Comparative Phenotypic Analysis of Anabaena sp. PCC 7120 Mutants of Porin-like Genes

Hannah Schätzle1,2,3, Eva-Maria Brouwer1, Elisa Liebhart1,2, Mara Stevanovic1, and Enrico Schleiff1,2,3,4*

1Institute for Molecular Biosciences, Goethe University, Frankfurt am Main, Germany
2FIERCE, Goethe University, Frankfurt am Main, Germany
3Buchmann Institute for Molecular Life Sciences, Goethe University, Frankfurt am Main, Germany
4Frankfurt Institute of Advanced Studies, Frankfurt am Main, Germany

Porins are essential for the viability of Gram-negative bacteria. They ensure the uptake of nutrients, can be involved in the maintenance of outer membrane integrity and define the antibiotic or drug resistance of organisms. The function and structure of porins in proteobacteria is well described, while their function in photoautotrophic cyanobacteria has not been systematically explored. We compared the domain architecture of nine putative porins in the filamentous cyanobacterium Anabaena sp. PCC 7120 and analyzed the seven candidates with predicted OprB-domain. Single recombinant mutants of the seven genes were created and their growth capacity under different conditions was analyzed. Most of the putative porins seem to be involved in the transport of salt and copper, as respective mutants were resistant to elevated concentrations of these substances. In turn, only the mutant of alr2231 was less sensitive to elevated zinc concentrations, while mutants of alr0834, alr4741 and alr4499 were resistant to high manganese concentrations. Notably the mutant of alr4550 shows a high sensitivity against harmful compounds, which is indicative for a function related to the maintenance of outer membrane integrity. Moreover, the mutant of all5191 exhibited a phenotype which suggests either a higher nitrate demand or an inefficient nitrogen fixation. The dependency of porin membrane insertion on Omp85 proteins was tested exemplarily for Alr4550, and an enhanced aggregation of Alr4550 was observed in two omp85 mutants. The comparative analysis of porin mutants suggests that the proteins in parts perform distinct functions related to envelope integrity and solute uptake.

Keywords: Cyanobacteria, β-barrel proteins, Omp85 function, outer membrane biogenesis, porins

Introduction

Cyanobacteria are Gram-negative bacteria, as they possess an outer membrane (OM) that acts as diffusion barrier, a peptidoglycan mesh (PG) and a plasma membrane (PM). The cyanobacterial envelope in certain aspects differs from that of other Gram-negative heterotrophs [1]. Exemplarily the PG layer in filamentous Anabaena sp. strain PCC 7120 (Anabaena sp.) is approximately 14 nm thick, and the distance between PM and OM is about 45 nm [2]. Here, the outer membrane continuously surrounds the whole filament by not penetrating into the septum area between two cells [2]. Compared to that the PG of E. coli is approximately 6 nm thick and the distance between PM and OM is in the range of 20 nm [3]. The denoted dimensions of the cell envelope might be even disparate within certain cyanobacterial species, as some multicellular cyanobacteria species possess different cell types with distinct cell envelope properties.

Membrane proteins regulate transport processes across membranes and define the susceptibility of an organism against harmful compounds and antibiotics. A protein class that is highly abundant in the OM are porins. Porins are membrane embedded β-barrel proteins that allow diffusion of substrates with low or high specificities (reviewed e.g. in [4, 5]). Their functions are not only related to diffusion and solute transport, but porins are also crucial for outer membrane integrity including antibiotic resistance [6] or pathogenesis [4]. Porins that lack specificity are termed general or non-specific porins. They typically consist of 16β-strands and facilitate diffusion of small hydrophilic compounds [7]. In addition, substrate specific porins exist, typically composed of 18β-strands [8]. Porins with less than 16β-strands are described as well, for example the monomeric OmpG from Escherichia coli is composed of 14β-strands [9]. Although monomeric porins have been reported, porins most often occur as trimers [5]. Porins and other outer membrane β-barrel proteins are integrated into the OM by the β-
barrel assembly machinery [10]. An Omp85 protein constitutes the main pore of this protein complex. The N-terminal part of the Omp85 protein bears polypeptide transport-associated domains (POTRA) that recognize the non-folded membrane proteins. Moreover, the POTRA domains interact with periplasmic chaperones or other proteins involved in the insertion and folding process [11].

The freshwater cyanobacterium Anabaena sp. is a model organism with regard to bacterial cell differentiation, as in the absence of a combined nitrogen source specialized cells called heterocysts are formed. Heterocysts are morphologically distinct from vegetative cells, including the structure of the cell envelope [12]. They are surrounded by a polysaccharide layer that mechanically protects the underneath glycolipid layer. The prevalence of certain proteins in the outer membrane only mildly differs between vegetative cells and heterocysts [13]. Information on cyanobacterial porins is rather limited compared to proteobacteria. Cyanobacterial porins were found to be relatively large with about 30–50 kDa per monomer, whereas most proteobacterial porin monomers have a molecular weight lower than 40 kDa [14-16]. The enlargement results from the presence of an additional cyanobacteria specific N-terminal domain [17]. This domain is related to the conserved surface layer homology (SLH) domain, which is involved in targeting and linking proteins to cell-wall associated components [18-21].

Examinations of SomA and SomB, two out of six porin-like proteins in the unicellular cyanobacterium Synechocystis sp. strain PCC 6803 (Synechocystis sp.), indicated that the permeability of organic compounds is relatively low (0.4 and 0.9 nS) [17]. Hence, it is hypothesized that Synechocystis sp. lacks classical porins. The major OM proteins did not allow diffusion of organic substances, whereas inorganic molecules could penetrate the pores [22]. Thus the overall permeability of the Synechocystis sp. OM was around 20-fold lower than that of E. coli. The authors claim that this might be a consequence of the photoautotrophic lifestyle of Synechocystis sp., which renders an import of sugars dispensable. Cyanobacteria that are living in symbiosis with plants seem to represent an exception, as the photosynthetic activity in those organisms is often diminished. In a symbiotic species of the cyanobacterium Nostoc muscorum the sugar-specific porin Omp25 was found to be required for proper uptake of sugars [23]. Notably an OprB-type porin together with the Omp85 protein and LptD, a protein that is involved in the transfer of lipid A, was found to be globally conserved among cyanobacterial OM proteins [24]. However, there is no indication whether those porins are truly carbohydrate-selective or not. Recently it was described that in Synechocystis sp. the porin Slr1908 mediates the transport of inorganic iron [25]. Slr1908-like sequences were identified in many cyanobacterial species, indicating a porin-dependent iron uptake in cyanobacteria.

Nine porin-like genes were assigned based on sequence alignments and amino acid sequence properties in the genome of Anabaena sp. [26-28], and previous analyses showed that porins might be involved in ethidium bromide and presumably erythromycin uptake in Anabaena sp. [29]. The central Omp85 proteins of Anabaena sp. are encoded by abr0075, alr2269 and alr4893. Proteomic analyses revealed that Alr2269 is the most abundant among the three Omp85 proteins, whereas Alr4550, Alr4499 and Alr3608 are the most abundant among the putative porins [28].

Here, we comparatively characterize the seven genes coding porin-like proteins in Anabaena sp. by analyzing the transcript abundance of the genes under standard and starvation conditions. Although the function of these protein has not been biochemically characterized, for simplification and easier reading we subsequently refer to the proteins as "porins" based on the sequence and motif similarity to characterized proteins in proteobacteria. Moreover, mutants of the single porin-like genes were phenotypically analyzed in presence of high metal, salt and drug concentrations. The results indicate that the highly abundant porin Alr4550 plays an important structural role in Anabaena sp., since the corresponding mutant was strongly defected in OM integrity. Further, a function in the transport of manganese, zinc or cobalt is suggested for specific porins. Interestingly a mutant of alls191 was altered in growth under diazotrophic conditions, which suggests an insufficient nitrogen fixation capacity or an enhanced nitrogen demand of the strain. Moreover, we show that Alr2269 and Alr4893 affect the membrane altered in growth under diazotrophic conditions, which suggests an insufficient nitrogen fixation capacity or an enhanced nitrogen demand of the strain. Moreover, we show that Alr2269 and Alr4893 affect the membrane

Materials and Methods

Bioinformatics

Sequences and sequence information were extracted from NCBI database [30]. Logo plots were created using the WebLogo online tool (http://weblogo.berkeley.edu/logo.cgi, [31]). The sequence alignment for determination of the sequence identities was performed with Clustal Omega [32, 33].

Generation of Anabaena sp. PCC 7120 Mutants

An internal fragment of the gene of interest was amplified by PCR with gene specific oligonucleotides (Table S1). BglII-sites were introduced at 5' and 3'-ends of the fragment. This fragment was inserted into BamHI-digested pCSV3 carrying a Cs.3 cassette (kind gift from Prof. E. Flores, [34, 35]) yielding the final plasmid for conjugation (Table S2). Single-recombinant insertion mutants of Anabaena sp. were created with the triparental mating method [36-38] utilizing E. coli strains HB101 (gets transformed with the plasmid of interest) and ED8654 (carries the conjugative plasmid). In short the two E. coli strains were mixed with Anabaena sp. wild type, and the plasmid of interest bearing a homolog region to the gene of interest is transferred to Anabaena sp. by conjugation. A single recombination event happens where the whole plasmid bearing the homolog region gets integrated into the genome, resulting in cells that are resistant towards spectinomycin and streptomycin.

The genotype of the exconjugants was tested by PCR utilizing oligonucleotides that anneal outside of the internal homologous fragment in combination with a vector-specific oligonucleotide (Table S1). The mutant strains are listed in Table S3.
Phenotypic Analysis of the Porin Mutants

Anabaena sp. strains were stored on BG11 plates [39] with 1% (w/v) bacto-agar (BD Biosciences) until use. For expression analysis by qRT-PCR Anabaena sp. wild type and mutants were grown in buffered liquid YBG11 medium [40]. In case of mutants the medium was supplemented with 5 μg ml⁻¹ spectinomycin dihydrochloride pentahydrate (Duchefa Biochemie) and streptomycin sulfate (Roth).

For the phenotyping on plates, cultures were washed and suspended at a final concentration OD₅₇₀nm = 1.5 μl of the non-diluted suspension and of a 1:10 dilution was spotted onto YBG11 plates containing given amounts of supplements or on YBG11 plates (YBG11: YBG11 medium without NaNO₃). Plates were incubated under constant illumination (70 μmol photons m⁻² s⁻¹) at 28-29°C for 7 days. Each spotting assay was repeated with independent cultures at least three times. Representative images are shown in the results. All strains depicted in one line were grown on the same plate.

Microscopy

Light microscope images were taken with Olympus CXX41 using a 40x objective and a Thorlabs DCC1645C-HQ camera. Heterocysts were stained with alcian blue to improve visibility. Anabaena sp. suspension was mixed in a 1:1 ratio with the alcian blue staining solution (0.5% alcian blue in 50% ethanol, [41]) and incubated for 5 min. After three washing steps with YBG11 medium (centrifugation at 1500 g, 5 min) filaments were inspected under the microscope.

RNA Isolation and qRT-PCR Analysis

RNA isolation, cDNA synthesis and qRT-PCR with the corresponding oligonucleotides (Table S1) was performed as previously described [42]. Three independent wild-type and mutant cultures grown for 5 days were used for RNA isolation, while for starvation experiments three week-old cultures were used. qRT-PCR was performed on cDNA from three biological replicates. The Ct values for the genes were normalized to Ct values of the rnpB transcript in the corresponding sample, yielding the ΔCt. For the calculation of the ΔΔCt value the ΔCt for each sample was normalized to the ΔCt of wild type grown under control conditions [43].

Membrane Isolation and Rate Zonal Centrifugation

OM from Anabaena sp. was isolated as described [30] with slight modifications. Anabaena sp. cultures were grown in BG11 to exponential growth phase, 500 ml were harvested and washed with 5 mM 4-(2-hydroxyethyl)-1-piperazinethanesulfonic acid (HEPES), pH 7.6 supplemented with 1 mM phenylmethylsulfonyl fluoride (PMSF). Cell lysis and harvesting of membrane fractions was performed as described [30]. The membrane fraction was resuspended in 30 % sucrose containing 20 mM HEPES, pH 7.6 and 0.2 mM PMSF. It was layered on top of a 55% (w/v) sucrose cushion. The OM was sedimented by centrifugation (130,000 x g, 16 h, 4°C). The OM pellet was washed with 20 mM HEPES, pH 7.6 supplemented with 0.2 mM PMSF and collected by centrifugation (130,000 x g, 1 h, 4°C). Membranes were resuspended in 20 mM HEPES, pH 7.6 with 0.2 mM PMSF and stored at -80°C.

100 μg protein of the membrane fraction was loaded on top of a linear sucrose gradient (10 to 70%; w/v) and centrifuged (100,000 x g, 1 h, 4°C). The gradient was fractionated into ten fractions of 1 ml. Proteins were precipitated with 0.02% (w/v) Na-deoxycholate and 15% (v/v) trichloroacetic acid. Precipitates were resolved in 6-fold SDS-Urea loading buffer containing 200 mM Tris HCl pH 6.8; 8M Urea; 0.1 mM ethylenediaminetetraacetic acid (EDTA); 5% sodium dodecyl sulfate (SDS) and 0.03% bromophenol blue and subjected to SDS-PAGE followed by immunoblotting. Blotting membranes were stained with direct blue 71 (DB71) as described [44].

For total protein extract 1 ml of exponential phase culture was harvested by centrifugation (10,000 x g, 1 h, 4°C). Membranes were resuspended in 20 mM HEPES, pH 7.6 with 0.2 mM PMSF and stored at -80°C.

100 μg protein of the membrane fraction was loaded on top of a linear sucrose gradient (10 to 70%; w/v) and centrifuged (100,000 x g, 16 h, 4°C). The gradient was fractionated into ten fractions of 1 ml. Proteins were precipitated with 0.02% (w/v) Na-deoxycholate and 15% (v/v) trichloroacetic acid. Precipitates were resolved in 6-fold SDS-Urea loading buffer containing 200 mM Tris HCl pH 6.8; 8M Urea; 0.1 mM ethylenediaminetetraacetic acid (EDTA); 5% sodium dodecyl sulfate (SDS) and 0.03% bromophenol blue and subjected to SDS-PAGE followed by immunoblotting. Blotting membranes were stained with direct blue 71 (DB71) as described [44].

For total protein extract 1 ml of exponential phase culture was harvested by centrifugation (10,000 x g, 5 min) and the pellet was resuspended in 6-fold SDS-Urea loading buffer (recipe given above). Samples were incubated at 42°C for 5 min and centrifuged afterwards to pellet insoluble cell debris, the supernatant was used for SDS PAGE. Antibodies against the peptides specific for Alr4450 were generated by Peptide Specialty Laboratories (Germany) and previously described [45], the following peptides were used: Peptide 1: ASGQGLQTFQVGSTGNNC, Peptide 2: PITEDTKVQDQVNRYSNENKGNAQ. Antibodies against Alr2269 and Tic22 have also been previously described [30,46].

Inductively Coupled Plasma Mass Spectrometry (ICP-MS)

The intracellular metal concentration of Anabaena sp. wild type grown for seven days in YBG11 medium was analysed. The procedure was adapted [47] with modifications. The cells were sedimented by centrifugation (3,000 x g, 16 h, 4°C). The OM was washed twice with 20 mM 2-(N-morpholino)ethanesulfonic acid (pH 5) and 20 mM EDTA. The final pellet was resuspended in double-distilled water (ddH₂O). For normalization the OD₅₇₀ was determined and the cells were counted using a Helber bacteria counting chamber (Hawksley). From here on experiments were conducted in a metal clean laboratory. 1 ml of each sample was incubated at 120°C in 7 M HNO₃ overnight until dryness. Before measurement, the samples were resolved in 5% HNO₃. As controls, samples of ddH₂O and cultivation medium were analysed. Glassware used during the experiment was incubated in 4% HNO₃ overnight prior to use.

Results

Seven Proteins in Anabaena sp. Contain a Porin-like Domain Architecture

In previous studies, the nine putative porin-candidates alr0834, alr2231, all4499, alr4550, alr4741, all5191, all7614, alr3917 and alr3608 were identified in the Anabaena sp. genome [30, 48]. In addition, All3289 and
Alr5049 were assigned as OmpA-like proteins [48]. OmpA-type porins are thought to perform structural functions related to membrane integrity [8, 49]. However, Alr5049 codes for a protein with 1289 amino acids. The C-terminal region might form an OM anchor for the large soluble domain that contains characteristics of glycoproteins. Hence, we suggest that this protein should not be considered as typical OmpA. In turn, Alr3289 codes for a protein with 169 amino acids lacking the characteristic domains of an OmpA [48]. Thus, based on the small size, the absence of domains characteristic for OmpA proteins and considering that the outer membrane localization can be confirmed in future, the protein might represent a functional OmpX [50].

Seven of the nine putative porins (Alr0834, Alr2231, Alr4499, Alr4550, Alr4741, All7614, Alr3608) contain an S-layer homology domain (SLH) which is also found in other cyanobacterial porins [19]. In Alr3608, the two predicted SLH-domains are located in the C-terminal region, whereas in other porins they are found in the N-terminal part of the protein (Fig. 1A). Moreover, seven of the putative porin proteins contain an OPR-B domain that is characteristic for carbohydrate-selective porins (Fig. 1A). This domain could not be identified in Alr3608 and Alr3917. In turn, a domain of unknown function (DUF) and a short region characteristic for a DNA translocase FtsK were identified in Alr3608 and Alr3917, respectively. However, a BLAST search with the Alr3917 sequence against cyanobacterial genomes in the NCBI-database did not yield a similarity to cyanobacterial FtsK sequences. Notably, Anabaena sp. FtsK is encoded by all7666.

The comparison of the amino acid sequence of the last β-strand in the seven porins with OPR-B fold shows a high degree of conservation, while Alr3608 and Alr3917 are somewhat distinct (Fig. 1B). It is proposed that the most C-terminal region of proteobacterial outer membrane proteins with β-barrel fold contains the signal for membrane insertion and Omp85 interaction [51-54]. In addition, the so called β-signal initiates the association with Omp85 (BamA; [55, 56]). The comparison of the identified motif to that of the 22 TonB-dependent transporters (TBDT) in the outer membrane uncovers a common motif in the last β-strand: hydrophobic/small amino acid (Ψ) – x – hydroxylated amino acids (Φ/η) – x – aromatic amino acids (Ω) – x – Ω. This motif might be important for the insertion of the proteins into the outer membrane, as it was reported that an aromatic amino acid at the last position of the sequence is important for proteobacterial outer membrane protein insertion [53]. In addition, the insertion of mitochondrial β-barrel proteins depends on a hydrophobic signal in the last strand as well [57]. Comparison of the amino acid sequence of the seven porin-like proteins to the amino acid sequence of the two most abundant E. coli porins, namely OmpC and OmpF [5], yielded a sequence identity between 18.8% to 24.6% (Fig. 1C). This indicates that the cyanobacterial proteins do not share a high sequence similarity to proteobacterial porins.

In summary, the bioinformatics inspection revealed that seven out of the nine predicted porins in Anabaena sp. contain a domain architecture consistent with a porin-like function, although no secretion signal was predicted for Alr4741. The other two proteins previously assigned to the porin-like family (Alr3608, Alr3917) contain a domain architecture that is rather atypical for porins. As it is questionable whether these two proteins constitute porins, they were excluded from the subsequent analyses.

### Genotyping of Porin Mutants and Growth under Standard Conditions

In order to assess specific mutant phenotypes, single-recombinant mutants were created by integration of a plasmid bearing the CS.3 cassette into the porin genes. Five mutants had the plasmid insertion in forward
direction, while the plasmid was inserted in the opposite direction into alr0834 and all4499 loci (Fig. 2A). Mutants of all4499 and alr2231 were not segregated, as wild-type copies of the genes were present after repeatedly diluting the cultures on medium containing antibiotics (Fig. 2A, first lane). This might indicate an essential function of the two gene products under the given conditions. For all other genes, segregated mutants were obtained.

The growth behavior of the porin mutants was compared to wild type. In addition, mutants exhibiting alterations in outer membrane integrity, bearing either an insertion in Anabaena sp. omp85 genes (AFS-I-alr0075, AFS-I-alr2269, and AFS-I-alr4893; AFS stands for Anabaena mutant created in Frankfurt by the Schleiff group) [58] or the tic22 gene (AFS-I-tic22, alr0114, [45, 46, 59]) were used as additional controls. As reported before, AFS-I-alr0075, AFS-I-alr2269 and AFS-I-alr4893 could not be segregated since wild-type copies could be amplified even after exposing the strains to increased concentrations of antibiotics [58] (Fig. S1).

On standard YBG11 plates only AFS-I-alr0075 and AFS-I-tic22 grew to a lower density compared to wild type (Fig. 2B, first and second panel). This behavior was not described before [45, 46, 59] as the effect was only visible when the culture was sufficiently diluted (Fig. 2B, second panel). However, the strains with plasmid insertion in the porin-like genes did not show any phenotype under this condition (Fig. 2B).

The Mutants of Three Porin-Like Genes Are Affected in Growth in Absence of Combined Nitrogen

On plates without a combined nitrogen source (YBG11$_0$), a condition where Anabaena sp. fixes atmospheric nitrogen in heterocysts, AFS-I-alr2269 and AFS-I-tic22, but not AFS-I-alr0075 and AFS-I-alr4893 were impaired in growth (Fig. 2B, third panel) as previously reported [46]. With respect to the with respect to the porin mutants, AFS-I-all5191 grew to a lower density than wild type on YBG11$_0$ plates (Fig. 2B, second panel). In addition, AFS-I-alr4741 and AFS-I-alr0834 grew to slightly higher densities under diazotrophic conditions compared to wild type.

Consequently, filaments of wild type, AFS-I-alr0834, AFS-I-alr4741 and AFS-I-all5191 grown in YBG11 or YBG11$_0$ were morphologically inspected. In standard YBG11 medium the filaments of all strains appeared morphologically comparable to wild-type cells (Fig. 3, +NO$_3$). Heterocysts were not detected in the mutant and wild-type strains grown in the presence of nitrate. After a seven-day cultivation in YBG11$_0$, medium heterocysts were visible in wild type, AFS-I-alr0834, AFS-I-alr4741 and AFS-I-all5191 (Fig. 3, -NO$_3$). Thus the defective growth of AFS-I-all5191 is not due to an inability of this strain to differentiate heterocysts. Though, since in general the cells of AFS-I-all5191 looked comparatively pale in the absence of nitrate (Fig. 2B), we assume that
Most of the Genes Coding for Porin-like Proteins are Upregulated by Metal Starvation

The regulation of porin-mediated solute diffusion in bacteria is governed by distinct parameters. For instance, the extracellular nutrient supply status and the osmolarity might trigger transcriptional responses [60]. *Anabaena* sp. performs photosynthesis and is therefore not obligatorily dependent on sugar uptake. Thus, the potential changes in gene expression of porins in response to altered metal concentrations in the growth medium were examined. For this, *Anabaena* sp. wild type was grown in YBG11 with modified metal concentrations by omitting either manganese (Mn), iron (Fe), zinc (Zn) or copper (Cu) from the medium. These metals were used since *Anabaena* sp. accumulates (5.0 ± 0.3) × 10¹⁴ atoms Mn in cells equivalent to 1 ml culture at OD₇₅₀ = 1, (0.5 ± 0.05) × 10¹⁴ atoms Cu/1 ml (OD₇₅₀ = 1) and (0.5 ± 0.03) × 10¹⁴ atoms Zn/1 ml (OD₇₅₀ = 1) as determined by ICP-MS. By that, the cellular concentration of these metals is in the similar range to that of iron ((3.4 ± 0.2) × 10¹⁴ atoms Fe/1 ml (OD₇₅₀ = 1)), which is considered to be a limiting factor for cyanobacterial growth and is especially important for N₂-fixing cyanobacteria [61, 62].

Thus, RNA was isolated from wild-type cells grown for 21 days in the indicated media (Table 1) and qRT-PCR

**Table 1. qRT-PCR analysis of expression of porin genes.**

| *Anabaena* sp. PCC 7120 | YBG11 | YBG11ₚ₃ | YBG11ₚ₄ | YBG11ₚ₅ | YBG11ₚ₆ |
|--------------------------|-------|---------|---------|---------|---------|
| **-ΔCt**                 | -ΔΔCt | -ΔΔCt   | -ΔΔCt   | -ΔΔCt   | -ΔΔCt   |
| *alr0834*                | -10.8 ± 0.3 | 1.7 ± 0.1 | 1.6 ± 0.1 | 0.1 ± 0.2 | 0.8 ± 0.1 |
| *alr2231*                | -12.3 ± 0.3 | 1.9 ± 0.1 | 0.0 ± 0.1 | 0.5 ± 0.3 | 0.9 ± 0.1 |
| *all4499*                | -9.4 ± 0.2 | 0.5 ± 0.1 | 1.4 ± 0.2 | -0.6 ± 0.2 | 0.2 ± 0.2 |
| *alr4550*                | -6.4 ± 0.1 | -0.5 ± 0.1 | -3.5 ± 0.2 | -3.3 ± 0.2 | -2.8 ± 0.2 |
| *alr4714*                | -11.5 ± 0.4 | 1.3 ± 0.2 | 0.2 ± 0.2 | -0.3 ± 0.2 | 0.6 ± 0.2 |
| *all5191*                | -12.1 ± 0.5 | 1.8 ± 0.1 | 1.1 ± 0.1 | 0.0 ± 0.2 | 1.7 ± 0.1 |
| *all7614*                | -8.3 ± 0.2 | 0.3 ± 0.1 | -0.5 ± 0.1 | -0.8 ± 0.2 | -0.3 ± 0.1 |

The -ΔCt value based on the housekeeping gene rnpB is given for YBG11 and the -ΔΔCt based on the YBG11 values is shown for the treatments, the standard deviation is indicated. Values in bold represent changes with $p < 0.05$ (Student’s t-test) and an absolute fold-change higher or lower (italics) than one.
was performed on cDNA. The high transcript abundance of \textit{alr4550} in standard YBG11 medium corresponded to proteomic analyses in which Alr4550 and All4499 were identified as the highest abundant porins in the OM of \textit{Anabaena} sp. [30].

The mRNA abundance of all7614 was not affected after 21 days of culturing wild type in media lacking either manganese, iron, copper or zinc (Table 1). Manganese deprivation resulted in an increase of the transcripts of \textit{alr0834}, \textit{alr2231}, \textit{alr4741} and \textit{alr5191} when compared to YBG11 (Table 1). The transcript of \textit{alr0834} was further increased under iron limitation, whereas the transcript of \textit{alr5191} was increased under iron and zinc limitation. Moreover, iron deprivation yielded in an elevated \textit{all4499} transcript level when compared to cultivation in YBG11.

Notably \textit{alr4550} demonstrated an exceptional behavior. On the one hand, it is highly expressed in YBG11 when normalized to \textit{rnpB}. On the other hand, among all tested genes, only \textit{alr4550} transcripts decreased in medium lacking Fe, Cu or Zn (Table 1). Only Mn deprivation did not significantly alter \textit{alr4550} transcript levels (Table 1).

In conclusion, the expression of the genes coding for the seven porin-like proteins does not show a common regulation after 21 days in media without individual trace metals. This might indicate distinct functions of the analyzed proteins. Altogether, manganese and iron depletion resulted in increased abundance of four and three transcripts, respectively. Zn depletion led to enhanced mRNA levels of only \textit{all5191}.

### The Mutants of Porin-like Genes Show a Differential Sensitivity to Divalent Metal Stress

The sensitivity against elevated metal concentrations was tested with the porin mutants, as it is expected that strains with defected transport capacities of certain substrates display hyper-resistance towards increased concentrations of these substrates. Interestingly, virtually all mutant strains except AFS-I-\textit{alr4741} exhibited a resistance towards elevated copper concentrations (35 μM) compared to the wild type. AFS-I-\textit{alr4550} and AFS-I-\textit{all7614} grew to a slightly lower density under those conditions compared to the other mutant strains (Fig. 2C, first panel). Also, the growth of the two \textit{omp85} mutants, AFS-I-\textit{alr2269} and AFS-I-\textit{alr4893}, was inhibited under elevated copper concentrations, while AFS-I-\textit{alr0075} grew to a lower density (Fig. 2C, first panel). Thus, the \textit{omp85} mutants exhibited a similar sensitivity against the selected copper concentration compared to wild type, while AFS-I-\textit{tic22} showed an enhanced resistance. It can be concluded that copper entry into the porin mutants is largely limited compared to wild type. Notably, none of the porin transcripts was increased after copper depletion, leading to the assumption that this effect might not be specifically related to a single porin (Table 1).

On plates with an excess of manganese (1.35 mM MnCl$_2$; Fig. 2C, second & third panel) AFS-I-\textit{alr4550} and AFS-I-\textit{all7614} grew to lower densities when compared to wild type, while AFS-I-\textit{alr0834}, AFS-I-\textit{alr4499} and AFS-I-\textit{alr4741} mutants, AFS-I-\textit{alr2269} and AFS-I-\textit{alr4893} were more resistant than wild type, while AFS-I-\textit{tic22} showed a higher sensitivity (Fig. 2C, second & third panel). Interestingly AFS-I-\textit{alr2231} grew better than wild type when the zinc concentration was enhanced (Fig. 2C, fifth panel). With respect to the control strains, only AFS-I-\textit{alr4893} was hyper-sensitive towards this zinc concentration (Fig. 2C, fourth panel).

In addition to the mentioned divalent metals, the growth in the presence of enhanced cobalt concentrations was determined as well. The ICP-MS measurements showed that cobalt was about ten-fold less abundant in cells then determined as well. The ICP-MS measurements showed that cobalt was about ten-fold less abundant in cells then in standard YBG11 medium corresponds to

### Table 2. qRT-PCR analysis of porin genes in individual porin mutants.

|        | \textit{alr0834} | \textit{alr2231} | \textit{alr4499} | \textit{alr4741} |
|--------|-----------------|-----------------|-----------------|-----------------|
| \textit{alr0834} | n.d.            | 1.8 ± 0.4       | 4.4 ± 0.5       | 0.9 ± 0.4       |
| \textit{alr2231} | 0.7 ± 0.3       | n.d.            | 0.8 ± 0.5       | 2.8 ± 0.5       |
| \textit{all4499} | -0.2 ± 0.1      | 3.9 ± 0.4       | n.d.            | 0.2 ± 0.2       |
| \textit{alr4550} | 1.5 ± 0.2       | 2.0 ± 0.4       | 0.7 ± 0.3       | n.d.            |
| \textit{alr4741} | -0.1 ± 0.2      | 2.5 ± 0.3       | 0.0 ± 0.5       | n.d.            |
| \textit{all5191} | 1.0 ± 0.1       | 2.2 ± 0.3       | 0.7 ± 0.4       | 1.3 ± 0.3       |
| \textit{all7614} | -2.5 ± 0.1      | 1.6 ± 0.4       | 0.5 ± 0.2       | 0.1 ± 0.2       |

The \(-ΔΔCt\) based on the \textit{rnpB} expression and BG11 values are shown for the mutants. Values in bold are changes with \(p < 0.05\) (Student's t-test) compared to wild type.

May 2021 | Vol. 31 | No. 5
in this mutant strain was analyzed. Interestingly, the transcript abundance of all other porin-coding genes was increased in this mutant (Table 2). This strengthens the hypothesis that zinc resistance is mediated by \textit{alr}2231-insertion and not by the diminished transcription of another putative porin gene.

In addition, in the presence of enhanced concentrations of manganese, the mutants of \textit{alr}0834, \textit{all}4499 and \textit{alr}4741 were more resistant than wild type. Interestingly, \textit{all}4499 was comparatively high expressed under standard conditions and the transcript abundance did not change after manganese deprivation. However, the expression of \textit{alr}0834 and \textit{alr}4741 was upregulated in the absence of manganese (Table 1). Analyzing the transcript abundance of the other porins in the respective mutants revealed a downregulation of \textit{alr}4550 and \textit{all}7614 expression in \textit{AFS-I-alr}0834 (Table 2). The mutants of \textit{alr}4550 and \textit{all}7614 were the only strains showing a hypersensitivity towards an elevated manganese concentration. Therefore, this downregulation does not explain the manganese resistance of \textit{AFS-I-alr}0834. In \textit{AFS-I-alr}4499 and \textit{AFS-I-alr}4741 no other gene coding for a porin-like protein was found to be downregulated (Table 2). Thus, the resistance of \textit{AFS-I-alr}4499 and \textit{AFS-I-alr}4741 against elevated manganese can be attributed to the mutated gene.

The Mutant of \textit{alr}4550 Shows a Reduced Integrity of the Outer Membrane

Next, the integrity of the outer envelope was analyzed in the porin mutants. To this end, lysozyme that catalyzes peptidoglycan hydrolysis was added to the medium (Fig. 4A, first panel). \textit{AFS-I-alr}4550, and \textit{AFS-I-tic}22 were hampered in growth in presence of 250 μg ml⁻¹ lysozyme when compared to wild type. For \textit{AFS-I-tic}22, this is in line with earlier findings [46]. In turn, \textit{AFS-I-alr}0075 grew similar to wild type, while all other porin mutants showed an improved growth compared to wild type. This was observed as well for the \textit{omp}85 mutants \textit{AFS-I-alr}2269 and \textit{AFS-I-alr}4893. On plates containing 10 μg ml⁻¹ SDS, the porin mutants followed a comparative trend as in presence of lysozyme; the porin mutants grew somewhat better than wild type, which in turn did grow better as in presence of lysozyme (Fig. 4A, second panel). Here \textit{AFS-I-alr}4550 did grow, but only to a low density. The mutants \textit{AFS-I-alr}2269 and \textit{AFS-I-tic}22 were inhibited in growth in the presence of SDS, while \textit{AFS-I-alr}4893 was less affected than wild type. In the presence of 50 μg ml⁻¹ proteinase K the porin mutants behaved again similar as in the presence of lysozyme, while only \textit{AFS-I-all}7614 behaved similar to wild type (Fig. 4A, third panel). \textit{AFS-I-alr}0075 and \textit{AFS-I-tic}22 were inhibited in growth in presence of protease K. Instead, \textit{AFS-I-alr}2269 growth was comparable to wild type and \textit{AFS-I-alr}4893 grew better than wild type. In presence of the antibiotic erythromycin again all porin mutants except \textit{AFS-I-alr}4550 showed an enhanced growth compared to wild type (Fig. 4A, fourth panel), with the exception of \textit{AFS-I-alr}2269 which exhibited growth inhibition.

Next, the cells were grown under salt stress, which is known to alter the abundance of porins [64]. In the presence of 100 mM KCl a similar growth of the porin mutants as in the presence of protease K was observed, while the growth of the \textit{omp}85 mutants was enhanced. Growth of \textit{AFS-I-tic}22 on the other hand was severely affected (Fig. 4A, fifth panel). The addition of 100 mM NaCl reduced the growth of \textit{AFS-I-alr}4550, \textit{AFS-I-all}7614 and \textit{AFS-I-tic}22 when compared to wild type (Fig. 4A, sixth panel). When 150 mM NaCl was added, wild type, \textit{AFS-I-alr}4550, \textit{AFS-I-all}7614, \textit{AFS-I-alr}2269 and \textit{AFS-I-tic}22 were defective in growth, while \textit{AFS-I-alr}0834, \textit{AFS-I-alr}2231, \textit{AFS-I-all}4499, \textit{AFS-I-alr}4714, \textit{AFS-I-all}5191, \textit{AFS-I-alr}0075 and \textit{AFS-I-all}4893 were able to grow on this medium (Fig. 4A, seventh panel).

Our results demonstrate that the mutants of five porin-like genes (\textit{alr}0834, \textit{alr}2231, \textit{all}4499, \textit{alr}4741 and \textit{all}5191) were more resistant towards the selected compounds, which is indicative for alterations in the outer membrane integrity. \textit{AFS-I-tic}22 for instance, a previously characterized mutant with alterations in OM defects (Fig. 4). In contrast, mutation of the highest expressed porin-like gene \textit{alr}4550 causes a phenotype that is consistent with an impaired outer membrane integrity in general (Figs. 2 and 4). \textit{AFS-I-all}7614 showed an intermediate phenotype. On the one hand, the strain was sensitive to divalent metals and sodium chloride (Figs. 2 and 4), on the other hand the mutant presented a higher resistance to SDS and lysozyme.

**Fig. 4. Outer membrane integrity of porin mutants.** (A) 5 μl of wild type, the porin, the \textit{omp}85 and the \textit{tic}22 insertion mutants (indicated on top) at OD₇₅₀ = 1.0 were spotted onto media composed of YBG11 supplemented with indicated divalent metals. Images were taken after 7 days of growth. Representative results (n = 3) are shown. (B) The increased (blue arrow) or decreased (red arrow) growth capacity of a mutant strain on YBG11 with the indicated ingredient in comparison to wild type is indicated. LYS = lysozyme, SDS = sodium dodecyl sulfate, PK = proteinase K, ERY = erythromycin.
To determine the importance of the Omp85 protein function for porin-insertion into the outer membrane, the protein abundance of a porin was examined in wild type and the \(\text{omp85}\) mutants. As Alr4550 was found to be the most abundant protein component of the \textit{Anabaena} sp. outer membrane [30] the abundance of this protein was tested. Total protein was extracted from wild type, AFS-I-alr0075, AFS-I-alr2269 and AFS-I-alr4893. The mRNA levels of alr2231 were not affected by the mutation of \(\text{omp85}\) genes.

### Table 3. qRT-PCR analysis of expression of porin genes in \(\text{omp85}\) mutants.

|                     | AFS-I- | AFS-I-alr0075 | AFS-I-alr2269 | AFS-I-alr4893 |
|---------------------|--------|---------------|---------------|---------------|
| \(\Delta\Delta C\)   |        |               |               |               |
| \(\text{alr0834}\)  | -1.2 ± 0.6 | -2.2 ± 0.1    | -0.2 ± 0.1    |               |
| \(\text{alr2231}\)  | -0.7 ± 0.1 | -2.5 ± 2.1    | 0.2 ± 0.1     |               |
| \(\text{all4499}\)  | -2.1 ± 0.6 | 0.0 ± 0.1     | 0.3 ± 0.1     |               |
| \(\text{alr4550}\)  | 1.6 ± 1.2 | 0.7 ± 0.3     | 2.0 ± 0.3     |               |
| \(\text{alr4741}\)  | 0.2 ± 0.1 | 0.5 ± 0.3     | 1.4 ± 0.3     |               |
| \(\text{all5191}\)  | -2.3 ± 0.7 | -2.3 ± 0.1    | -0.8 ± 0.5    |               |
| \(\text{all7614}\)  | 1.5 ± 0.9 | 1.5 ± 0.1     | -13 ± 11      |               |

The \(\Delta\Delta C\) based on \(\text{rnpB}\) expression and BG11 values is shown for the mutants. Values in bold are changes with \(p < 0.05\) (Student’s t-test) compared to wild type.

**Omp85 Proteins are Distinct in Their Function In Porin Biogenesis**

Altogether these results can be seen as first hint towards a putative functional relation between the Omp85 protein Alr4893 and most of the porins, as the mutant phenotypes were consistent under many conditions (Figs. 2 and 4). In order to assess the influence of the \(\text{omp85}\) gene mutations on the porins, the expression of the putative porin-coding genes was analyzed in the three \(\text{omp85}\) mutants (Table 3). In AFS-I-alr0075 a reduction of the transcripts of all4499 and all5191 was observed. In AFS-I-alr2269 a reduction in transcript abundance of alr0834 and all5191 was detected, while the abundance of all7614 was enhanced (Table 3). In turn, the transcript abundance of all7614 was strongly reduced in AFS-I-alr4893, while the transcript levels of alr4550 and alr4741 were increased. The mRNA levels of alr2231 were not affected by the mutation of \(\text{omp85}\) genes.

Fig. 5. Relation between Omp85 function and porin biogenesis. (A) A representative result (\(n = 3\)) for the outer membrane protein content in wild type (first lane) and the three indicated \(\text{omp85}\) mutants (lane 2-4) is shown. Total protein extract was probed with specific antibodies against Alr2269 or Alr4550 as indicated. DB71 staining of the large subunit of Rubisco (LSU) was used as loading control. The average of the ratio of the protein density in mutants and wild type is shown as analyzed with ImageJ. The standard deviation is smaller than 25%. All lanes come from the same gel and have been developed simultaneously. (B, C) Outer membrane vesicles were isolated from the indicated strains and subjected to rate zonal ultracentrifugation. Fractions 2-10 of the sucrose gradient (25% to 70% (v/w)) were probed with an antibody against Alr2269 (B) or Alr4550 (C). Note: in B the result for AFS-I-alr2269 was longer exposed than other blots to visualize the bands. In (B) and (C) one of two repetitions is shown. The densitometric protein distribution of both experiments normalized to the highest intensity is shown on the bottom of each panel as bar diagram. In (A) and (C) the blue arrowhead marks the presumed degradation product of Alr4550, the grey arrowhead marks the full length protein and the red arrowhead points at a fragment that likely represents a non-native state of the protein. The migration of the 66 kDa molecular weight standard is indicated.
presence of outer membranes in the total protein fractions was confirmed by detection of Alr2269 using a specific antibody. The protein was detected in the cell lysate of all strains except AFS-1-\(alr2269\) (Fig. 5A, panel 1). Thus, Alr2269 could not be detected in AFS-1-\(alr2269\)-lysate although the strain is not segregated (Fig. S1). The relatively small amount of outer membrane proteins in total cell lysate and the reduced protein abundance in the mutant are possible explanations for this. The loading of lysate was comparable between all four strains as controlled by DB71 staining (Fig. 5A, panel 3, LSU). Probing for Alr4550 in wild type led to two apparent signals (Fig. 5A, panel 2). The upper band corresponds to the full length protein, whereas the lower molecular weight signal most likely represents a degradation product. Both fragments have been detected in wild type in previous experiments [45]. The size of the lowest fragment is consistent with the molecular weight of Alr4550 lacking the periplasmic 5-layer homology domain.

Alr4550 was more abundant in the \(omp85\) mutants AFS-1-\(alr2269\) and AFS-1-\(alr4893\) compared to the wild type. Especially in AFS-1-\(alr4893\) the protein amount was comparatively increased. Moreover, the protein band of higher molecular weight than the mature protein was more abundant in the mutants when compared to wild type (Fig. 5A, panel 2). This apparent fragment likely represents an SDS-resistant unfolded or aggregated intermediate, which likewise has been reported for other outer membrane β-barrel proteins [65, 66]. In AFS-1-\(alr2269\) the high molecular weight fragment was the dominating species, whereas in AFS-1-\(alr4893\) the form migrating as intermediate predominated. Hence, our results might suggest that the plasmid-insertion in \(alr2269\) or \(alr4893\) leads to an increased protein production of Alr4550 as well as an increased detection of misfolded or degraded proteins. The increase in Alr4550 protein amount was found to be reflected by an enhanced mRNA abundance of \(alr4550\) in AFS-1-\(alr4893\), but not AFS-1-\(alr2269\) (Table 3).

Alterations in the outer membrane integrity and biogenesis as well as an alteration of the protein to lipid ratio can result in an aberrant membrane density which can be examined by rate zonal centrifugation [67-69]. The effects of the \(omp85\) mutations on membrane protein integrity were examined by analyzing the outer membrane protein density in the single insertion mutants. The sedimentation behavior of outer membranes was analyzed by rate zonal centrifugation. Gradient fractions were collected and subjected to SDS-PAGE. To detect non-porin type OM-proteins the antibody against Alr2269 was utilized (Fig. 5B). As AFS-1-\(alr2269\) is not segregated a signal was obtained in this sample as well.

Isolated outer membranes from \(Anabaena\) sp. that contained Alr2269 sedimented in the range of 49% to 61% sucrose. The largest quantity of Alr2269 was detected between 49% and 53% (Fig. 5B, top panel). The same distribution of Alr2269 was detected when outer membrane preparations of the \(omp85\) mutants or AFS-1-\(tic22\) were analyzed. However, in AFS-1-\(alr0075\), AFS-1-\(alr2269\) and AFS-1-\(tic22\) the majority of the protein migrated between 53% and 56% sucrose (Fig. 5B).

In wild type outer membrane fractions Alr4550 was more broadly dispersed than Alr2269. Similar to Alr2269, the largest quantity of Alr4550 was found between 49% to 61% sucrose as well. The peak was shifted to the fractions between 53% and 56% sucrose (Fig. 5C, top). Interestingly, the distribution of Alr4550 in the membrane fractions isolated form AFS-1-\(alr0075\) (Fig. 5C, fourth panel) and AFS-1-\(tic22\) (Fig. 5C, bottom) was comparable to that of wild type. Hence, Alr4550 synthesis and integrity seems not affected in AFS-1-\(alr0075\). This is also underlined by the fact that the Alr4550-fragment which putatively represents the aggregated form of the protein was not detectable in AFS-1-\(alr0075\) (Fig. 5A and 5C, fourth panel). In turn, in AFS-1-\(tic22\) the aggregated form was detectable (Fig. 5C, bottom, red arrowhead).

The sedimentation behavior of Alr4550 in samples from AFS-1-\(alr2269\) and AFS-1-\(alr4893\) showed clear distinctions compared to wild type. While the general distribution profile of the membrane fractions containing Alr4550 from AFS-1-\(alr2269\) was more disperse than in wild type (ranging from 43% to 61%), the peak fraction was found at a comparable density (Fig. 5C, second panel). However, an enrichment of the porin in fractions of higher density that possibly represents aggregated protein was observed (Fig. 5C, second panel, fraction 10). The migration of the membrane fractions isolated from AFS-1-\(alr4893\) that contained Alr4550 was shifted to higher density compared to wild type (56%-61% sucrose; Fig. 5C, third panel). Just like observed in AFS-1-\(alr2269\) samples, a significant portion of Alr4550 was found in the last gradient fractions. In both strains the high molecular-weight fragment that likely represents the aggregated form was identified, whereas in wild type samples it was not detected in this experiment. This is consistent with earlier reports documenting the aggregation of unfolded, not inserted proteins at the outer membrane [70].

Taken together these results indicate that outer membrane biogenesis or integrity is affected in AFS-1-\(alr2269\) and AFS-1-\(alr4893\) as judged from the sucrose gradient centrifugation analysis and detection of Alr2269 and Alr4550, whereas AFS-1-\(alr0075\) samples only exhibited an aberrant distribution of Alr2269, but not Alr4550. Factors that could cause the increased density of the vesicles could be for example an increased protein to lipid ratio, an altered (enhanced) co-migration of peptidoglycan with the proteins or variations in LPS production [69, 71, 72].

**Discussion**

Porins have an important function in the regulation of solute uptake and further contribute to the maintenance of envelope integrity [8]. The degree of specificity of certain porin classes is highly diverse [6]. In proteobacteria a distinction between e.g. OmpA-type porins (structural function) and OmpF-type porins (transport function) can be made [8]. In the genome of the filamentous cyanobacterium \(Anabaena\) sp. a typical OmpA-like porin, namely Alr4550, was identified. Expression analysis revealed that among all the porins in \(Anabaena\) sp. the \(alr4550\) transcript was high abundant under standard conditions (Table 1). Remarkably AFS-1-\(alr4550\) was hypersensitive compared to wild type under virtually all tested conditions (except elevated copper levels). Growth of AFS-1-
alr4550 in the absence of fixed nitrogen was not inhibited, hence Alr4550 dysfunction seems not to affect heterocyst development (Figs. 2 and 3). Thus, Alr4550 might have a prominent structural function in Anabaena sp. that is distinct from the function of the other porin-like proteins, considering the exceptionality of the mutant phenotype. This could be for instance mediated by the interaction of Alr4550 with cell wall (components) and thereby connecting the OM to the peptidoglycan, as described for OmpA in E. coli [49, 73].

Most of the porin mutants were increasingly resistant towards high potassium chloride and sodium chloride concentrations compared to the wild type. The same was observed when the macrolide erythromycin was added. In contrast to other antibiotics, macrolide antibiotics are generally not thought to penetrate through porins, therefore the enhanced resistance might point towards altered membrane properties (Fig. 3) [18, 74]. Interestingly, AIS-1-all5191 was the only strain that did barely grow in the absence of a combined nitrogen source (Fig. 2). Apparently, this strain was able to differentiate heterocysts as judged from microscopic analyses, showing that the protein might not play a role in heterocyst development. We rather suggest that the absence of All5191 generates a condition that complicates nitrogen fixation. Consequently, the percentage of heterocysts increases is AIS-1-all5191 filaments. AIS-1-omp834 and AIS-1-omp4741 grew better on YBG11, compared to wild type (Fig. 2). An alteration in the heterocyst pattern of the two strains was not observed, thus the reason for the enhanced growth of the mutants remains to be elucidated. Noteworthy the transcript of all5191 was increasingly abundant in AIS-1-alr0834 and AIS-1-omp4741. Since AIS-1-all5191 was defected in growth in absence of a combined nitrogen source, an increased production of the protein might on the other hand have beneficial effects on the growth capacity in YBG11, as observed in AIS-1-omp834 and AIS-1-omp4741.

With respect to the uptake of divalent ions, a relation to iron uptake could not be established in this study. Ferric iron ions are prevalently complexed to organic ligands like siderophores [75]. However, the transport of siderophores in Anabaena is rather dependent on functional TBDTs instead of porins [63, 76]. It was shown that inorganic iron is highly bioavailable to cyanobacteria [77, 78]. Moreover, homologues of the iron-transporting porin Srl1908 in Synechocystis sp. are found in many freshwater and marine cyanobacteria species [27]. This generally suggests an important role of porin-mediated iron transport among cyanobacteria, which needs to be further examined in Anabaena sp.

Also for cobalt uptake, no relation to a porin in Anabaena sp. could be made, as mutation of porins did not lead to a resistance against elevated concentrations (Fig. 2). Cobalt also might be taken up in Anabaena sp. in form of cobalamin. This uptake is dependent on the BtuB proteins of the TBDT-family, which are predicted to exist in Anabaena sp. as well [79]. Similarly, two TBDTs but none of the porin-like genes investigated here have been found to be regulated by the zinc starvation sensor Zur in Anabaena sp. [80]. This is consistent with the earlier identification of the zinc transporter ZnuD in proteobacteria, which belongs to the TonB-dependent transporter (TBDT) family [81, 82]. However, AIS-1-omp2231 shows a high resistance to elevated levels of zinc (Fig. 2), which might link the Alr2231 function to the uptake of zinc when the trace metal is present in sufficient amounts. Such function would be consistent with the identification of a cyanobacterial porin as zinc binding protein [83]. An adequate zinc supply is important for organisms as for example the carbonic anhydrase depends on zinc. This enzyme is involved in conversion of CO₂ to bicarbonate, which is important for the regulation of the pyruvate conversion to oxaloacetate and thus central for the cyanobacterial metabolism.

The uptake of other divalent metals like manganese and copper is discussed to depend on porin-like proteins in Gram-negative bacteria [84–88]. The importance of manganese is associated with the regulation of the pyruvate pool as phosphoenolpyruvate carboxykinase and pyruvate carboxylase are Mn-dependent metalloenzymes. Moreover, the oxygen evolving complex in photosystem II bears manganese as cofactor [89]. This uptake is dependent on the BtuB proteins of the TBDT-family, which are predicted to exist in Anabaena sp. as well [79]. Similarly, two TBDTs but none of the porin-like genes investigated here have been found to be regulated by the zinc starvation sensor Zur in Anabaena sp. [80]. This is consistent with the earlier identification of the zinc transporter ZnuD in proteobacteria, which belongs to the TonB-dependent transporter (TBDT) family [81, 82]. However, AIS-1-omp2231 shows a high resistance to elevated levels of zinc (Fig. 2), which might link the Alr2231 function to the uptake of zinc when the trace metal is present in sufficient amounts. Such function would be consistent with the identification of a cyanobacterial porin as zinc binding protein [83]. An adequate zinc supply is important for organisms as for example the carbonic anhydrase depends on zinc. This enzyme is involved in conversion of CO₂ to bicarbonate, which is important for the regulation of the pyruvate conversion to oxaloacetate and thus central for the cyanobacterial metabolism.

The uptake of other divalent metals like manganese and copper is discussed to depend on porin-like proteins in Gram-negative bacteria [84–88]. The importance of manganese is associated with the regulation of the pyruvate pool as phosphoenolpyruvate carboxykinase and pyruvate carboxylase are Mn-dependent metalloenzymes. Moreover, the oxygen evolving complex in photosystem II bears manganese as cofactor [89]. This uptake is dependent on the BtuB proteins of the TBDT-family, which are predicted to exist in Anabaena sp. as well [79]. Similarly, two TBDTs but none of the porin-like genes investigated here have been found to be regulated by the zinc starvation sensor Zur in Anabaena sp. [80]. This is consistent with the earlier identification of the zinc transporter ZnuD in proteobacteria, which belongs to the TonB-dependent transporter (TBDT) family [81, 82]. However, AIS-1-omp2231 shows a high resistance to elevated levels of zinc (Fig. 2), which might link the Alr2231 function to the uptake of zinc when the trace metal is present in sufficient amounts. Such function would be consistent with the identification of a cyanobacterial porin as zinc binding protein [83]. An adequate zinc supply is important for organisms as for example the carbonic anhydrase depends on zinc. This enzyme is involved in conversion of CO₂ to bicarbonate, which is important for the regulation of the pyruvate conversion to oxaloacetate and thus central for the cyanobacterial metabolism.

The uptake of other divalent metals like manganese and copper is discussed to depend on porin-like proteins in Gram-negative bacteria [84–88]. The importance of manganese is associated with the regulation of the pyruvate pool as phosphoenolpyruvate carboxykinase and pyruvate carboxylase are Mn-dependent metalloenzymes. Moreover, the oxygen evolving complex in photosystem II bears manganese as cofactor [89]. This uptake is dependent on the BtuB proteins of the TBDT-family, which are predicted to exist in Anabaena sp. as well [79]. Similarly, two TBDTs but none of the porin-like genes investigated here have been found to be regulated by the zinc starvation sensor Zur in Anabaena sp. [80]. This is consistent with the earlier identification of the zinc transporter ZnuD in proteobacteria, which belongs to the TonB-dependent transporter (TBDT) family [81, 82]. However, AIS-1-omp2231 shows a high resistance to elevated levels of zinc (Fig. 2), which might link the Alr2231 function to the uptake of zinc when the trace metal is present in sufficient amounts. Such function would be consistent with the identification of a cyanobacterial porin as zinc binding protein [83]. An adequate zinc supply is important for organisms as for example the carbonic anhydrase depends on zinc. This enzyme is involved in conversion of CO₂ to bicarbonate, which is important for the regulation of the pyruvate conversion to oxaloacetate and thus central for the cyanobacterial metabolism.
Our results imply diverse functions of single porin-like proteins, reflected by the specific phenotypes of the mutants. Most prominently, AFS-1-alr4550 exhibited a unique phenotype by sowing hypersensitivity towards metals, salt and harmful compounds. Also the strong transcription of alr4550 suggests a crucial function of the protein, which apparently involves the maintenance of envelope integrity. First associations between single porin candidates and manganese, zinc or copper transport as well as diazotrophic growth capacity could be drawn, thus these connections need to be further examined. The density gradient centrifugation experiments showed that the three omp85 mutants in Anabaena sp. are affected in the migration of Alr2269 and Alr4550, but to different extents. The presence of aggregated intermediates of Alr4550 in the two omp85 mutants AFS-1-alr2269 and AFS-1-alr4893, but not in AFS-1-alr0075, suggests a functional specification on certain substrates for membrane insertion.

Author Contributions
ES conceptualized and HS, EMB and ES designed the study. HS, EMB, EL, and MS conducted experiments. HS, EMB and ES performed analysis and interpretation of data. HS and ES wrote the original manuscript. All authors critically revised and approved the article.

Acknowledgments
We would like to thank Prof. Enrique Flores for providing material used for mutant generation. We thank Dr. Sotirios Fragkostefanakis and Julia Graf for comments on the manuscript. Moreover, we thank Dr. H.-Michael Seitz and Prof. Dr. Horst Marschall for support with the ICP-MS analysis. The work was funded by the Deutsche Forschungsgemeinschaft DFG SCHL585/7-2 to ES. HS received a stipend of the Buchmann Foundation.

Conflict of Interest
The authors have no financial conflicts of interest to declare.

References
1. Jürgens UJ, Drews G, Weckesser J. 1983. Primary structure of the peptidoglycan from the unicellular cyanobacterium Synechocystis sp. strain PCC 6714. J. Bacteriol. 154:471-478.
2. Wilk L, Strauss M, Rudolf M, Nicolsen K, Flores E, Kühlbrandt W, et al. 2011. Outer membrane continuity and septosome formation between vegetative cells in the filaments of Anabaena sp. PCC 7120. Cell. Microbiol. 13:1744-1754.
3. Matias VRF, Al-Amoudi A, Dubochet J, Beveridge TJ. 2003. Cryo-transmission electron microscopy of frozen-hydrated sections of Escherichia coli and Pseudomonas aeruginosa. J. Bacteriol. 185:6112-6118.
4. Baldridge S, Falso M, Cantisani M, Tarallo R, Elena Della Pepa M, D’Oriano V, et al. 2013. Microbe-Host Interactions: Structure and Role of Gram-Negative Bacterial Porins. Curr. Protein Pept. Sci. 13:843-854.
5. Nakaïo H. 1994. Porins and specific diffusion channels in bacterial outer membranes. J. Biol. Chem. 269:3905-3908.
6. Choi U, Lee CR. 2019. Distinct roles of outer membrane porins in antibiotic resistance and membrane integrity in Escherichia coli. Front. Microbiol. 10:953.
7. Novikova OD, Solovyeva TF. 2009. Non-specific porins of the outer membrane of Gram-negative bacteria: Structure and functions. Biochem. Suppl. Ser. A: Membr. Cell. Biol. 3:3-15.
8. Schirmer T, Wang YE, Rosenbusch JP. 1995. Structural basis for sugar translocation through maltoporin channels at 3.1 Å resolution. Science 267:512-516.
9. Connan S, Zhang Y, Chelley S, Bayley H. 2000. Biochemical and biophysical characterization of OmpG: A monomeric porin. Biochemistry 39:11845-11854.
10. Kim KH, Aulakh S, Paetzel M. 2012. The bacterial outer membrane β-barrel assembly machinery. Protein Sci. 21:751-768.
11. Zimmerman RE, Dave AM, Bruce BD. 2014. Structure and Function of POTRA Domains of Omp85/TPS Superfamily. pp. 1-34. In International Review of Cell and Molecular Biology, Elsevier Inc.
12. Kumar K, Meda Herrera RA, Golden JW. 2010. Cyanobacterial heterocysts. Cold Spring Harb. Perspect. Biol. 2:a003315.
13. Moslavac S, Reisinger V, Berg M, Mirus O, Vosyka O, Ploščer M, et al. 2007. The proteome of the heterocyst cell wall in Anabaena sp. PCC 7120. Biol. Chem. 388:823-829.
14. Hancock REW. 1987. Role of porins in outer membrane permeability. J. Bacteriol. 169:929-933.
15. Jap BK, Watanuki Y. 1990. Biophysics of the structure and function of porins. Q. Rev. Biophys. 23:367-403.
16. Nakaïo H. 2003. Molecular basis of bacterial outer membrane permeability revisited. Microbiol. Mol. Biol. Rev. 67:593-656.
17. Hansel A, Tadros MH. 1998. Characterization of two pore-forming proteins isolated from the outer membrane of Synechococcus PCC 6301. Curr. Microbiol. 36:321-326.
18. Brechtel E, Bahl H. 1999. In Thermoanaerobacterium thermosulfurigenes EM1 S-layer homology domains do not attach to peptidoglycan. J. Bacteriol. 181:5017-5023.
19. Mk N, Kosma P, Puchberger M, Egelseer EM, Mayer HF, Sletten UB, et al. 1999. Structural and functional analyses of the secondary cell wall polypeptide of Bacillus subtilis serineni CCM 2177 that serves as an S-layer-specific anchor. J. Bacteriol. 181:7643-7646.
20. Kern H, Ryan C, Fassl K, Schneewind O. 2010. Bacillus anthracis surface-layer proteins assemble by binding to the secondary cell wall polysaccharide in a manner that requires csb and tagO. J. Mol. Biol. 401:757-775.
21. Lups A, Engelhardt H, Peters J, Santarius U, Volker S, Baumeister W. 1994. Domain structure of the Acetogenium kivui surface layer revealed by electron crystallography and sequence analysis. J. Bacteriol. 176:1224-1233.
22. Kowalz H, Tochigi S, Takahashi H, Kojima S. 2017. Outer membrane permeability of cyanobacterium Synechocystis sp. strain PCC 6803: Studies of passive diffusion of small organic nutrients reveal the absence of classical porins and intrinsically low permeability. J. Bacteriol. 199:e00371-17.
23. Ekman M, Picossi S, Campbell EL, Meeks JC, Flores E. 2013. A Nostoc punctiforme sugar transporter necessary to establish a cyanobacterium-plant symbiosis. Plant. Physiol. 161:1984-1992.
24. Simm S, Keller M, Selymes M, Schleiff E. 2015. The composition of the global and feature specific cyanobacterial core-genomes. *Front. Microbiol.* 6:219.

25. Qi G, Jiang H, Li H, Li Z, Deng B, Shang J, et al. 2020. A unique porin mediates iron-selective transport through cyanobacterial outer membrane proteins. *Environ. Microbiol.* 23:376-390.

26. Hahn A, Schleiff E. 2014. The Cell Envelope. In *The Cell Biology of Cyanobacteria*, Flores E, Herrero A, eds., pp. 29-87. Caister Academic Press, UK, Norfolk.

27. Oliveira P, Martins NM, Santos M, Couto NAS, Wright PC, Tamagnini F. 2015. The *Anabaena* sp. PCC 7120 exoproteome: Taking a peek outside the box. *Life Sci.* 5:130-163.

28. Moslavac S, Bredemeier R, Mirus O, Granvogl B, Eichacker LA, Schleiff E. 2005. Proteomic analysis of the outer membrane of the cyanobacterium *Anabaena* sp. strain PCC 7120. *J. Proteome Res.* 4:1330-1338.

29. Hahn A, Stevanovic M, Mirus O, Schleiff E. 2012. The TolC-like protein HgdD of the cyanobacterium *Anabaena* sp. strain PCC 7120 is involved in secondary metabolite export and antibiotic resistance. *J. Biol. Chem.* 287:41126-41138.

30. Acland A, Agarwala R, Barrett T, Beck J, Benson DA, Bollin C, et al. 2014. Database resources of the National Center for Biotechnology Information. *Nucleic Acids Res.* 42:D7-D17.

31. Crooks GE, Hon G, Chandonia J-M, Brenner SE. 2004. WebLogo: A sequence logo generator. *Genome Res.* 14:1188-1190.

32. Sievers F, Wilm A, Dineen D, Gibson TJ, Karplus K, Li W, et al. 2011. Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. *Mol. Syst. Biol.* 7:539.

33. Madeira F, Park YM, Lee J, Buso N, Gur T, Madhusoodanan N, et al. 2019. The EMBL-EBI search and sequence analysis tools APIs in 2019. *Nucleic Acids Res.* 47:W636-W641.

34. Valladares A, Rodriguez V, Camargo S, Martinez-Noel GMA, Herrero A, Luque I. 2011. Specific role of the cyanobacterial pipX gene in Synechocystis sp. strain PCC 7002. *Protoc. Microbiol. Physiol.* 5:W636-W641.

35. Rudolf M, Kranzler C, Lis H, Margulis K, Stevanovic M, Keren N, Schleiff E. 2015. Multiple modes of iron uptake by the filamentous, nitrogen-fixing filamentous cyanobacteria *Anabaena* sp. strain PCC 7120. *Environ. Microbiol.* 17:58-70.

36. Struyve M, Moons M, Tommassen J. 1991. Carboxy-terminal phenylalanine is essential for the correct assembly of a bacterial outer membrane protein. *J. Biol. Chem.* 266:1329-1335.

37. Liu X, Ferenci T. 2008. Regulation of porin-mediated outer membrane permeability by nutrient limitation in *Escherichia coli*. *J. Bacteriol.* 190:3917-3922.

38. Berman-Frank I, Cullen JT, Shaked Y, Sherrill RM, Falkowski PG. 2001. Iron availability, cellular iron quotas, and nitrogen fixation in Trichodesmium. *Limnol. Oceanogr.* 46:1249-1260.

39. Paerl HW, Crocker KM, Praetler L-E. 1987. Limitation of N fixation in coastal marine waters: Relative importance of molybdenum, iron, phosphorus, and organic matter availability. *Limnol. Oceanogr.* 32:525-536.

40. Rudolf M, Kranzler C, Lis H, Margulis K, Stevanovic M, Keren N, Schleiff E. 2015. Multiple modes of iron uptake by the filamentous, siderophore-producing cyanobacterium, *Anabaena* sp. PCC 7120. *Mol. Microbiol.* 97:577-588.

41. Berman-Frank I, Cullen JT, Shaked Y, Sherrill RM, Falkowski PG. 2001. Iron availability, cellular iron quotas, and nitrogen fixation in Trichodesmium. *Limnol. Oceanogr.* 46:1249-1260.
64. Barron A, May G, Bremer E, Villarejo M. 1986. Regulation of envelope protein composition during adaptation to osmotic stress in *Escherichia coli*. *J. Bacteriol.* 167: 433-438.

65. Dekker N, Tommassen J, Lustig A, Rg J, Rosenbusch P, Verheij HM. 1997. Dimerization regulates the enzymatic activity of *Escherichia coli* outer membrane Phospholipase A. *J. Biol. Chem.* 272: 3179-3184.

66. Volokhina EB, Beckers E, Tommassen J, Bos MP. 2009. The β-barrel outer membrane protein assembly complex of *Neisseria meningitidis*. *J. Bacteriol.* 191: 7074-7085.

67. Braun M, Silhavy TJ. 2002. Imp/OnA is required for cell envelope biogenesis in *Escherichia coli*. *Mol. Microbiol.* 45: 1289-1302.

68. Genevois S, Steebs L, Rohloll P, Letesson J-J, van der Ley P. 2003. The Omp85 protein of *Neisseria meningitidis* is required for lipid export to the outer membrane. *EMBO J.* 22: 1780-1789.

69. Steebs L, de Cock H, Evers E, Zomer B, Tommassen J, van der Ley P. 2001. Outer membrane composition of a lipopolysaccharide-deficient *Neisseria meningitidis* mutant. *EMBO J.* 20: 6937-6945.

70. Sklar IG, Wu T, Kahne D, Silhavy TJ. 2007. Defining the roles of the periplasmic chaperones SurA, Skp, and DegP in *Escherichia coli*. *Genes Dev.* 21: 2473-2484.

71. Smit J, Kamio Y, Nikaido H. 1975. Outer membrane of *Salmonella typhimurium*: chemical analysis and freeze fracture studies with lipopolysaccharide mutants. *J. Bacteriol.* 124: 942-958.

72. Osborn MJ, Gander JE, Parisi E, Carson J. 1972. Mechanism of assembly of the outer membrane of *Salmonella typhimurium*. Isolation and characterization of cytoplasmic and outer membrane. *J. Biol. Chem.* 247: 3962-3972.

73. Smith SGJ, Mahon V, Lambert MA, Fagan RP. 2007. A molecular Swiss army knife: OmpA structure, function and expression. *FEMS Microbiol. Lett.* 273: 1-11.

74. Pagès J-M, James CE, Winterhalter M. 2008. The porin and the permeating antibiotic: a selective diffusion barrier in Gram-negative bacteria. *Nat. Rev. Microbiol.* 6: 893-903.

75. Gledhill M, Buck KN. 2012. The organic complexation of iron in the marine environment: A review. *Front. Microbiol.* 3: 69.

76. Rudolf M, Stevanovic M, Kranzler C, Pernil R, Keen N, Schleiff E. 2016. Multiplicity and specificity of siderophore uptake in the cyanobacterium *Anabaena*. sp. PCC 7120. *Plant. Mol. Biol.* 92: 57-69.

77. Lutkenhaus JF. 1977. Role of a major outer membrane protein in *Salmonella typhimurium*: chemical analysis and freeze fracture studies with lipopolysaccharide mutants. *J. Bacteriol.* 131: 631-637.

78. Napolitano M, Rubio MÁ, Santamaría-Gómez J, Olmedo-Verd E, Robinson NJ, Luque I. 2012. Characterization of the response to Zinc deficiency in the Cyanobacterium *Anabaena* sp. strain PCC 7120. *J. Bacteriol.* 194: 2426-2436.

79. Fresenborg LS, Graf J, Schätzle H, Schleiff E. 2020. Iron homeostasis of cyanobacteria: advancements in siderophores and metal transporters. pp. 85-117. In *Advances in Cyanobacterial Biology*. Elsevier.

80. Napolitano M, Rubio MÁ, Santamaría-Gómez J, Olmedo-Verd E, Robinson NJ, Luque I. 2012. Characterization of the response to Zinc deficiency in the Cyanobacterium *Anabaena* sp. strain PCC 7120. *J. Bacteriol.* 194: 2426-2436.

81. Calmettes C, Ing C, Buckwalter CM, El Bakkouri M, Chieh-Lin Lai C, Pogoutse A, et al. 2015. The molecular mechanism of Zinc acquisition by the neisserial outer-membrane transporter ZnuD. *Nat. Commun.* 6: 7996.

82. Stork M, Bos MP, Jongers IA, de Kok N, Schilders I, Weynants VE, et al. 2012. An outer membrane receptor of *Neisseria meningitidis* involved in zinc acquisition with vaccine potential. *PLoS Pathog.* 6: e1000969.

83. Barnett JP, Scanlan DJ, Blindauer CA. 2014. Identification of major zinc-binding proteins from a marine cyanobacterium: Insight into metal uptake in oligotrophic environments. *Metallomics* 6: 1254-1268.

84. Crespo-Ferreira R, Cederstrom CR, Pina-Correa AM, Cardoso-Dias JD, de Souza RM. 2012. The role of unchelated Fe in the iron nutrition of phytoplankton. *Limnol. Oceanogr.* 57: 400-404.

85. Haeili M, Speer A, Rowland JL, Niederweis M, Wolschendorf F. 2015. The role of porins in copper acquisition by mycobacteria. *Int. J. Mycobacteriol.* 4: 91-92.

86. Hohle TH, Franck WL, Stacey G, O'Brian MR. 2011. Bacterial outer membrane channel for divalent metal ion acquisition. *Proc. Natl. Acad. Sci. USA* 108: 15390-15395.

87. Puttenhoven JF. 1977. Role of a major outer membrane protein in *Escherichia coli*. *J. Bacteriol.* 131: 631-637.

88. Speer A, Rowland JL, Haeili M, Niederweis M, Wolschendorf F. 2013. Porins increase copper susceptibility of *Mycobacterium tuberculosis*. *J. Bacteriol.* 195: 5133-5140.

89. Eisenhut M. 2015. Manganese homeostasis in cyanobacteria: prevalence of reductive Iron uptake. *Front. Microbiol.* 6: 100573.

90. Hohle TH, Franck WL, Stacey G, O'Brian MR. 2011. Bacterial outer membrane channel for divalent metal ion acquisition. *Proc. Natl. Acad. Sci. USA* 108: 15390-15395.