepared according to the method described by Gonzales \textit{et al.},\textsuperscript{41} with some minor modifications. Briefly, cell pellets were resuspended in 3 ml of 20 mM HEPES pH 7.2 containing a Complete\textsuperscript{TM} EDTA-free protease inhibitor cocktail tablet. Cells were disrupted by sonication and cell debris was removed by centrifugation at 12 000 × g for 10 minutes.
The collected supernatant was centrifuged at 40 000 × g at 4 °C for 30 minutes using a Beckman TLA100.3 rotor. The dark green pellet was resuspended in 3 ml of 20 mM HEPES pH 7.2, 20 mM MgSO₄, 2% (v/v) Triton X-100 and incubated on ice for 45 minutes with occasional agitation. This step solubilises membrane lipids, whilst promoting carboxysome aggregation. The sample was centrifuged again at 40 000 × g to generate a yellow-brown carboxysome enriched pellet that was resuspended in 20 mM HEPES pH 7.2.

Inductively coupled plasma mass spectrometry (ICP-MS)

3% (v/v) trace metal grade nitric acid purified in house by sub-boiling point distillation was used as the sample matrix. For quantitation, calibration was performed in the range 0–500 ppb using external Zn and P ICP-MS standards. Er (Agilent Technologies, USA) was used as an internal standard. Measurements were taken using an Agilent 7500 series ICP mass spectrometer (Agilent Technologies, USA), equipped with a cross flow nebulizer, quartz spray chamber, and an Octopole Reaction System (ORS) cell. All samples were measured in triplicate in helium gas-mode to remove matrix interferences. Cell lysates were prepared as described below from the combined cell pellets of 3 replicate cultures, and were diluted 1:20 in matrix for analysis.

Two-dimensional liquid chromatography (2D-LC)

Two-dimensional liquid chromatography (2D-LC) was performed essentially as described previously.³⁷ 5 mg total protein from the soluble, insoluble, or carboxysome enriched cell fraction was applied to a Biosep S2000 HPLC gel filtration column (Phenomenex, UK) equilibrated in 10 mM HEPES pH 7.2 (containing 0.01% (w/v) octyl β-D-glucopyranoside for the insoluble fraction). Protein was eluted isocratically in the same buffer using a flow rate of 1 ml min⁻¹ and collected in 1.0 ml fractions. The column was calibrated using 1 mg ml⁻¹ solutions of bovine serum albumin – 66 kDa, Carbonic anhydrase – 29 kDa, lysozyme – 14 kDa, and substance P – 1 kDa standards in column buffer. Dextran blue dye (GE Healthcare, UK) was used to determine the void volume (V₀), which was 5.1 ml. Protein containing fractions [5–13 ml] from 4 separate column runs (except for the carboxysome enriched cell fraction where just 1 column run was performed) were combined and further separated using strong anion exchange mini-spin columns (Thermo Scientific, UK). The spin columns were equilibrated using 2 mM HEPES pH 7.2 (plus 0.01% (w/v) octyl β-D-glucopyranoside for the insoluble fraction). Protein was eluted using a NaCl gradient of 0–2 M in 80 μl fractions. 50 μl of these samples were used for ICP-MS analysis and 30 μl were retained for analysis by SDS-PAGE and peptide mass fingerprinting. All stock solutions were prepared using 18 MΩ water and treated with Chelex-100 resin (Bio-Rad, UK) to remove traces of contaminating metal ions. Acid-washed plastic-ware was used throughout to prevent contamination of samples with external sources of zinc or other metals. For the 2D-LC experiments for soluble and insoluble fractions, two biological replicates were carried out, with reproducible results.

Immobilized metal affinity chromatography (IMAC)

1 ml IMAC columns (GE Healthcare, UK) were prepared according to the manufacturer’s instructions and either charged with Zn²⁺ or left un-charged. Columns were equilibrated with 10 mM HEPES pH 7.2, 0.5 M NaCl, 0.01% (w/v) octyl β-D-glucopyranoside. 20 mg of protein from crude cell lysates were loaded onto each column and the flow-through (FT) fraction collected. Unbound protein was washed through the column using 4 × 1 ml of buffer containing 2 mM imidazole (W1–W4), before bound proteins were eluted using 2 × 1 ml of buffer with 20 mM imidazole (E1–E2) and 2 × 1 ml of buffer containing 200 mM imidazole (E3–E4).

SDS-polyacrylamide gel electrophoresis

SDS-PAGE was performed using mini-Protean III TGX™ precast 4–15% gels (Bio-Rad, UK) using standard protocols.⁴² Samples were mixed with an equal volume of gel loading buffer and heated to 80 °C for 5 minutes. 50 μl of sample was loaded per lane except where otherwise stated. Gels were stained using Coomassie brilliant blue R-250 (National Diagnostics, USA).

Peptide mass fingerprinting

Protein bands were excised from SDS-PAGE gels using a scalpel blade and subjected to in-gel tryptic digestion using a commercially produced kit (Pierce, Thermo Scientific, UK). Peptide masses were determined by MALDI-TOF MS analysis. 2 μl of sample matrix (10 mg ml⁻¹ azacyano-4-hydroxycinnamic acid in 50% acetonitrile, 0.1% trifluoroacetic acid) was mixed with 2 μl of sample and spotted onto a steel MALDI-target plate. Peptide masses were determined using a Bruker Ultraflex II MALDI-TOF/TOF mass spectrometer (Bruker Daltonics, Coventry, UK) with a 337 nm laser and operated in reflectron mode. Mass calibration was performed using PEG2000 and verified using bradykinin and substance P peptide standards. Internal mass accuracy was confirmed by the presence of the autolytic trypsin peaks at 845.2 and 2211.1 Da. Mass spectra were acquired over the range of 800–3500 Da. Mass lists were generated using Bruker Flex-analysis software with default parameters, and searched against either the NCBI or SwissProt databases using Mascot (Matrix Science, UK). The following search criteria were selected: Fixed modification of carbamidomethyl on cysteine, variable modification of oxidation of methionine, maximum of 1 missed cleavage, <50 ppm mass accuracy, “other bacteria” was selected for taxonomy. Only searches giving significant MOWSE scores were recorded.

Structural models

Comparative modelling was employed to provide insight into the likely structures of the ZnuA, FutA, CynA, and cyanobacterial porin (CBP) proteins. Suitable templates were identified using the remote homology recognition server Phyre²,⁴³ which also produces multiple and pairwise sequence alignments (see ESI†), and in favourable cases sound protein models. All initial models were generated by Phyre². The model for ZnuA was based on pdb 2OV3 (ZnuA from Synechocystis sp. PCC6803; mutant devoid of...
His-rich loop, 44 that for CynA on pdb 3UN6 (Ligand Binding Component of ABC-type Import System from Staphylococcus aureus), 23 and that for FutA was based on pdb 2PT1 (FutA from Synecocystis sp. PCC6803). 46 The ZnuA and CynA models were further developed using the molecular modelling programme MOE2011.10. Hydrogen atoms, zinc ions, and a cyanate ion in the case of CynA were introduced, and the metal sites were then energy-minimised, using an in-house customised version of the AMBER99 force-field, in which Zn-specific parameters were incorporated. After energy minimisation of the position of the immediate zinc ligands, the adjacent environment was optimised, and finally, the entire molecule was subjected to energy minimisation. All minimisations were terminated based on the steepness of the RMS gradient ( < 0.5). Models were submitted to the WHATIF web interface (http://swift.cmbi.ru.nl/servers/html/index.html), ensuring that a physically reasonable model had been produced.

The 3D model for the CBP SYNW2224 was generated using the “Intensive mode” on the Phyre server, and is based on three overlapping templates: residues 46–98 are based on pdb 3PYW (SLH domain from Bacillus anthracis), 47 residues 92–141 on pdb 3SWF (a helix from a rod cyclic nucleotide-gated ion channel), 48 and residues 134–501 on pdb 4GF4 (Pseudomonas putida OprB, a porin for carbohydrate uptake). 49 Structural images were generated in MOLMOL v.2k.50

Results

Synechococcus sp. WH8102 grows well in zinc-depleted media

To investigate if depletion of zinc may affect the growth of Synechococcus sp. WH8102, cultures were studied in an artificial seawater medium based on Aquil 38,39 that was supplemented with either 0 (zinc-depleted medium) or 80 nM (designated here as “replete” medium, although it should be noted that the resulting free zinc concentration is still very low) zinc. The presence of 100 μM EDTA leads to a theoretical free Zn$^{2+}$ concentration of 16 pM in the replete medium, with 98.98% of the added Zn$^{2+}$ complexed. Free Zn$^{2+}$ concentrations in open ocean surface waters vary from 1 to 71 pM. 11–13,16 Hence, the free Zn$^{2+}$ concentration in our replete medium is not dissimilar to those that Synechococcus sp. WH8102 might encounter in its natural habitat. The free ion concentrations are considered the most relevant factor governing uptake. 51,52

The growth data in Fig. 1(a) show that the cultures grown with 0 or 80 nM zinc exhibited no significant difference in growth rate or final cell yield. This suggests that Synechococcus sp. WH8102 either has no absolute requirement for zinc, or that it is able to scavenge trace amounts of zinc from the depleted growth medium, which are unavoidably introduced as a contaminant with other media components.

Efficient zinc uptake at extremely low zinc concentrations

In a subsequent experiment, following 19 days growth in either zinc depleted or replete medium, cells were harvested and the zinc and phosphorus content of crude cell lysates was measured by Inductively-Coupled-Plasma-Mass-Spectrometry (ICP-MS). The phosphorus levels were used as a proxy for biomass. The cells grown under zinc-depleted conditions had accumulated significant quantities of zinc, and no significant difference in the Zn : P ratio was recorded between the two conditions tested (Fig. 1(b)).

The fact that significant amounts of zinc were captured by Synechococcus sp. WH8102 even at extremely low concentrations points to the existence of highly efficient uptake mechanisms. Significantly, analysis of 1-dimensional SDS-PAGE gels of the complete proteomes from cells grown under the two regimes revealed that the amount of an outer-membrane protein (SYNW2224) was markedly increased under zinc-depleted conditions suggesting a potential role in zinc acquisition for this protein (Fig. 1(c)). A second protein band was visibly diminished

Fig. 1 Effect of zinc on the growth of Synechococcus sp. WH8102 and subcellular distribution of zinc. (a) Growth of Synechococcus sp. WH8102 in Aquil medium containing either 0 nM (dashed line), or 80 nM (solid line) added zinc. (b) Zn : phosphorus ratios in whole cells. (c) CBB-stained SDS-PAGE gel comparing the protein profiles of crude cell lysates prepared from Synechococcus sp. WH8102 cells grown in zinc-depleted or replete media. 32 μg of protein was loaded per lane. Proteins showing a marked difference in expression are boxed. Box 1 corresponds to a putative CBP (SYNW2224; identified with 14 matched peptides, 33% sequence coverage and a MOWSE score of 78). Box 2 yielded the large subunit of RuBisCO as possible hit (6 matched peptides, 14% sequence coverage, MOWSE score 68). The large deviation between the observed and expected molecular weight could be due to persistence of the homo-dimer or a larger complex. 115
under zinc-depletion conditions; this was tentatively identified as the large subunit of RuBisCO, although it is not clear why the observed molecular weight was considerably larger than expected, or why differences in zinc levels should affect the abundance of this protein.

**Fractionation of the zinc proteome by 2-dimensional liquid chromatography**

In an attempt to isolate intact Zn-bound proteins from *Synechococcus* sp. WH8102, the soluble and insoluble proteomes, as well as carboxysomal fractions, were further fractionated using native 2-dimensional liquid chromatography (2D-LC). This approach was selected as it allows proteins to be separated rapidly and under mild conditions that promote the likelihood for keeping proteins folded and intact including moderately strong metal cofactors. It should therefore in principle be suitable for the separation of Zn-containing proteins that form relatively stable complexes. However, a drawback is the resolution of separation achievable by 2D-LC, which is considerably lower than that obtained using traditional denaturing 2D-gel electrophoresis methods.

In the present study, proteins were initially separated using size exclusion chromatography (SEC), followed by anion exchange chromatography (AEX). The “insoluble” fraction was solubilised and separated in the presence of the mild detergent octyl glucoside, which has been used extensively for the isolation of native membrane proteins. In each case, 5 mg total protein was applied to a BioSep S2000 HPLC gel filtration column and protein was eluted in 13 × 1 ml fractions. The elution fractions from four separate SEC runs were combined and applied to mini-spin anion exchange columns. These columns allowed the rapid and simultaneous separation and concentration of the gel-filtration fractions. Bound proteins were eluted from the anion exchange columns in 80 µl fractions in a step-wise fashion using a NaCl gradient of 0–2 M. Each of the collected fractions was then analysed by ICP-MS for zinc content (Fig. 2). In the soluble fraction, zinc eluted from the gel filtration column as a single broad peak between ca. 14 kDa and 70 kDa. In the insoluble fraction, two distinct peaks were observed, one at ca. 30–70 kDa and a second at around 1 kDa, which contains peptides, other small molecules, and possibly also free metal ions. The proteins present in fractions with the highest zinc concentrations were analysed by SDS-PAGE (Fig. 2). For the soluble fraction, the anion exchange samples obtained from separation of the 9 ml gel filtration fraction were analysed. Despite the concentration step, very few proteins were observed on the gel, but one protein migrating to ca. 18 kDa on the gel could be observed even before staining by its light pink colour. This protein was identified by peptide mass fingerprinting and found to be a subunit of c-phycoerythrin, a component of the light harvesting phycobilisome complex. After the gel was stained with Comassie, only one further protein (at ca. 10 kDa) could be visualised. This protein was identified as ribosome recycling factor, and neither of these two proteins is predicted to bind metal ions. However, like other biliproteins, c-phycoerythrin contains linear tetrapyrrrole chromophores (phycobilins) with considerable metal-binding ability. It appears likely that cellular zinc has partially been redistributed to the chromophores on these proteins. In support of this hypothesis, several biliproteins were also captured on Zn$^{2+}$-IMAC columns (see below). The redistribution of zinc in our samples might explain why no predicted zinc binding proteins including alkaline phosphatase or carbonic anhydrase were detected in the soluble fractions following 2D-LC. Similar results were obtained from the analysis of the insoluble cell fraction, with c-phycoerythrin again the most abundant protein present in the zinc enriched cell fractions (Fig. 2). The only other protein detected in the peak zinc fractions was PstS, a periplasmic phosphate binding protein. Whilst this protein is not predicted to bind zinc or any other metals *in vivo*, it was captured using a Zn$^{2+}$-IMAC column (see below), suggesting that it has an affinity for zinc *in vitro*. Full details of the proteins identified are given in Table S2 (ESI†).

In the carboxysome fraction, zinc was only detected in lower molecular weight sub-fractions (20 kDa and below), but the concentrations of proteins were too low for identification by peptide mass fingerprinting.

In conclusion, despite the rapid and mild separation conditions used for the chromatography and the relatively high stability of Zn$^{2+}$ complexes according to the Irving-Williams series, no major zinc-binding proteins in any of the sub-cellular fractions could be identified by the 2D-LC approach. Major drawbacks were the low resolution of the separation steps and insufficient sensitivity in protein detection, both of which are exacerbated by the presence of highly abundant biliproteins. These pose a significant challenge for native metalloproteomics in this organism and likely other related cyanobacteria. In terms of classical mass-spectrometry-based proteomics, biliproteins cause dynamic range problems, since they can account for as much as 60% of total cellular protein; hence, they are inherently likely to impede the detection of low abundance proteins. In terms of native metalloproteomics, their demonstrated metal-binding ability (also see below) causes additional problems, as this may lead to metal redistribution in cell lysates – a problem unlikely to be solvable by depletion strategies.

**Enrichment of zinc-binding proteins by immobilized metal ion affinity chromatography**

Metal-trafficking proteins often bind their cargo in a kinetically labile fashion and close to the protein surface, to enable facile transfer to and from other molecules. This is an ideal prerequisite to capture such proteins by immobilized metal ion affinity chromatography (IMAC). Similar approaches have been used in previous studies to isolate proteins that have an affinity for the immobilised metal ion *in vitro*, e.g. proteins with copper affinity from human liver cells, *Arabidopsis thaliana*, and microalgae, and metal-binding proteins from plant mitochondria.

Eluates from a Zn$^{2+}$-IMAC column were analysed by 1D SDS-PAGE (Fig. 3), and proteins were identified by peptide mass fingerprinting. A total of 30 different proteins with zinc-binding ability were identified (Table 1), including several enzymes, biliproteins, carboxysomal shell proteins, periplasmic binding proteins, and two outer membrane proteins. Based on biochemical
data from homologous proteins, some of these proteins are predicted to bind metal ions; others, including biliproteins and carboxysomal shell proteins, are not. Identified proteins are described and discussed below.

**Enzymes and regulatory proteins.** Amongst the proteins captured, there were several enzymes predicted to require zinc for function, including fructose bisphosphate aldolase (SYNW0791), transketolase (TktA; SYNW0141), dihydrolipoyl dehydrogenase (LdpA; SYNW1630), RNA polymerase, and a putative alkaline phosphatase. A related fructose bisphosphate aldolase from the protozoan *Giardia lamblia* (pdb 3GAK) coordinates a zinc ion via two His and one Glu residue, with a further Asp residue that can contribute to an alternative binding site nearby. 59 All of these zinc-binding residues are conserved in the protein from *Synechococcus* sp. WH8102, which otherwise is 43% identical to the enzyme from *G. lamblia*. The zinc site in the *G. lamblia* protein is mobile and at least the two His residues are fairly surface-exposed, with the potential for exchange between any eventual protein-bound metal and immobilised zinc on the IMAC column. Whilst we also note that fructose bisphosphate aldolase...
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The enzymes transketolase (TktA) and dihydrolipoyl dehydrogenase (LpdA) from E. coli were previously found to bind Zn after separation of the proteome by denaturing 2D gel electrophoresis. More recently, in vivo zinc binding was demonstrated for both enzymes by in vivo labelling with Zn and subsequent separation by native 2D gel electrophoresis. TktA from E. coli (pdb 2R80) contains at least six His residues in its substrate-binding pocket, which are fully conserved in the homologue from Synechococcus sp. WH8102. These His residues are involved in binding the diphosphate moiety, the sugar and the phosphate of the D-fructose-6 phosphate group which are fully conserved in the substrate-binding pocket, which are conserved in the enzyme from Synechococcus sp. WH8102. Whether the latter site is of significance regarding enzymatic activity is unknown.

In addition, several further enzymes were also captured on the IMAC column. Using structural models, CHED analysis, and manual inspection, possible reasons for this are explored below. Glutaredoxin (modelled on pdb 3QMX, from Synechocystis) contains three Cys residues, and one of them is flanked by a His and an Asp residue, both surface-accessible, hence in principle suitable for IMAC capture. Ruberythrin from Desulfovibrio vulgaris (pdb 1DVB) contains binding sites for three iron ions, one Cys site, and a binuclear site, which are conserved in RbrA from Synechococcus sp. WH8102. As a relatively weakly-binding metal ion, Fe is prone to be lost during protein separation. Thus, in principle, sites in Fe-proteins may become available for binding to immobilised metal. The most surface-accessible residues are two Cys residues, whereas the His and Glu residues in the binuclear site are buried in the folded protein, and are not clustered in the primary sequence, so relatively unlikely to be responsible for IMAC binding.

SYNW2391 is annotated as a putative alkaline phosphatase, is structurally related to 7-bladed β-propeller oxidoreductases, and contains an abundance of potential metal sites (CHED detected seven sites in total). Similarly, at least five metal-binding sites can be identified in glutamine synthetase (GlnA). The protein is 55% identical to GlnA from Salmonella typhimurium (pdb 1FPY). One large, surface-exposed site involving several Glu and His residues is the ATP-binding site, which also requires two M ions for binding and ATP hydrolysis. Essentially similar considerations hold for the beta subunit of RNA polymerase RpoC2 which also harbours a Mg-requiring ATP-binding site. In addition, a structural ZnCys site is present, but this is deeply buried (RpoC2 was modelled on pdb 4G7O, RNA polymerase from Thermus thermophilus). Similarly, the large subunit of RubisCO harbours three potential metal sites, one of them coinciding with the binding site for ribulose-1,5-bisphosphate, which also requires a Mg ion. It is hence conceivable that similar ternary complexes can also be formed with NTA-immobilised Zn (or indeed other immobilised metal ions). O-Acetyl-homoserine sulfhydrylase also displayed three potential metal sites, one of them containing a Cys residue, but not related to enzymatic activity. Phosphoglucomutase, glucose-6-phosphate isomerase, and the oligosaccharide phosphorylase GlgP are all part of sugar metabolism, and are enzymes that work in sequence. GlgP is required for the breakdown of oligosaccharides, resulting in glucose-1-phosphate. This is converted to glucose-6-phosphate by phosphoglucomutase, and this is converted to fructose-6-phosphate by the isomerase. Several oligosaccharide phosphorylases are reported to be

Fig. 3 Isolation of zinc binding proteins by Zn -IMAC. CBB-stained gel of Zn-IMAC column fractions. 20 mg of crude cell lysate was applied to a 1 ml Zn-IMAC column and the flow-through (FT) collected. Unbound protein was washed through the column using buffer containing 2 mM imidazole (W1–W4). Strongly bound protein was eluted using buffer containing 20–200 mM imidazole (E1–E4). Proteins present in the elution fractions were identified by peptide mass fingerprinting (Table 1).
activated, stimulated, or inhibited by various metal ions (http://www.brenda-enzymes.info/php/result_flat.php?ecno=2.4.1.1), suggesting the presence of metal binding sites, and indeed, six potential sites were detected by CHED. Most phosphoglucomutases require Mg²⁺ for activity, but show limited activity with various other metal ions. The Synechococcus from the ciliate WH8102 phosphoglucomutase shares 51% identity with that

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**Components of phycobilisomes**

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Note: This is the protein cluster number in the Cyanorak database which is publically accessible at http://www.sb-roscoff.fr/Phyto/cyanorak/.

The two-component response regulator RpaB is a two-domain protein, and two potential metal-binding sites in its N-terminal domain were detected by CHED. It shares 42% identity with the RegX3 regulator from *Mycobacterium tuberculosis* (pdb 2OQR7⁴), and the crystal structure of the latter has been stabilised by La³⁺ ions, although metals are not thought to play a role in the activity of this protein. The metal-binding residues in RpaB and RegX3 are not identical, but the location of the exposed N-terminal sites is roughly similar. Elongation factor Tu displays an abundance of clustered, surface-exposed His and Glu residues forming at least three sites, with no clear functional significance, and not related to the GTP-binding site.⁷⁵ Two potential metal binding sites can be predicted for elongation factor G, again without relationship to known protein function.⁷⁶ CHED analysis of structural models of subunit beta of ATP synthase and porphobilinogen deaminase did not reveal any pertinent metal binding sites.

**Carboxysomal proteins.** Two carboxysomal shell proteins, annotated as CcmK/CsoS1 (SYNW1719) and CcmK2 (SYNW1712).
were captured on the Zn-IMAC column.‡ In a previous study, we found that CcmK1 also had a strong affinity for Co\textsuperscript{2+}-IMAC columns.\textsuperscript{37} Considering that the carboxysomes also house carboxic anhydrase, the unexpected affinity for zinc and cobalt displayed by the CsoS1 proteins was intriguing. Fig. S1 (ESI\textsuperscript{†}) shows a homology model for CcmK1 (103 aa), which adopts a typical bacterial microcompartment domain fold. The larger Ccmk2 (183 aa, modeled using pdb 3NWG, a PduT homolog from a novel bacterial microcompartment from \textit{Desulfotobacterium hafniense})\textsuperscript{77} contains two such domains (see Fig. S2, ESI\textsuperscript{†}). Both proteins display similar surface-exposed patches of potential metal ligands (Glu, His), although the CHED program did not report any metal sites in either protein, even under the least stringent search conditions. Since these proteins form multimeric complexes as part of the carboxysome shell, a hexameric assembly of a closely related CsoS1 protein was also inspected (pdb 2EWH; CsoS1A from \textit{Halothiobacillus neapolitanus}),\textsuperscript{78} but no additional inter-protein metal sites were detected. It is conceivable that the loop containing E86, H88 and E90 (CcmK1 numbering; the same motif is also present in the C-terminal portion of CcmK2) is flexible enough to provide a metal-binding motif with sufficient affinity for binding to an IMAC column, at least in the monomeric protein. We note that the pore structures formed by the structurally related ethanolamine-utilising microcompartment shell protein, EutL from \textit{E. coli}, were affected by the presence of Zn\textsuperscript{2+},\textsuperscript{79} but in neither case is it clear yet whether an interaction between microcompartment shell proteins and Zn\textsuperscript{2+} is of physiological relevance, or only occurs \textit{in vitro}.

**Periplasmic binding proteins.** The ZnuA component (SYNW2481) of a putative ZnuABC zinc uptake transporter was identified in the Zn-IMAC eluate, confirming its expression, and supporting the hypothesis that this system could operate in zinc uptake in \textit{Synechococcus} sp. WH8102. True ZnuA proteins can often be distinguished\textsuperscript{28} from the closely related Mn\textsuperscript{2+} binding proteins (MntC)\textsuperscript{80} by the presence of a His-rich loop (in ZnuA), but the putative ZnuA orthologues from \textit{Synechococcus} sp. WH8102§ are unusual in that they do not contain this His-rich loop. Both MntCs and ZnuAs contain three strictly conserved His residues in their folded part that are involved in forming the major metal-binding site. A homology model for SYNW2481 (Fig. 4(a)) shows the protein fold including this binding site.

However, several other periplasmic binding proteins were also captured on the Zn-IMAC column, including putative iron (FutA/IdiA; SYNW1797), cyanate (CynA; SYNW2487), and phosphate (PstS; SYNW1018) transporters. The interaction of the iron-binding protein FutA with the immobilised Zn\textsuperscript{2+} is unsurprising, because a certain degree of affinity for a non-cognate metal ion can be expected based on simple coordination chemistry principles. In contrast, the capture of the predicted cyanate (CynA) and phosphate (PstS) binding proteins was surprising.

‡ Since WH8102 contains π carboxysomes, the term CsoS1 should be used (Badger and Price, 2003) – ccm (for carbon-concentrating mechanism) proteins are components of β carboxysomes, but this nomenclature is widely ignored in genome annotations; therefore, we have retained the names as they appear in relevant databases.

§ SYNW0971 is also predicted as ZnuA.

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**Fig. 4** Modelled metal-binding sites in periplasmic binding proteins captured on a Zn-IMAC column. (a) Putative ZnuA (SYNW2481) from \textit{Synechococcus} sp. WH8102. The zinc ion is coordinated to three His residues (H67, H139, H198) and a fourth non-proteinaceous ligand, modelled as water. (b) Putative cyanate transporter CynA (SYNW2487). The model (residues 197–520) is based on pdb 3UN6,\textsuperscript{45} an uncharacterised protein from \textit{Staphylococcus aureus}. Zinc is coordinated to three Cys residues and a cyanate ion. The fold of the modelled CynA protein is very similar to that of periplasmic nitrate and hydrogen carbonate binding proteins, but none of the metal-binding residues are conserved in either nitrate or hydrogen carbonate transporters from \textit{Synechocystis} sp. PCC6803. (c) Detailed view of the metal- and cyanate-binding site, with His316 and Trp237 as potential H-bond donors.
Therefore, homology models for both latter proteins were generated to inspect potential metal-binding sites. The model for the phosphate binding protein PstS did not reveal any clear metal sites (not shown). In contrast, one of the templates identified for the cyanate transporter CynA contained zinc ions in the binding cleft, one of them coordinated to two Cys and one His residue plus one water molecule. The template (pdb 3UN6)\(^\text{35}\) refers to an uncharacterised protein from *Staphylococcus aureus*. Like CynA, this protein is most closely related to the COG0715 cluster which contains periplasmic components of ABC-type nitrate–sulfonate–bicarbonate transport systems. The two Cys residues are conserved in CynA, and the His residue is replaced by another Cys residue (Fig. 4(b) and (c)). It is likely that at least some of these residues are responsible for the observed interaction with the immobilised Zn\(^{2+}\), and considering the nature of the ligands, \emph{in vivo} Zn\(^{2+}\) and/or Cd\(^{2+}\) binding is most likely. It is plausible that the metal ion facilitates the binding of the cyanate anion to be transported. Considering that Zn\(^{2+}\) (or any other metal ion) is needed only in catalytic quantities inside cells, whilst cyanate provides the macronutrient nitrogen, co-transport of the metal appears less likely. It is noteworthy that the orthologous bicarbonate transporter CmpA from *Synechocystis* sp. PCC6803 utilises a Ca\(^{2+}\) ion to enhance bicarbonate binding.\(^{41}\)

**Outer membrane proteins.** Perhaps most intriguingly, a predicted outer membrane protein, (SYNW2224) that was found to be more abundant under low-zinc conditions (Fig. 1(c)), was also one of the proteins captured on the Zn-IMAC column, along with a second putative outer membrane protein (SYNW2227). Both proteins belong to the family of cyanobacterial porins (CBP; TC 1.B.23). Two homologues from the freshwater strain *Synechococcus* sp. PCC7942 (SYNPC7942 1464 and SYNPC7942 1463; termed SomA and SomB, for *Synechococcus* outer-membrane protein A and B) have been extensively studied.\(^{82,83}\) In contrast to carbohydrate-translocating porins from heterotrophic bacteria, the single-channel conductance of SomA suggested that small solutes, most likely ions, are transported,\(^{44}\) but the actual substrates have not been identified.

To enable the inspection of SYNW2224 for potential metal-binding sites, we generated a coarse model with the aid of the Phyre2 server (Fig. 5). The model shows a 16-stranded \(\beta\)-barrel, an N-terminal S-layer homology domain, connected by a long amphipathic helix. The S-layer domain, the helix, as well as several loops (both extra- and intracellular) abound in carboxylic groups, which might provide metal ion interaction sites, although it should be noted that the number of positively and negatively charged side-chains in SYNW2224 is overall similar. Many other marine CBPs (Table S3, ESIF) have a pronounced over-abundance of negatively charged side-chains. The model also suggests that the ‘pore’ is probably constricted by several loops, consistent with the hypothesis that these porins transport simple inorganic ions.

**Discussion**

The ability of *Synechococcus* sp. WH8102 cultures to grow at extremely low zinc concentrations and maintain cellular zinc homeostasis, as judged by total zinc quotas, suggests that *Synechococcus* sp. WH8102 has extremely efficient zinc uptake mechanisms capable of scavenging zinc even from depleted media. Only slight, if any, growth limitation at extremely low free zinc concentrations has previously been observed for *Synechococcus bacillaris*\(^{23}\) and *Prochlorococcus* MED4,\(^{24}\) especially when cobalt concentrations were also low, pointing to potential co-limitation by these two metal ions, and to possible substitution of zinc by cobalt but not \emph{vice versa}.\(^{23,24}\)

The mechanisms for metal ion uptake in marine cyanobacteria are only partially understood. Genes for DNA-binding metal sensor proteins of the Fur family are present in all available genomes,\(^{29}\) as are those for ABC transporters for Fe, Mn, and Zn.\(^{26,30}\) Little is known regarding transport across the outer membrane, but considering that metal ions such as Fe\(^{3+}\) and Zn\(^{2+}\) must be bio-accumulated by a factor of at least \(10^5\), it follows that uptake mechanisms, including transport across the outer membrane, must be highly efficient.

One principal possibility is that zinc is transported into the cell as a complex with a chelating ligand. This option typically involves TonB-dependent receptors,\(^{56}\) which actively (\emph{i.e.} under consumption of ATP) transport metal complexes with organic ligands across the outer membrane. TonB-dependent receptors are widely involved in bacterial iron uptake, but have also been implicated in bacterial zinc uptake, including four examples in *Pseudomonas protegens* PF-5 in zinc-depleted soil,\(^{87}\) two examples
in the opportunistic pathogen *Acinetobacter baumannii*,88 and a protein named “ZnuD” produced by pathogenic *Neisseria meningitidis* under zinc limitation.89 In the freshwater cyanobacterium *Anabaena* sp. PCC 7120, zinc starvation caused the upregulation of a Zur-regulated TonB-like receptor.90 However, even though TonB-like genes have been discovered in *Prochlorococcus*,90 most marine *Synechococcus* strains lack the respective proteins.30,31,91,92 This absence of TonB-dependent receptors means that no specific outer membrane transporters for any M+ or M++ metal ion, including Fe++, are known for marine *Synechococcus*. Thus, our discovery that the putative porins SYNW2224 and SYNW2227 have metal-binding ability offers a new hypothesis regarding the uptake of scarce trace metal ions by *Synechococcus* from oligotrophic waters, namely that at least some cyanobacterial porins of the CBP family play a central role in this process. The SYNW2224 model generated (Fig. S) does not reveal any clearly defined binding sites, but many surface-accessible negatively charged carboxylate residues. On the basis of the nature and distribution of these potential metal-binding residues, the specificity of CBPs is expected to be limited, since all M+ or M++ species have significant affinities for clusters of carboxylate groups. Indeed, SYNW2224 was also captured on a Co2+-IMAC column in our previous study.37 The S-layer homology domain portion, and the amphipathic helix of SYNW2224, are also rich in negative charges, and hence may contribute to the permeability of the outer membrane for enhanced nitrogen and carbon uptake,102 or that they might transport phosphate.103 Cation- and anion-selective porins from the same organism, *e.g.* OmpF and PhoE from *E. coli*, share a high degree of similarity. Without direct biophysical studies, it is not possible to assign or predict particular selectivities for CBPs, but it may be expected that separate membrane channels for cationic and anionic nutrients should exist.106 If SYNW2224 proves to function in metal transport, then its increased expression under P-limitation105 could be an indirect consequence of the increased requirement for metal-dependent alkaline phosphatases, which are also up-regulated under these conditions.103 The increased expression of these enzymes would likely also increase cellular demand for zinc and/or calcium.107,108 Upregulation of the definitively zinc-related metallothionein SmtA in response to P-limitation in *Synechococcus* sp. WH8102 has recently been observed,25 indicating that Zn and P metabolic processes are linked in this strain. A more general link between Zn and P metabolism is also reflected in global biogeochemical cycling.109 Regarding the transported species, for eukaryotic phytoplankton zinc uptake, the free ion concentration has been thought to determine bioavailability,51 but a recent study demonstrated that weak organic ligands in the presence of a much stronger ligand such as EDTA increased the rate of zinc uptake by zinc-limited cultures of *Emiliania huxleyi* and *Thalassiosira weissflogii*, via a mechanism that likely involves the formation of ternary complexes between the weak metal–ligand complexes and a cell surface uptake molecule.110

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Evidence that porins may function in metal uptake is available at the transcriptional level for other bacteria; for example, the expression of the porin OmpT in *Vibrio cholerae* is dependent on the level of iron in the environment, and is positively regulated by the ferric uptake regulator, Fur.95 The outer-membrane protein MnoP in *Bradyrhizobium japonicum* is expressed under conditions of manganese limitation, and is required for high-affinity manganese uptake.96 In *Mycobacterium tuberculosis* the porin MsPA has been shown to be required for copper uptake across the outer cell membrane, with msPA deletion mutants showing severe growth defects when grown in a trace copper medium.97 The expression of several porins was also zinc-dependent in *Pseudomonas protegens*,87 and that of the OprD porin in *Pseudomonas aeruginosa* is down-regulated by excess zinc.98 Significantly, in freshwater *Synechococcus* sp. PCC7942, the gene for the CBP somB, but not its neighbouring CBP gene somA, has been found to be up-regulated under iron starvation.99 somA and somB do not form an operon, but each have their own transcription start sites.100 Importantly, the upstream region of the somB gene in the PCC7942 strain is predicted by RegPrecise101 to contain a recognition site for the zinc-uptake regulator protein Zur. Hence, its expression is likely also zinc-regulated. RegPrecise also predicts zur boxes for CBP homologues in *Thermosynechococcus elongatus* (tbl246), *Microcystis aeruginosa* (MAE_10010), and *Synechocystis* sp. PCC6803 (sll1550), and therefore proposed the name OmpZ (for zinc-regulated outer-membrane protein) for these homologues. Most cyanobacterial strains, including marine strains, have multiple CBP genes (Table S3, ES1), with *Synechococcus* sp. WH8102 having at least four (synw2128, synw2223, synw2224, synw2227). The most divergent cyanobacterial strain, *Gloeobacter violaceus*, also harbours six CBP genes; four of them with and two without an SLH domain, indicating that duplication and divergence of these porins occurred even before the evolution of chloroplasts. *Prochlorococcus* sp. CCMP1375 (SS120), a strain with one of the smallest genomes, contains only two CBP genes, but there is no simple relationship between genome size and the number of CBP genes. Inspection of the genomic neighbourhoods of these genes (Table S3, ES1) reveals that they are frequently localised in the vicinity of genes for the periplasmic binding proteins for Zn, Fe, or phosphate, or associated with genes suggesting a relationship with the metabolism of other metal ions, including Mn, Co and Ni. Whilst analyses of genomic neighbourhoods, or regulation by a particular nutrient cannot directly infer function, we suggest that our demonstrated zinc-binding ability of a CBP, and the high abundance of SYNW2224 under zinc-depleted conditions, adds two strong pieces of experimental evidence towards at least some of these porins playing an important role in the transport of essential metal ions such as zinc.

CBPs have previously been found to be up-regulated under nitrogen or sulfur starvation conditions in *Synechococcus* sp. PCC7942,102 and under phosphorus starvation in *Synechococcus* sp. WH8102,25,103,104 and *Prochlorococcus* spp. MED4 and MIT9313.105 This has led to suggestions that these proteins might increase the permeability of the outer membrane for enhanced nitrogen and sulfur uptake,102 or that they might transport phosphate.103 Cation- and anion-selective porins from the same organism, *e.g.* OmpF and PhoE from *E. coli*, share a high degree of similarity. Without direct biophysical studies, it is not possible to assign or predict particular selectivities for CBPs, but it may be expected that separate membrane channels for cationic and anionic nutrients should exist.106 If SYNW2224 proves to function in metal transport, then its increased expression under P-limitation105--107 could be an indirect consequence of the increased requirement for metal-dependent alkaline phosphatases, which are also up-regulated under these conditions.103 The increased expression of these enzymes would likely also increase cellular demand for zinc and/or calcium.107,108 Upregulation of the definitively zinc-related metallothionein SmtA in response to P-limitation in *Synechococcus* sp. WH8102 has recently been observed,25 indicating that Zn and P metabolic processes are linked in this strain. A more general link between Zn and P metabolism is also reflected in global biogeochemical cycling.109
Such a mechanism is also conceivable for zinc uptake by CBPs. SYNW2224 and SYNW2227 clearly have the ability to interact with partially complexed \( \text{Zn}^{2+} \), as presented by a nitrilotriacetic acid-based IMAC column. The conductance data characteristic of small solutes found for porins closely similar to SYNW2224 and SYNW2227 suggests that free \( \text{Zn}^{2+} \) (and likely other metal ions), or complexes with water, chloride, and other small anions, may be the entities transported, hence transport through the outer membrane could involve a decomplexation step.

The periplasm could then act as the first sorting point for different ions. Four periplasmic binding proteins were captured by Zn-IMAC, only two of them predicted to bind metal ions. Besides the periplasmic ZnuA component of a predicted zinc ABC transporter (SYNW2481), the \( \text{Fe}^{3+} \)-binding protein FutA (SYNW1797) was also identified. The genome of *Synechococcus* sp. WH8102 contains an additional putative *znuABC* gene cluster (SYNW0969, 0970, and 0971) which according to RegPrecise harbours a predicted Zur recognition sequence in its promoter region. The periplasmic component of this cluster is *symw0971*; the respective protein was not detected in our metalloproteomics study.

The phosphate-binding protein PstS (SYNW1018) and the cyanate-binding protein CynA (SYNW2487) were also captured by Zn-IMAC. Whilst the observed metal affinity of PstS remains enigmatic, a very obvious metal-binding site ideally suited to bind \( \text{Zn}^{2+} \) (or related ions) was detected in CynA. Considering the scarcity of \( \text{Zn}^{2+} \) (and related ions such as \( \text{Co}^{2+} \) and \( \text{Cd}^{2+} \)) in the natural habitat of *Synechococcus* sp. WH8102, it appears counter-intuitive that the bacterium should “waste” a potentially catalytically active metal ion for a merely supporting role in transport of an unusual nitrogen source. It is noteworthy that *Synechococcus* sp. WH8102 is one of the few strains that contains an active uptake system for cyanate, and is also able to utilise cyanate as sole nitrogen source. Several *Prochlorococcus* strains also have this capability. Transcripts for cyanA (likely most closely related to HLII types) were abundant in stratified surface waters of the Gulf of Aqaba. These waters are characterised by nitrogen depletion, but also by high Prochlorococcus abundance. It has been suggested that the ability to utilise an additional nitrogen source may give a competitive advantage in extremely N-depleted surface layers of stratified ocean waters. It is intriguing that this may involve the participation of zinc or a closely related metal.

**Conclusions**

Our understanding of the mechanisms of zinc uptake into Gram-negative bacteria is incomplete, even though these are of particular interest for environments in which the concentration of essential zinc is extremely low, for example in oligotrophic oceans, but also during the acute phase response of a mammalian host to bacterial infection.

This study has demonstrated the increased expression of a putative cyanobacterial porin (CBP) under conditions of zinc depletion, and its zinc-binding ability – a property not previously demonstrated for any bacterial porin. Together with bioinformatic evidence for likely regulation of CBP homologues by the zinc uptake regulator Zur, this suggests a role for at least some of these proteins in \( \text{Zn}^{2+} \) uptake across the outer membrane of cyanobacteria, although the transport of other...
inorganic cations is also likely. CBPs have previously been implicated in the uptake of carbon, nitrogen, and phosphorus. This work has shown that their expression may not only be regulated by lack of macronutrients, but also of micronutrients.

Furthermore, the isolation of a putative periplasmic zinc-binding protein, ZnUA, by Zn²⁺-IMAC has provided the first experimental evidence that this protein is expressed by *Synechococcus* sp. WH8102, and has the ability to bind zinc in vitro. Together with the finding that *Synechococcus* sp. WH8102 accumulates appreciable quantities of zinc even under extreme zinc depletion, and further bioinformatic information, this augments our understanding of zinc homeostasis in this and other marine cyanobacterial strains (Fig. 6). In addition, combining Zn-IMAC with comparative modelling has led to the discovery of a novel metal-binding site in the periplasmic cyanate-binding protein CynA that enables *Synechococcus* sp. WH8102 to exploit cyanate as nitrogen source.

Although no intact zinc-bound enzymes or other proteins were identified from any of the fractions analysed by native liquid chromatography, the alternative approach, IMAC, has enabled the enrichment of at least one protein involved in zinc uptake, as well as several candidates that warrant further detailed biophysical studies, including the porins SYN2224 and 2227, the carboxysomal proteins CcmK1 and CcmK2, and the periplasmic cyanate binding protein CynA. It should be emphasised again that binding to a particular IMAC column does not allow any firm conclusions regarding any in vivo binding partners. It is possible that in several instances, the observed interaction with Zn²⁺ corresponds to adventitious binding. Careful inspection of each candidate, as a minimum via homology modelling and metal site analysis, should be carried out to eliminate false positives. Even then, caution must be applied regarding any statements pertaining to metal specificity, and further meta-data (such as genomic neighbourhood, synteny, and presence of transcription factor recognition sites) must be considered. Taking all these caveats into consideration, the experimental metalloproteomics in conjunction with bioinformatic approaches employed in the present work have uncovered expected and unexpected players in the zinc-binding network of a representative of an environmentally important class of marine phytoplankton.

**Acknowledgements**

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