The Signal Transduction Protein $P_{II}$ Controls Ammonium, Nitrate and Urea Uptake in Cyanobacteria

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$P_{II}$ signal transduction proteins are widely spread among all domains of life where they regulate a multitude of carbon and nitrogen metabolism related processes. Non-diazotrophic cyanobacteria can utilize a high variety of organic and inorganic nitrogen sources. In recent years, several physiological studies indicated an involvement of the cyanobacterial $P_{II}$ protein in regulation of ammonium, nitrate/nitrite, and cyanate uptake. However, direct interaction of $P_{II}$ has not been demonstrated so far. In this study, we used biochemical, molecular genetic and physiological approaches to demonstrate that $P_{II}$ regulates all relevant nitrogen uptake systems in *Synechocystis* sp. strain PCC 6803: $P_{II}$ controls ammonium uptake by interacting with the Amt1 ammonium permease, probably similar to the known regulation of *E. coli* ammonium permease AmtB by the $P_{II}$ homolog GlnK. We could further clarify that $P_{II}$ mediates the ammonium- and dark-induced inhibition of nitrate uptake by interacting with the NrtC and NrtD subunits of the nitrate/nitrite transporter NrtABCD. We further identified the ABC-type urea transporter UrtABCDE as novel $P_{II}$ target. $P_{II}$ interacts with the UrtE subunit without involving the standard interaction surface of $P_{II}$ interactions. The deregulation of urea uptake in a $P_{II}$ deletion mutant causes ammonium excretion when urea is provided as nitrogen source. Furthermore, the urea hydrolyzing urease enzyme complex appears to be coupled to urea uptake. Overall, this study underlines the great importance of the $P_{II}$ signal transduction protein in the regulation of nitrogen utilization in cyanobacteria.

Keywords: $P_{II}$ signaling protein, GlnB, cyanobacteria, nitrogen regulation, nitrate uptake, ammonium uptake, urea uptake, ABC transporters

INTRODUCTION

The emergence of the oxygenic photosynthesis by ancestors of present cyanobacteria (Soo et al., 2017) laid the ground for the evolution of present days life on planet earth. Until today, cyanobacteria occupy a high variety of illuminated habitats, where they represent one of the most abundant primary producers (Whitton, 2012). Accordingly, cyanobacteria are essential contributors to the global carbon cycle. Many cyanobacterial strains have acquired the ability to fix atmospheric nitrogen, making them key players in the global nitrogen turnover.
Nitrogen represents a necessary macronutrient for all living organisms and therefore constitutes an important growth-limiting factor in most ecosystems (Vitousek and Howarth, 1991). The regulation of nitrogen metabolism in cyanobacteria mainly depends on the fine-tuned network of the signal transduction protein PII, the global nitrogen transcription factor NtcA and the NtcA co-activator PipX (Vegapalas et al., 1992; Espinosa et al., 2006, 2007, 2014; Forchhammer, 2008; Luque and Forchhammer, 2008; Forcada-Nadal et al., 2018).

PII signal-transduction proteins are widespread in all three domains of life, where they represent one of the largest and most ancient families of signaling proteins (Chellamuthu et al., 2013; Forchhammer and Luddecke, 2016). PII proteins are involved in the regulation of various nitrogen- and carbon-anabolic processes (Forchhammer, 2004, 2008). Canonical PII proteins are homotrimeric with three characteristic loop regions, designated as B-, C-, and T-loops, which compose the effector molecule binding sites (Cheah et al., 1994; Xu et al., 2003; Forchhammer, 2004, 2008; Llacer et al., 2007; Fokina et al., 2010a; Zhao et al., 2010; Zeth et al., 2014). The large surface exposed T-loop is the prevailing protein interaction module of PII. The PII proteins sense the energy status of the cell by the competitive binding of ADP or ATP (Zeth et al., 2014). Binding of ATP and synergistic binding of 2-oxoglutarate (2-OG) allows PII to sense the current carbon/nitrogen status of the cell (Fokina et al., 2010a). 2-OG is an intermediate of the TCA-cycle that provides the carbon skeleton for inorganic nitrogen incorporation by the glutamine synthetase/glutamate synthase (GS/GOGAT) cycle. Due to this, 2-OG links carbon and nitrogen metabolism and acts as an indicator for the intracellular carbon/nitrogen balance (Muro-Pastor et al., 2001; Fokina et al., 2010a). Besides effector molecule binding, post-translational modification of PII represents a second level of regulation (Forchhammer et al., 2004; Merrick, 2014). Depending on nitrogen availability, cyanobacterial PII can be phosphorylated at the apex of the T-loop at position Ser49 (Forchhammer and Tandeau de Marsac, 1995; Forchhammer and Hedler, 1997). In other prokaryotes, like E. coli, PII is modified by uridylylation in response to nitrogen availability instead of phosphorylation (Jiang et al., 1998). Binding of effector molecules as well as post-translational modifications lead to various PII conformations. Depending on the conformational state, PII can interact with a variety of interaction partners and thereby regulate the cellular C/N balance (Radchenko and Merrick, 2011; Forchhammer and Luddecke, 2016). In cyanobacteria, PII indirectly regulates the global nitrogen control transcriptional factor NtcA through binding of the NtcA co-activator PipX (Espinosa et al., 2007). In common with other bacteria, the cyanobacterial PII protein can control the acetyl-CoA levels by interacting with the biotin carboxyl carrier protein (BCCP) of acetyl-CoA carboxylase (ACC) (Hauf et al., 2016). Furthermore, PII regulates arginine biosynthesis by interacting with the enzyme N-acetylglutamate kinase (NAGK), which catalyzes the rate-limiting step of this pathway (Caldovic and Tuchman, 2003; Heinrich et al., 2004; Llacer et al., 2007; Watzer et al., 2015). If sufficient energy and nitrogen is available, indicated by a high intracellular ATP and low 2-OG level, non-phosphorylated PII interacts with NAGK, enhancing its catalytic efficiency and relieving it from feedback inhibition by arginine (Heinrich et al., 2004; Maheswaran et al., 2004; Llacer et al., 2007). At high intracellular arginine levels, the carbon/nitrogen storage polymer cyanophycin (multi-L-arginyl-poly-L-aspartate) accumulates in Synechocystis sp. strain PCC 6803 (hereafter Synechocystis) (Maheswaran et al., 2006; Watzer et al., 2015). A PII variant was identified with a single amino acid substitution, Ile86 to Asn86 [hereafter referred as PII(I86N)], which constitutively binds NAGK in vitro (Fokina et al., 2010b). Replacing the wild-type PII with a I86N variant in Synechocystis generated a mutant strain, which strongly overproduced arginine and cyanophycin (Watzer et al., 2015). On the other hand, the PII(I86N) strain showed a growth defect in ammonium-supplemented medium (Watzer et al., 2015).

Cyanobacteria use nitrogen sources in a hierarchical order, with ammonium being the preferred nitrogen source. As a consequence, when ammonium is provided together with other suitable nitrogen sources, ammonium will be utilized first (Muro-Pastor et al., 2005). In most natural habitats, the ammonium availability is low, so that high affinity ammonium permeases are required for efficient ammonium uptake (Rees et al., 2006). In Synechocystis, the Amt1 permease is mainly responsible for ammonium uptake (Montesinos et al., 1998). However, elevated intracellular ammonium concentrations are toxic to the cells (Drath et al., 2008; Dai et al., 2014), and therefore, ammonium transport must be tightly controlled. The ammonium transporter family (Amt) is widespread among all domains of life (Wirén and Merrick, 2004). In E. coli, the PII homolog GlnK regulates the ammonium permease AmtB by direct protein–protein interaction. Under ammonium excess conditions, ADP-complexed GlnK blocks an uncontrolled influx of ammonium by inserting the apex of the T-loop into the cytoplasmic exit pores of AmtB (Conroy et al., 2007).

For the assimilation of nitrate, a active nitrate transporter, a nitrate reductase (NR) and a nitrite reductase (NiR) are required (Ohashi et al., 2011). Two types of nitrate transporter systems have been found among cyanobacteria, a high-affinity nitrate/nitrite permease NrtP and the ABC-type transporter NrtABCD (NRT) (Omaña et al., 1993; Luque et al., 1994; Sakamoto et al., 1999; Ohashi et al., 2011). NRT is a bispecific nitrate/nitrite transporter showing high affinity for both substrates (Maeda and Omata, 1997). Intracellular nitrate is first reduced to nitrite by NR and subsequently reduced to ammonium by NiR. Subsequently, ammonium is assimilated in the GS/GOGAT cycle (Flores and Herrero, 1994). Both, NR and NiR, use photosystem I reduced ferredoxin as an electron donor, indicating a coupling of photosynthesis and nitrate assimilation (Manzano et al., 1976; Flores and Herrero, 1994). Addition of ammonium to nitrate adapted cells results in an immediate inhibition of nitrate uptake and a repression of proteins involved in nitrate assimilation (NR and NiR). The ammonium-induced inhibition of NRT is regulated by the PII protein and the C-terminal domain of NrtC (Kobayashi et al., 1997; Lee et al., 1998). Phosphomimetic variants of PII and a PII phosphatase (PphA) deletion mutant, in which PII is constitutively phosphorylated, showed...
ammonium promoted inhibition of nitrate uptake like the wild-type (Lee et al., 2000; Kloft and Forchhammer, 2005). However, this response was abolished in a PII deficient mutant (Kloft and Forchhammer, 2005).

The ability to utilize urea as nitrogen source is widely distributed among bacteria, fungi and algae (Baker et al., 2009; Solomon et al., 2010; Esteves-Ferreira et al., 2018). In common with other bacteria, the cyanobacteria Synechocystis and Anabaena sp. PCC 7120 possess a high affinity urea ABC-type transporter, which is capable of urea import at concentrations lower than 1 µM (Valladares et al., 2002). The gene cluster urtABCDE, encoding all subunits of this ABC-type urea transporter, is transcriptionally controlled by the global transcription factor NtcA (Valladares et al., 2002).

The present study was inspired by the phenotype of the cyanophycin-accumulating strain variant Synechocystis PII(186N), which was impaired in ammonium utilization. Starting with analyzing a possible regulation of the cyanobacterial AmT permease by PII, we found additional evidence for a direct regulation of the nitrate/nitrite transporter NrtABCD and the urea transporter UrtABCDE by the PII signaling protein during this study.

MATERIALS AND METHODS

Cultivation Conditions

Standard cloning procedures were performed in Escherichia coli NEB 10-beta (NEB). Strains were grown in LB-medium at 37°C with constant shaking at 300 rpm.

Synechocystis strains were grown photoautotrophically in BG-11 medium supplemented with 5 mM NaHCO₃ and nitrate, ammonium, or urea as nitrogen source (Rippka et al., 1979). BG-11 agar plates were used by adding 1.5% (w/v) Bacto-agar (Difco), 0.3% (w/v) sodium thiosulfate pentahydrate and 10 mM TES-NaOH pH 8 (Roth) to liquid BG-11 medium. Antibiotics were added when required. Cultivation of liquid cultures for physiological experiments occurred in 50, 100, or 500 mL Erlenmeyer flasks, at 28°C and with constant shaking of 120 rpm. Cultures were continuously illuminated with a photon flux rate of 40–50 µmol photons m⁻² s⁻¹. Growth rates were determined by measuring the optical density at 750 nm (OD₇₅₀).

For induction of nitrogen starvation conditions, exponentially growing cells (OD₇₅₀ 0.4–0.8) were harvested by centrifugation (3,000 × g for 10 min at room temperature), washed and resuspended in BG-11 medium lacking a suitable nitrogen source (BG-11G).

For spot assays (drop plate method), Synechocystis cultures were adjusted to an OD₇₅₀ of 1. A dilution series to the power of 10 was made using BG-11G. Then, 5 µL of every dilution step (10⁰–10⁻⁴) were dropped on BG-11 agar plates. Plates were cultivated at 28°C with constant illumination of 40 µmol of photons s⁻¹ m⁻².

Bacterial Two-Hybrid Assay

Plasmids were constructed by PCR amplification using high-fidelity Q5 polymerase (NEB) and oligonucleotides with overlapping regions. Genomic Synechocystis DNA or plasmids served as templates. PCR fragments were inserted in linearized bacterial two-hybrid vectors pUT18 and pKT25 containing the genes for either the T18 or T25 subunit of the adenylate cyclase CyaA (Karimova et al., 2001) by isothermal, single-reaction DNA assembly according to Gibson et al. (2009). Since the multiple cloning site of pKT25 is located downstream of the T25 subunit, allowing only N-terminal localization of the tag, we constructed plasmid pKT25n (Table 1; pKT25n-fw and pKT25n_rev) to achieve a C-terminal fusion of the tag to the gene of interest. Therefore, plasmid pKT25 was linearized using PCR and the gene of interest was fused upstream of the T25 subunit.

Primers, plasmids, and strains used in this study are listed in Tables 1–3, respectively.

Escherichia coli BTH101 cells were co-transformed with plasmid pUT18 and plasmid pKT25 or pKT25n (Table 2). Plasmids pKT25 or pKT25n contained the possible PII interaction partner fused N- or C-terminal to the T25 subunit. Plasmid pUT18 contains a gene-fusion of the PII-encoding glnB gene or a genetically modified glnB gene containing the I86N mutation [Ile (5′ATC) at codon position 86 to Asn (5′AAC)] (Watzer et al., 2015), the R9L mutation [Arg (5′CGG) at codon position 9 to Leu (5′CTG)] (Fokina et al., 2010a) or the S49D mutation [Ser (5′TCG) at codon position 49 to Asp (5′GAT)] (Lee et al., 2000) with the T18 subunit. The glnB gene and the modified glnB genes were always fused N-terminal with the T18 subunit. Co-transformants were plated on LB-plates (supplemented with 100 µg mL⁻¹ ampicillin and 50 µg mL⁻¹ kanamycin) and cultivated for 2 days at 30°C.

To reduce the level of heterogeneity, five clones from each plate were picked to inoculate 5 mL LB-medium (containing 100 µg mL⁻¹ ampicillin and 50 µg mL⁻¹ kanamycin). Cultures were cultivated overnight at 37°C. Overnight cultures were diluted 1:100 in 3 mL of fresh LB-medium (containing 100 µg mL⁻¹ ampicillin and 50 µg mL⁻¹ kanamycin) and grown to an OD₆₀₀ of 0.7. Three µL of each culture were plated on X-Gal (containing 100 µg mL⁻¹ ampicillin, 50 µg mL⁻¹ kanamycin, 1 mM IPTG, 40 µg mL⁻¹ X-Gal) and MacConkey (containing 100 µg mL⁻¹ ampicillin, 50 µg mL⁻¹ kanamycin, 1 mM IPTG, 1% maltose) reporter plates. Reporter plates were incubated for 3–4 days at 25°C.

Construction and Cultivation of Synechocystis PII-3xFLAG Tag Strains

The previously described pPD-NFLAG and pPD-CFLAG plasmids were used to construct Synechocystis PII (GlnB) fusion proteins with an N- or C-terminal 3xFLAG tag, respectively (Hollingshead et al., 2012; Chidgey et al., 2014). Together with a kanamycin resistance cassette, this construct was inserted in the Synechocystis wild-type genome by homologous recombination, replacing the psbAII gene (Chidgey et al., 2014). Transformants were selected and segregated by kanamycin resistance. For pull-down experiments, 2 L batch cultures were inoculated at OD₇₅₀ = 0.2 in BG-11 medium and propagated as described above, with magnetic stirring at 120 rpm and bubbling with 2% CO₂ (v/v) supplemented ambient air. Cell harvesting was
### Table 1 | Oligonucleotides used in this study.

| Primer | Sequence (5′–3′ direction) |
|--------|----------------------------|
| pKT25n_fw | ACCATGCACGACATCGACATCA |
| pKT25n_rev | CATAGCTTGCTCTGTCGTAAGAATG |
| glnB_fw | TGATGCTGAGAAGCGAGAATGGATATG |
| glnB_rev | CTGCGAGGGAAGCGAGGCAATGGATATG |
| pipX-fw | TCAGAAGAAGCGAGGCAATGGATATG |
| pipX_rev | GCTGCGAGGGAAGCGAGGCAATGGATATG |
| pigX_pKT25n_fw | ACCATGCACGACATCGACATCA |
| pigX_pKT25n_rev | CATAGCTTGCTCTGTCGTAAGAATG |
| amt1_pKT25n_fw | ACCATGCACGACATCGACATCA |
| amt1_pKT25n_rev | CATAGCTTGCTCTGTCGTAAGAATG |
| nrtC_pKT25n_fw | ACCATGCACGACATCGACATCA |
| nrtC_pKT25n_rev | CATAGCTTGCTCTGTCGTAAGAATG |
| urtD_pKT25n_fw | ACCATGCACGACATCGACATCA |
| urtD_pKT25n_rev | CATAGCTTGCTCTGTCGTAAGAATG |
| urtE_pKT25n_fw | ACCATGCACGACATCGACATCA |
| urtE_pKT25n_rev | CATAGCTTGCTCTGTCGTAAGAATG |

Performed at OD<sub>750</sub> = 0.6 by mixing the cultures with ice in a 2:1 ratio (w/w) for rapid metabolic inactivation, and centrifugation at 7,477 × g for 10 min. Cell pellets were subsequently washed with nitrogen-free BG-11<sup>0</sup> at 4°C and snap frozen in liquid nitrogen. For experimental controls, the wild-type strain was similarly cultivated and subjected to pull-down assays followed by mass spectrometry analyses as described below. Two independent replicates were prepared per condition.

### Preparation of Cell Extracts and Anti-FLAG Pull-Down

Frozen cell pellets were washed with 5 mL IP buffer, containing 25 mM MES/NaOH; pH 6.5, 5 mM CaCl<sub>2</sub>, 10 mM MgCl<sub>2</sub>, and 20% (v/v) glycerol and subsequently resuspended in 3 mL IP buffer, including protease inhibitors (complete EDTA-free; Roche) for cell lysis. Therefore, an equal volume of glass beads (0.1–0.15 mm diameter) was added and cells were disrupted using a FastPrep-24 Ribolyser (MP Biomedicals) with five cycles of 20 s at 6.5 m s<sup>–1</sup> and 4°C. After centrifugation for 5 min at 3,314 × g and 4°C, the supernatant was transferred and adjusted with IP buffer to a final volume of 6 mL. For membrane protein solubilization, 1.5% (w/v) dodecyl-β-D-maltoside (DDM; Carl Roth) was added and incubated under agitation for 1 h at 4°C, before insoluble cell debris was removed by centrifugation for 20 min at 20,000 × g and 4°C. The soluble cell extracts (from 2 L cultures) were subjected to the P<sub>II</sub>-FLAG pull-down procedure: for this, 400 µL of resuspended anti-FLAG-M2-agarose (Sigma) was added into empty SPE-columns and washed twice with 1 mL IP buffer, supplemented with 0.04% (w/v) DDM, before
the supernatants were incubated for 5 min and removed by gravity flow. Repeated washing steps with each 5 mL DDM supplemented IP buffer were performed similarly, until the flow through appeared colorless. Elution of coupled proteins from the anti-FLAG resin was performed by incubation in 600 µL DDM supplemented IP buffer containing 0.3 µg/mL 3xFLAG peptide mixtures were subjected to stage tip purification digestion as described elsewhere (Spät et al., 2015). Resulting were resuspended in denaturation buffer for subsequent tryptic acetone/methanol precipitation and resulting protein pellets Eluates from the pull-down workflow were subjected to Analysis, and Data Processing

Proteomics Workflow, NanoLC-MS/MS, and Data Processing
Eluates from the pull-down workflow were subjected to acetone/methanol precipitation and resulting protein pellets were resuspended in denaturation buffer for subsequent tryptic digestion as described elsewhere (Spät et al., 2015). Resulting peptide mixtures were subjected to stage tip purification (Rappsilber et al., 2007). Protein detection by mass spectrometry (MS) was performed as described previously (Spät et al., 2015); in brief, peptide mixtures were loaded onto a 15 cm reversed-phase C18 nanoHPLC column on an EASY-LC system (Proxeon Biosystems) and separated in a 90 min segmented linear gradient. Eluted peptides were ionized by the on-line coupled ESI source and injected in a LTQ Orbitrap XL mass spectrometer (Thermo Scientific). MS spectra were acquired in the positive-ion mode. Per scan cycle, one initial full (MS) scan was followed by fragmentation of the 15 most intense multiply charged ions by collision induced dissociation (CID) for MS/MS scans. The scan range was 300–2,000 m/z for precursor ions at resolution 60,000 and sequenced precursors were dynamically excluded for fragmentation for 90 s. The lock mass option was enabled for real time mass recalibration (Olsen et al., 2005). All raw spectra were processed with the MaxQuant software (version 1.5.2.8) (Cox and Mann, 2008) at default settings. Peak lists were searched against a target-decoy database of Synechocystis sp. PCC 6803 with 3,671 protein sequences, retrieved from Cyanobase (Nakao et al., 2010) (July 2014), plus the sequence of the N- or C-terminal 3xFLAG tagged PII fusion protein (MDYKDDDDKDYKDDDDKDYKDDDDKAAAKKVEAIIRPFKLDYKIALVNAGIVGMVTSEVRGGFRQGKQTERYRSYG VEFQKQLEIIEVQVDEGQVDMVVDKLVSAAARTGEIGDGKIFIS PVDSVVRIRGTEKDEA1 or MKKVEAIIRPKLDYKIALVN AGIVGMVTSEVRGGFRQGKQTERYRSYGVEFQKQLEIIEVQ VDEGQVDMVVDKLVSAAARTGEIGDGKIFISPVDSVVRIRGTEKDEA1 SYKDDDDKDYKDDDDKDYKDDDDK), respectively, as well as 245 common contaminants with the following database search criteria: trypsin was defined as a cleaving enzyme and up to two missed cleavages were allowed; carbamidomethylation of cysteines was defined as a fixed modification.

### TABLE 2 | Plasmids used in this study.

| Plasmid | Tag localization | Description | References |
|---------|-----------------|-------------|------------|
| pPD-nFLAG | N-terminal | Encoding the N-terminal 3xFLAG tag | Hollingshead et al., 2012 |
| pPD-CFLAG | C-terminal | Encoding the C-terminal 3xFLAG tag | Chidgey et al., 2014 |
| pCT25 | | Encoding T25 fragment of adenylate cyclase CyaA (amino acids 1–224) | Karimova et al., 2001 |
| pCT25n | | Derived from pCT25. Upstream of the T25 fragment | This study |
| pUT18 | | Encoding T18 fragment of adenylate cyclase CyaA (amino acids 225–399) | Karimova et al., 2001 |
| pUT18 glnB | N-terminal | Derived from pUT18. Encoding glnB | This study |
| pUT18 glnB (I86N) | N-terminal | Derived from pUT18. Encoding glnB containing the I86N mutation | This study |
| pCT25 pipX | C-terminal | Derived from pCT25. Sequence encoding pipX. | This study |
| pCT25 pipX | | Positive control | This study |
| pCT25 amr1 | C-terminal | Derived from pCT25. Sequence encoding amr1 | This study |
| pCT25 amr1 | | Derived from pCT25. Sequence encoding amr1 | This study |
| pCT25 ntrC | C-terminal | Derived from pCT25. Sequence encoding ntrC | This study |
| pCT25 ntrC | | Derived from pCT25. Sequence encoding ntrC | This study |
| pCT25 nrdD | C-terminal | Derived from pCT25. Sequence encoding nrdD | This study |
| pCT25 nrdD | | Derived from pCT25. Sequence encoding nrdD | This study |
| pCT25 urtD | C-terminal | Derived from pCT25. Sequence encoding urtD | This study |
| pCT25 urtD | | Derived from pCT25. Sequence encoding urtD | This study |
| pCT25 urtE | C-terminal | Derived from pCT25. Sequence encoding urtE | This study |
| pCT25 urtE | | Derived from pCT25. Sequence encoding urtE | This study |
| pCT25 | N-terminal | Derived from pCT25. Sequence encoding urtE | This study |

### TABLE 3 | Strains used in this study.

| Strains | Description | References |
|---------|-------------|------------|
| E. coli NEB 10-beta | Cloning strain | NEB |
| E. coli BTH101 | Bacterial two-hybrid host strain | Euromedex |
| Synechocystis sp. PCC 6803 | Wild type | Pasteur Culture Collection |
| Synechocystis sp. PCC 6803 (I86N) | Genomic PII (I86N) mutant | Watzer et al., 2015 |
| Synechocystis sp. ΔPI | Chromosomal deletion of glnB | Hisbergues et al., 1999 |
| Synechocystis sp. ΔPI + PII-Venus | Synechocystis sp. ΔPI transformed with pVZ232 encoding a PII-Venus fusion | Hauf et al., 2016 |
| Synechocystis sp. ΔPI + PII-Venus | | This study |
and methionine oxidation and protein N-terminal acetylation as variable modifications. Peptide and protein false discovery rates retrieved from MaxQuant were limited to 1%, each. Raw data acquired by mass spectrometry was deposited at the ProteomeXchange Consortium via the Pride partner repository (Vizcaino et al., 2013) under the identifier PXD013411.

**Determination of Nitrate, Nitrite, Ammonium, and Urea in Cell-Free Culture Medium**

To determine the nitrate, ammonium, or urea uptake, exponentially growing cells (OD$_{750} = 0.4–0.8$) were harvested by centrifugation (3,000 x g for 10 min at room temperature) and washed twice with BG-11 medium lacking combined nitrogen sources. Subsequently, the cultures were adjusted to OD$_{750} = 1$ with BG-11 medium. The assays were started by adding either 200 µM NaNO$_3$, 200 µM NH$_4^+$ or 150 µM urea, respectively. Cells were cultivated under constant shaking of 120 rpm and illumination of 40–50 µE. 1 mL aliquots of the cell suspensions were taken and subsequently centrifuged (13,000 x g for 5 min at room temperature) to remove the cells.

For nitrate quantification, the absorbance at 210 nm was measured in the cell-free medium. Since both nitrate and nitrite absorb at 210 nm, the apparent nitrate values were corrected for the presence of nitrite (Kloft and Forchhammer, 2005). Nitrite concentration of the cell-free medium was determined using the modified Griess reaction according to Fiddler (1977).

The ammonium concentration of cell-free medium was measured by using the Nessler reaction (Vogel et al., 1989). Urea was quantified by using the urea nitrogen (BUN) colorimetric detection kit (Invitrogen).

**Microscopy Procedures and Image Evaluation**

Fluorescence microscopy was performed using a Leica DM5500B microscope with a 100×1.3 oil objective. For the detection of Venus fluorescence, an ET500/20× excitation filter and an ET535/30m emission filter were used referred as YFP-channel. To detect cyanobacterial autofluorescence, an excitation filter BP 535/50 and a suppression filter BP 610/75 were used. Image acquisition was done with a Leica DFC360FX black-and-white camera. Captured images were colored with the Leica Application Suite Software (LAS AF) provided by Leica Microsystems. Bright-field images were exposed for 6 ms, Venus fluorescence images for 150 ms and autofluorescence images for 100 ms. Microscope slides covered with dried 2% (w/v) agarose solution were used to immobilize the cells during all microscopical examinations.

Quantitative image evaluation by fluorescence intensity measurements was performed using the open-source software ImageJ (Fiji) (Schindelin et al., 2012). To estimate the fluorescence intensities in different cell compartments, a linear profile of the gray values across the cells through the plasma membrane and cytoplasm was determined. The average gray level of the cytoplasm was subtracted from the maximum gray level of the plasma membrane signal to yield the "Δ fluorescence" value, which indicates the cytoplasmic membrane-localized signal.

An example of this profile data quantification is given in Supplementary Figure 1.

**RESULTS**

**P$_II$ Signaling Mutants Show Increased Sensitivity to Ammonium**

P$_II$ homologs of the GlnK subfamily from heterotrophic bacteria are known to regulate cellular ammonium influx by direct protein–protein interaction with the ammonium permease AmtB (Huergo et al., 2012). The observation of impaired ammonium utilization in the *Synechocystis* strain, which harbors the P$_II$(I86N) variant, suggested that this P$_II$ variant either blocked ammonium uptake or abolished proper ammonium uptake regulation with consequent ammonium sensitivity. The Ile 86 to Asn point mutation in this P$_II$ variant causes a specific alteration of the T-loop conformation, which facilitates interaction with NAGK and thus, results in an over-activation of this P$_II$ target (Fokina et al., 2010b). Conversely, other P$_II$ targets might by negatively affected by the altered T-loop conformation. To gain further insights into the role of P$_II$ in ammonium uptake, we tested ammonium utilization of a P$_II$ deletion mutant (ΔP$_II$) (Hisbergues et al., 1999) and of the ΔP$_II$ strain complemented with P$_II$-Venus (P$_II$-Venus-comp.) (Hauf et al., 2016). The complementation strain was previously generated by introducing a shuttle vector (pVZ322), encoding a P$_II$-Venus fusion under control of the native glnB promoter into the *Synechocystis* ΔP$_II$ mutant (Hauf et al., 2016). The Venus fluorescent protein was fused to the C-terminus of P$_II$ to avoid sterical hindrance of the T-loop.

When grown in presence of 5 mM ammonium in liquid medium, the *Synechocystis* wild-type, the ΔP$_II$ and the P$_II$-Venus-comp. strains showed similar growth rates and ammonium consumption (Figures 1A,B). However, with an elevated ammonium concentration of 10 mM, only the wild-type and P$_II$-Venus-comp. strains could maintain normal growth, while the ΔP$_II$ mutant arrested growth (Figure 1C). Ammonium utilization of wild-type and P$_II$-Venus-comp. were similar under this condition (Figure 1D) while the ΔP$_II$ strain ceased ammonium utilization (Figure 1D). The ammonium-sensitive phenotype of the ΔP$_II$ strain, in particular the response toward 10 mM ammonium, resembled the previously reported phenotype of the *Synechocystis* P$_II$(I86N) strain (Watzer et al., 2015). One possibility to explain the inability of the P$_II$ mutant strains to grow in 10 mM ammonium-supplemented medium is an increased sensitivity toward toxic effects of ammonium. Although ammonium represents the preferred nitrogen source, it becomes toxic for many photosynthetic organisms at higher concentrations (Drath et al., 2008). To find out whether the reduced growth and impaired ammonium utilization results from ammonium intoxication, we assayed the growth of the various strains in the presence of increasing concentrations of ammonium using the spot assay method (Figure 2). Over a range from 5 to 40 mM ammonium, the wild-type and P$_II$-Venus-comp. strains showed a similar growth
and ammonium tolerance. However, the $P_{\Pi}(I86N)$ mutant showed impaired growth at 25 mM while the $\Delta P_{\Pi}$ mutant was already strongly affected at a concentration of 10 mM ammonium (Figure 2). This demonstrated that $P_{\Pi}$ is indeed required to cope with elevated ammonium concentrations. The $P_{\Pi}(I86N)$ mutant can only partially replace the wild-type $P_{\Pi}$ protein, whereas $P_{\Pi}-Venus$ perfectly complemented the $\Delta P_{\Pi}$ strain.

Identification of Novel $P_{\Pi}$ Interaction Partners by FLAG-Tag $P_{\Pi}$ Pull-Down Assays

Uncontrolled influx of ammonium and resulting intoxication of Synechocystis $\Delta P_{\Pi}$ and $P_{\Pi}(I86N)$ mutants is likely a cause for the elevated ammonium sensitivity in these strains. This suggested the involvement of $P_{\Pi}$ in the regulation of
Synechocystis ammonium permeases. In order to verify such an interaction, we performed pull-down assays using a C-terminal 3xFLAG-tagged \( P_{II} \) fusion protein that was expressed in *Synechocystis* under control of the strong \( psbA \) promoter. Protein extracts were prepared in presence of detergent, to solubilize membrane-bound proteins, and were subjected to anti-FLAG-tag immunoprecipitation. The proteome composition of the immunoprecipitate was subsequently analyzed by MS to reveal the proteins that co-purify with FLAG-tag \( P_{II} \). As an initial experiment, the immunoprecipitation was performed with cells grown in BG-11 medium, containing nitrate as unique nitrogen source. To discriminate between \( P_{II} \) interaction partners and unspecific background proteins, the non-transformed wild-type strain was used as an experimental control. Each experiment was further validated by an independent replicate.

As a proof of concept, we were able to identify \( P_{II} \) as the most abundant protein in the \( P_{II} \) 3xFLAG pull-down (thereafter \( P_{II} \) pull-down). Furthermore, some reported interaction partners, such as PipX, were either exclusively identified in the \( P_{II} \) pull-down, or, as in the case of the \( P_{II} \) phosphatase PphA, were strongly enriched compared to the background control (Supplementary Figure 2) (Kloft and Forchhammer, 2005; Espinosa et al., 2007; Forcada-Nadal et al., 2014). The identification of a large number of background proteins in the experiments depends on the wash-stringency, the abundance of individual proteins and the high sensitivity of MS-detection. To distinguish between specific \( P_{II} \)-enriched interaction partners and background, we determined relative protein abundance ratios between the \( P_{II} \) and control experiments by using label free quantification (LFQ) (Cox et al., 2014). Proteins with a significantly higher abundance in the \( P_{II} \) pull-down were identified by significance analysis \((p\text{-value} = 0.01)\). Since both independent replicates from nitrate-grown cultures yielded very similar results, we correlated the data (Figure 3). Using this approach, the major ammonium permease Amt1 (Montesinos et al., 1998) indeed appeared among the significantly enriched proteins, supporting our suggestion that Amt1 represents a \( P_{II} \) binding target in cyanobacteria. Surprisingly, we could also identify subunits of other nitrogen import complexes besides Amt1 among the

![Nitrate growth conditions](image)

**FIGURE 3** | Proteins enriched in anti-FLAG-tag \( P_{II} \) immunoprecipitates from nitrate growth conditions. (Left) Scatterplot of the log\(_2\) transformed LFQ protein ratios \(( P_{II}/\text{control pull-down experiment})\) of 355 quantified proteins detected in two independent replicates. \( P_{II} \) pull-down specific outliers which are significantly enriched in both replicates are indicated in red, and significant outliers enriched only in Replicate 1 are indicated in magenta \((p\text{-value} = 0.01)\). (Right) Heat map representation of the \( P_{II} \)-specific enrichment of nitrogen metabolism related proteins. Log\(_2\) transformed LFQ protein ratios from both replicates are color coded and stars indicate significance. Proteins with missing LFQ ratios, detected exclusively in the \( P_{II} \) but not in the control pull-down, are striped in red/gray.
significantly enriched proteins, in particular subunits of the bispecific NrtABCD nitrate/nitrite and the UrtABCDE urea transporter complexes. For the NRT and URT transporters, all subunits except the periplasmic substrate binding proteins NrtA and UrtA were either exclusively present or strongly enriched in the PII pull-downs. This suggests that the cytoplasmic components of the transporters together with the pore forming subunit are still associated in a complex, detached from the loosely associated periplasmic binding proteins (NrtA or UrtA).

To further validate a potential interaction between PII and the URT transporter complex, we repeated the experiment with cells grown in urea-supplemented medium. In agreement with the results from the PII pull-down at nitrate growth conditions, we detected an enrichment of the PII target PipX, the ammonium permease Amt1, subunits of the NRT nitrate/nitrite transporter, and most prominent, subunits of the URT urea transporter (Supplementary Figure 3). By correlating the data from both independent replicates (Figure 4), it became evident that UrtD and UrtE are the strongest enriched proteins under this condition besides PII. This suggests that the presence of urea in the medium influences the interaction of the URT transporter complex with PII.

Besides proteins associated with nitrogen import, we also identified two small hypothetical proteins to be significantly enriched in all experiments, Sll0944 and Ssr0692. Interestingly, both proteins are under transcriptional control of the global nitrogen regulator NtcA (Giner-Lamia et al., 2017), and reveal opposed dynamics in response to nitrogen availability. Under nitrogen-limiting conditions, Sll0944 was found to be strongly up-regulated at transcriptome and proteome levels, whereas Ssr0692 was strongly down-regulated (Spät et al., 2015; Giner-Lamia et al., 2017). Intriguingly, Ssr0692 seems to be up-regulated under carbon limitation (Battchikova et al., 2010). Overall, our results imply a direct linkage of Sll0944 and Ssr0692 to nitrogen metabolism with a regulatory involvement of PII. Both proteins are currently under investigation to validate a potential interaction with PII and to determine their function.

FIGURE 4 | Proteins enriched in anti-FLAG-tag PII immunoprecipitates from urea growth conditions. (Left) Scatterplot of the log2 transformed LFQ protein ratios (PII/control pull-down experiment) of 110 quantified proteins detected in two independent replicates. PII pull-down specific outliers which are significantly enriched in both replicates are indicated in red, and significant outliers enriched in only one replicate are indicated in magenta (Replicate 1) or orange (Replicate 2) (p-value = 0.01). (Right) Heat map representation of the PII-specific enrichment of nitrogen metabolism related proteins. Log2 transformed LFQ protein ratios from both replicates are color coded and stars indicate significance. Proteins with missing LFQ ratios, detected exclusively in the PII but not in the control pull-down, are striped in red/gray, and n.d. indicates no detection in PII and control pull-down.
Surprisingly, the well-studied P\textsubscript{II} interactor NAGK was not clearly enriched in these experiments. A low percentage of P\textsubscript{II}-NAGK complexes was expected to be present under nitrate or urea growth conditions (Burillo et al., 2004), although the strongest interaction occurs under conditions of nitrogen oversupply. Possibly, the C-terminal 3xFLAG-tag fusion of this P\textsubscript{II} variant could negatively affect complex formation with NAGK. Therefore, pull-down experiments were also performed with an N-terminal 3xFLAG-tag fusion to P\textsubscript{II}. There, we could clearly identify NAGK in the pull-down of nitrate grown cells in two independent replicates (Table 4). The identification of NAGK in both N-terminal P\textsubscript{II}-fusion pull-down replicates implies that the localization of the 3xFLAG-tag has an influence on the interaction between P\textsubscript{II} and its targets. Comparing the results from N- or C-terminal FLAG-tag fused P\textsubscript{II} pull-down experiments can give additional information to identify potential P\textsubscript{II} interaction partners. In contrast to the experiments utilizing the C-terminal FLAG-tag, where subunits of the heterotrimeric urease complex UreABC were not enriched, to our surprise, the N-terminal FLAG-tagged P\textsubscript{II} protein co-purified the entire urease-complex including associated urease accessory proteins D, F, and G. This remarkable finding implies a possible direct interaction between the urease transport complex and the succeeding enzymatic machinery urea metabolism, indicating possible metabolic channeling of urea (Sweetlove and Fernie, 2018).

### Bacterial Two-Hybrid Assay

To further confirm the interaction of P\textsubscript{II} with the identified nitrogen transporters, we performed bacterial two-hybrid assays using the *Bordetella pertussis* adenylate cyclase two-hybrid system (BACTH) (Battesti and Bouveret, 2002). We tested the cytoplasmic localized ATP-binding proteins of the ABC-type transporters NrtC, NrtD, UrtD, UrtE (Omata et al., 1993; Valladares et al., 2002) as well as Amt1 for possible interactions with various P\textsubscript{II} proteins. For each interaction candidate, an N- and a C-terminal fusion to the T25 fragment was constructed. In case of a positive interaction with a P\textsubscript{II}-T18 fusion, CAMP is formed within a CAMP deficient *E. coli* host cell. The interactions were tested by plate assays on X-Gal and MacConkey reporter plates for highest detection sensitivity. The P\textsubscript{II} – PipX and leucine zipper interactions were used as positive controls. P\textsubscript{II}-T18 fusions with an empty pKT25 vector were used as negative controls. Next to wild-type P\textsubscript{II}-T18 fusion, we also included T18 fusions of the P\textsubscript{II}(I86N), P\textsubscript{II}(S49D), and P\textsubscript{II}(R9L) variants to find out how these different P\textsubscript{II} variants, including a phosphomimetic T-loop variant and variants with different effector binding properties would differ in potential P\textsubscript{II} interactions. Figure 5 shows the observed interactions on X-Gal plates in the bacterial two-hybrid assays.

### Table 4

Identified proteins in pull-down experiments at nitrate growth conditions utilizing the N-terminal 3xFLAG-tagged P\textsubscript{II} fusion protein. Displayed are the identified proteins from two independent replicates (Repl. I and Repl. II).

| Protein name/complex | Cyanobase ID | Mol. weight (kDa) | iBAQ intensity |
|----------------------|--------------|------------------|----------------|
| P\textsubscript{II}-N-3xFLAG P\textsubscript{II} | Ssl0707 | 15.43 | 4.2*10\(^7\) 1.1*10\(^9\) |
| PipX | Ssl0105 | 10.45 | 3.1*10\(^6\) 9.1*10\(^7\) |
| PphA | Ssl1771 | 28.47 | 4.2*10\(^4\) 9.1*10\(^5\) |
| NAGK | Ssr1898 | 31.53 | 9.4*10\(^4\) 6.5*10\(^5\) |
| Amt1 | Ssl0108 | 53.58 | 1.2*10\(^8\) 2.9*10\(^9\) |
| NrtA | Ssl1450 | 48.97 | 1.1*10\(^5\) 1.4*10\(^6\) |
| NrtB | Ssl1451 | 29.72 | 8.8*10\(^5\) 5.1*10\(^6\) |
| NrtC | Ssl1452 | 75.10 | 1.4*10\(^6\) 6.1*10\(^6\) |
| NrtD | Ssl1453 | 36.56 | 1.9*10\(^5\) 2.9*10\(^6\) |
| UreA | Ssr0447 | 48.36 | 1.0*10\(^5\) 2.8*10\(^5\) |
| UreB | Ssr1200 | 41.68 | 2.4*10\(^5\) 1.6*10\(^6\) |
| UreC | Ssr1201 | 45.08 | 3.5*10\(^5\) 4.9*10\(^6\) |
| UreD | Ssr1639 | 27.16 | 1.2*10\(^5\) 6.4*10\(^5\) |
| UreF | Ssr1899 | 20.23 | 3.1*10\(^4\) 5.2*10\(^5\) |
| UreG | Ssr0643 | 22.01 | 1.3*10\(^5\) 1.2*10\(^6\) |
| Ssr0692 | Ssr0692 | 18.15 | 8.4*10\(^5\) 9.8*10\(^7\) |
| Ssl0707 | 15.43 | 4.2*10\(^7\) 1.1*10\(^9\) |

Protein intensities were calculated using the MaxQuant iBAQ algorithm (Schwanhausser et al., 2011).
All P$_{II}$ variants showed interaction with the known P$_{II}$ interaction partner PipX. Wild-type P$_{II}$ showed clear interaction with Amt1 C-terminally fused to the T25 fragment, while the variants P$_{II}$(I86N) and P$_{II}$(R9L) did not interact. The loss of interaction of P$_{II}$(I86N) is in agreement with our previous conclusions regarding the impaired ammonium utilization of *Synechocystis* sp. strain P$_{II}$(I86N). The phosphomimetic variant P$_{II}$(S49D) showed weak interaction with Amt1. In case of the nitrate transport ATP-binding subunits NrtC and NrtD, the BACTH assay revealed clear interaction of wild-type P$_{II}$ with both subunits. Intriguingly, P$_{II}$ interacted only with NrtC when it was fused at its C-terminus with T25, whereas in the case of NrtD, P$_{II}$ interacted with the N-terminal fusion. In contrast to wild-type P$_{II}$, the P$_{II}$(R9L) variant was completely unable to interact with any of the Nrt-proteins, while weak interactions were observed with variants P$_{II}$(S49D) and P$_{II}$(I86N). With the ATP-binding subunits of the urea transporter, no interaction was observed with the UrtD protein, neither with wild-type P$_{II}$ nor with any of the tested variants. By contrast, the C-terminal T25 fusion of UrtE showed clear interaction with wild-type and all P$_{II}$ variants. The fact that positive interactions only occurred with selective combinations of P$_{II}$ variants and N-or C-terminal T25 fusions of target proteins supports the conclusion that the observed interactions are indeed specific.

**P$_{II}$ Deletion Mutant and P$_{II}$(I86N) Mutant Excrete Nitrite**

The pull-down and BACTH assays shown above clearly indicated interaction of wild-type P$_{II}$ protein with components of the uptake systems for ammonium, nitrate, and urea. The interaction with Amt1 likely resembles the known interaction of the P$_{II}$ protein GlnK from heterotrophic bacteria with AmtB (Conroy et al., 2007; Gruswitz et al., 2007) and perfectly explains the physiological response of *Synechococcus* PCC 7942 (hereafter *Synechococcus*) and *Synechocystis* P$_{II}$ mutants toward ammonium. Interaction of P$_{II}$ with components of the NrtABCDEF nitrate/nitrite transporter had in fact been suggested previously by several physiological studies, which documented altered nitrate utilization properties in various *Synechococcus* and *Synechocystis* P$_{II}$ mutants (Kobayashi et al., 1997; Lee et al., 1998, 2000; Kloft and Forchhammer, 2005). A characteristic phenotype of a P$_{II}$ deletion mutant grown in the presence of nitrate is the uncontrolled uptake of nitrate and subsequent excretion of nitrite into the medium (Kloft and Forchhammer, 2005). Reduction of nitrate requires two electrons whereas six electrons are needed for the reduction of nitrite to ammonium. Because of the lower electron demand for nitrate reduction, nitrite reduction becomes limiting when insufficient reductant is available. Therefore, when nitrate uptake is uncontrolled, nitrite accumulates and is excreted. We wondered if the *Synechocystis* P$_{II}$(I86N) strain shows a similar nitrite excretion phenotype than the P$_{II}$ deficient mutant when grown on nitrate and if P$_{II}$-Venus complements the nitrite excretion phenotype of the P$_{II}$-deficient mutant.

To answer this question, *Synechocystis* wild-type strain, P$_{II}$(I86N), ΔP$_{II}$ and the P$_{II}$-Venus comp. strains were grown with nitrate as sole nitrogen source under constant illumination (40 µmol photons m$^{-2}$ s$^{-1}$) and samples were removed for nitrite determination. Under these conditions, the various strains showed similar growth rates (Figure 6). Both mutant strains expected to be impaired in proper control of the NRT complex, the ΔP$_{II}$ mutant and the P$_{II}$(I86N) strain, indeed excreted nitrite into the medium. Notably, the P$_{II}$(I86N) mutant excreted much lower nitrite amounts than the P$_{II}$ deletion mutant, in agreement with the partial interaction of the P$_{II}$(I86N) variant observed in the BACTH assays with NrtC and NrtD subunits. By contrast, P$_{II}$-Venus perfectly complemented the nitrite excretion phenotype. Together, these results indicated that the BACTH interactions of P$_{II}$ with the NrtC and NtCD subunits are physiologically meaningful.

**P$_{II}$ Is Responsible for Inhibition of Nitrate Uptake**

To gain further insights in the P$_{II}$-dependent regulation of nitrate uptake, we measured nitrate consumption rates of the various P$_{II}$-mutants under different conditions. For this purpose, cells from exponential phase of growth in standard BG-11 medium were washed and subsequently incubated in BG-11$_0$ medium containing 200 µM NO$_3$ as nitrogen source. Nitrate utilization was quantified by measuring the nitrate concentration in the culture supernatant over time. Under constant illumination of 40 µmol photons m$^{-2}$ s$^{-1}$, *Synechocystis* wild-type and the P$_{II}$-Venus comp. strain showed lower nitrate consumption rates, while both, the P$_{II}$(I86N) and ΔP$_{II}$ mutant, showed higher nitrate removal (Figure 7A). In agreement with our previous results, we observed excretion of nitrite by the P$_{II}$(I86N) and ΔP$_{II}$ mutants (Figure 7B). Interestingly, while the nitrate utilization rate of the P$_{II}$(I86N) and ΔP$_{II}$ mutant appeared almost identical, the two mutants differed in their rate of nitrite excretion. While nitrite excretion of the ΔP$_{II}$ mutant accelerated over time, the P$_{II}$(I86N) initially excreted nitrite faster but then slowed it down over time, explaining the lower amounts of nitrite in the exponentially growing culture (Figure 6) as compared to the ΔP$_{II}$ mutant.

![Figure 6](https://example.com/figure6.png)
Addition of ammonium to nitrate grown cells leads to an immediate inhibition of nitrate uptake. This ammonium-dependent inhibition of nitrate uptake requires the presence of PIPII protein or PIPII phosphomimetic variants (Lee et al., 1998). Here, we tested if the PIPII(I86N) mutant was able to perform the ammonium-dependent nitrate uptake inhibition. For this purpose, nitrate consumption rates were determined after the addition of 2 mM ammonium. Both, Synechocystis wild-type and the PIPII-Venus comp. strains showed a complete inhibition of nitrate uptake in response to ammonium (Figure 7C). By contrast, the PIPII(I86N) and ΔPIPII mutant strains maintained nitrate uptake after ammonium addition. However, nitrate uptake in the PIPII(I86N) mutant was clearly diminished as compared to the ΔPIPII mutant (Figure 7C). The uncontrolled nitrate uptake of the PIPII(I86N) and ΔPIPII mutant caused excretion of nitrite. Due to the diminished uptake of nitrate by the PIPII(I86N) mutant, its nitrite excretion was correspondingly lower as compared to the ΔPIPII mutant (Figure 7D).

In addition to the absence of ammonium, active nitrate transport also requires photosynthetic CO2 fixation.
Nitrogen assimilation is tightly regulated by light/dark transitions. A transition from light to dark causes an immediate inhibition of nitrate uptake and an inhibition of the ammonium assimilating GS (Romero et al., 1985; Marques et al., 1992). To reveal a role of PII in this response, we tested the dark-switch-off of nitrate uptake in the various PII mutant strains. Although the overall nitrate consumption was very low in darkness (Figure 7E) clear differences could be resolved. Notably, the PII(I86N) and ΔPII mutant strains showed slow but constant nitrate uptake as compared to the wild-type and the PII-Venus comp. strain, which did not take up any nitrate. Corresponding to the uptake of nitrate, PII(I86N) and ΔPII mutant excreted small amounts of nitrite (Figure 7F), supporting the notion that the PII(I86N) and ΔPII mutants are unable to completely switch-off nitrate uptake in the dark.

PII Signaling Mutants Show Impaired Urea Utilization

To validate the physiological relevance of the interaction of PII with the UrtE subunit of the urea transporter UrtABCDE, we analyzed urea utilization in the different PII mutants. For this purpose, exponentially growing cells were washed and incubated in BG-11 medium, containing 150 μM urea as sole nitrogen source with constant illumination of 40 μmol photons m⁻² s⁻¹. Under these experimental conditions, Synechocystis wild-type and PII(I86N) mutant strains utilized similar amounts of urea (Figure 8A), whereas both, the ΔPII and PII-Venus comp. strains consumed considerably higher amounts of urea (Figure 8A). The wild-type property of the PII(I86N) variant is in agreement with the interaction of the PII(I86N) protein with the UrtE subunit, which appeared to be T-loop independent according to BACTH assays. Urea is hydrolyzed to CO₂ and two molecules of ammonium by the enzyme urease (Mobley and Hausinger, 1989; Esteves-Ferreira et al., 2018). We wondered if an uncontrolled influx of urea could lead to an excretion of ammonium. Therefore, we measured ammonium concentrations in culture supernatant of those cells, which were incubated with 150 μM urea as the sole nitrogen source. Indeed, we detected ammonium excretion of the ΔPII mutant proportional to its urea utilization (Figure 8B). In contrast, only minor amounts of ammonium could be detected in the culture supernatant of the wild-type, wild-type, and PII-Venus comp. strain (Figure 8B).

FIGURE 8 | Urea utilization and ammonium excretion of various Synechocystis strains as indicated. Exponentially growing and nitrate supplemented Synechocystis cells were washed and resuspended in a medium containing 150 μM urea. Shown are the means of three biological replicates. (A) Urea utilization and (B) ammonium excretion under constant illumination of 40 μmol photons m⁻² s⁻¹. (C) Urea utilization in presence of 2 mM ammonium and under constant illumination of 40 μmol photons m⁻² s⁻¹.
**Synechocystis** utilizes available nitrogen sources in a hierarchical order. In presence of ammonium, the uptake of external nitrogen sources is blocked (Muro-Pastor et al., 2005). To test ammonium-dependent inhibition of urea utilization, we determined urea consumption in presence of 2 mM ammonium. The **Synechocystis** wild-type, **P**II(186N) and **P**II-Venus comp. strains showed a clear inhibition in urea utilization in response to the addition of ammonium. In contrast, urea uptake in the Δ**P**II mutant was completely unaffected by ammonium addition (Figures 8A,C). This demonstrates that **P**II also controls urea uptake, as suggested by the interaction assays.

**P**II Localization Changes Upon Addition of Nitrate, Ammonium, and Urea

Our previous pull-down experiments and bacterial two-hybrid assays clearly showed interaction of **Synechocystis** **P**II with AmtI as well as with nitrate and urea transport components. All interactions appeared to be physiologically relevant. Since all these transporters are localized to the plasma membrane, we suspected that this interaction might affect the cellular localization of **P**II. Therefore, we monitored **P**II localization using the **Synechocystis** **P**II-Venus comp. strain grown under different nitrogen regimes.

Nitrate-replete cells in the mid-exponential growth phase (OD750 ~ 0.5) showed heterogeneous distribution of **P**II-Venus fluorescence in the cell. The majority of cells displayed a strong fluorescence signal in their center and peripheral cell boundary (Figures 9A,B,D). The signal in the center corresponds to the central cytoplasmic space, inside the multiple thylakoid layers. In this region, **P**II might interact with different soluble proteins, like NAGK or PipX. The area with low Venus fluorescence is occupied by the thylakoid membranes. The **P**II-Venus fluorescence at the cell boundary corresponds to the plasma membrane, where Amt1, NrtABCD and UrtABCDE are located (Hahn and Schleiff, 2014) (Figure 9C).

When cells are shifted to nitrogen-depleted conditions, they activate the NtcA regulon, including the *glnB* gene and various uptake systems for nitrogen compounds. During prolonged starvation, they undergo chlorosis, which includes a reduction of thylakoid membranes (Forchhammer and Schwarz, 2018). The chlorotic cells rapidly respond to the addition of combined nitrogen sources. Therefore, investigation of **P**II-Venus fluorescence in chlorotic cells and following the addition of combined nitrogen sources was expected to reveal further insights in the *in vivo* dynamics of **P**II interactions. To monitor the **P**II localization in the chlorotic, nitrogen depleted state, cells were grown to an OD750 of 0.4–0.6, washed, and resuspended in BG-11. After 1 week of nitrogen starvation, the **P**II-Venus signal was evenly distributed throughout the whole cell and its localization on the plasma membrane was not as distinct as during nitrate-supplemented exponential growth (Figure 10A). To test the localization following the addition of different nitrogen sources, 1 week nitrogen starved **Synechocystis** cultures were supplemented with 5 mM NO3−, 5 mM NH4+ or 5 mM urea, respectively. Immediately thereafter, a change in the **P**II localization became visible. The majority of cells showed a clearly visible, distinct plasma membrane localized **P**II-Venus fluorescence, while the remaining cytosol showed homogenously distributed fluorescence. The re-localization of **P**II to the plasma...
membrane appeared most clearly in response to ammonium addition as compared to nitrate or urea (Figures 10B–D).

Since the fluorescence distribution is subject to a certain degree of heterogeneity with the bulk of single cells observed in the microscope, we attempted to quantitatively describe the average distribution of PII-Venus in a representative set of cells. Therefore, a profile of the fluorescence intensities across median cell sections was determined from 54 individual cells per investigated time point. From these profiles, the relative fluorescence intensity distribution between plasma membrane localized signal to the average cytoplasmic signal was determined and displayed as box-plot (Figure 11). Under nitrogen depletion, the majority of the cells showed a higher cytoplasmic PII-Venus localization than plasma membrane localization. Addition of a nitrogen source directly induced a re-localization of PII toward the plasma membrane. In the first 2 h after addition of nitrate or urea, PII-Venus showed a similar localization change (Figures 11A,C,D). Two hours after the addition of nitrate, the PII-derived signal from the plasma membrane decreased again and re-appeared in the cytoplasmic space (Figure 11A), whereas in urea treated cells, PII continued to accumulate at the plasma membrane (Figure 11C). Addition of ammonium induced the most prominent migration of PII-Venus to the plasma membrane in the first hour as compared to nitrate or urea, but thereafter, PII-Venus started to move back to the cytoplasm (Figure 11B).

DISCUSSION

The present investigation unraveled a so far unrecognized global function of the cyanobacterial PII signaling protein in controlling the uptake of the major nitrogen sources ammonium, nitrate and urea. Previous studies already suggested occasionally involvement of PII in the regulation of ammonium and nitrate transport (Kobayashi et al., 1997; Lee et al., 1998, 2000; Kloft and Forchhammer, 2005; Conroy et al., 2007), however, this issue was never addressed systematically using a combined biochemical, genetic and physiological approach. Starting from a series of pull-down experiments using FLAG-tagged PII variants, we observed a putative interaction of PII with various nitrogen transport systems. The interactions were verified by bacterial-two hybrid assays and the biological significance of the interaction was validated by physiological
experiments. Finally, the proposed dynamic interaction of \( P_{P_{II}} \) with cytoplasmic membrane bound transporters was corroborated by fluorescence microscopy.

**Significance Statement**

\( P_{P_{II}} \) signaling proteins play versatile roles in the coordination of carbon- and nitrogen anabolism in prokaryotes and plant chloroplast. In different phylogenetic lineages, \( P_{P_{II}} \) controls a variety of different target proteins. In cyanobacteria, the \( P_{P_{II}} \) paralog GlnK has been shown to control global nitrogen-responsive gene expression by interacting with the transcriptional co-activator PipX. Furthermore, \( P_{P_{II}} \) controls nitrogen storage metabolism in cyanobacteria by regulating the key enzyme of arginine synthesis, \( N \)-acetylglutamate kinase. Finally, a key enzyme in fatty acid metabolism, the acetyl-CoA carboxylase, was shown to be a target of \( P_{P_{II}} \). Several lines of evidences suggested that \( P_{P_{II}} \) might also be involved in the control of ammonium and nitrate uptake, however, direct involvement of \( P_{P_{II}} \) has not yet been shown. In this study, we revealed the interactome of \( P_{P_{II}} \) from the cyanobacterium *Synechocystis* by immunoprecipitating FLAG-tagged \( P_{P_{II}} \) protein. We found prominent enrichment of components of ammonium, nitrate and urea uptake systems. Direct protein–protein interaction was confirmed by bacterial-two hybrid analysis and the physiological relevance was verified by analyzing ammonium-, nitrate-, and urea-uptake in various \( P_{P_{II}} \) mutant strains of *Synechocystis*. This study, therefore, demonstrates that \( P_{P_{II}} \) is the master regulator of the most prominent nitrogen transport systems in cyanobacteria.

**Synechocystis** \( P_{P_{II}} \) **Regulates Ammonium Uptake by Interacting With the Amt1 Ammonium Permease**

In *E. coli*, the \( P_{P_{II}} \) homolog GlnK regulates AmtB by direct protein–protein interaction to control the influx of ammonium (Conroy et al., 2007). In the GlnK-AmtB complex, the nucleotide binding pockets of GlnK are occupied with ADP and the T-loop of GlnK adopts a vertically extended conformation that closes the ammonium gas channel (Conroy et al., 2007; Maier et al., 2011; Forcada-Nadal et al., 2018). Since the conformation of the T-loop changes upon binding of ATP-Mg\(^{2+}\)–2-oxoglutarate, the GlnK-AMT complex only forms under conditions of low 2-OG concentrations, which allows formation of the alternative GlnK-ADP complex (Radchenko et al., 2014).

The present results suggest that in *Synechocystis*, \( P_{P_{II}} \) regulates the major ammonium permease Amt1 in a similar manner than GlnK the AmtB channel. A weak interaction of Amt1 could still be detected with the phosphomimetic variant S49D. The reduced affinity indicates that the negative charge at position 49 reduces the affinity to Amt1. Therefore, S49 phosphorylated wild-type \( P_{P_{II}} \) is expected to have an even weaker affinity to Amt1 (two negative charges of phosphoserine as compared to one negative charge of aspartate). In this respect, phosphorylation of S49 would be analogous to uridylylation of Y31 in enterobacterial GlnK proteins, which prevents AmtB interaction (Conroy et al., 2007). Under conditions of strong \( P_{P_{II}} \) phosphorylation (nitrogen-poor conditions or high CO\(_2\) to nitrate ratio), the Amt1 channel would remain open, allowing unrestricted uptake of ammonium ions. When cells are shifted to excess ammonium conditions, \( P_{P_{II}} \) becomes dephosphorylated (Ruppert et al., 2002) and diminished 2-OG levels allow formation of the \( P_{P_{II}} \)-ADP complex, which then can close the Amt1 pores.

In BACTH assays, the \( P_{P_{II}}(R9L) \) variant was completely impaired in Amt1 interaction. This variant is also unable to interact with NAGK, presumably due to a stabilizing function of the R9 residue at the interface to the binding partner (Fokina et al., 2010a). The failure of this variant to interact with Amt1 indicates that also for Amt1 interaction, the lower face of \( P_{P_{II}} \) with the protruding T-loops is involved in this protein–protein interaction, in agreement with the structure of the GlnK-AmtB complex.

The failure of the \( P_{P_{II}}(I86N) \) variant to interact with Amt1 corresponds to the loss of its affinity toward ADP (Fokina et al., 2010b). \( P_{P_{II}}(I86N) \) has one order of magnitude higher affinity toward ATP than toward ADP and thus, is expected to reside almost exclusively in the ATP-bound state, which would abrogate the interaction with Amt1.

Despite that ammonium uptake in the *Synechocystis* \( P_{P_{II}}(I86N) \) variant is no more under \( P_{P_{II}} \) control, this strain exhibits a higher ammonium tolerance than the \( \Delta P_{P_{II}} \) mutant. This difference could be explained by different ammonium assimilation properties. In the \( P_{P_{II}} \) deletion mutant, the ammonium-scavenging arginine synthesis pathway is not active due to lacking activation of the key enzyme NAGK by \( P_{P_{II}} \) (Maheswaran et al., 2006). By contrast, *Synechocystis* \( P_{P_{II}}(I86N) \) highly enhances NAGK activity, resulting in high intracellular arginine concentrations that leads to accumulation of cyanophycin granules (Watzer et al., 2015). This pathway will foster the metabolic removal of excess ammonium and thus results in increased ammonium tolerance.

Analysis of the subcellular localization of \( P_{P_{II}} \) in response to ammonium stimuli agrees with the above-depicted concept. Only 1 h after the addition of ammonium to chlorotic cells, \( P_{P_{II}} \) shows a strong migration toward the plasma membrane. Providing ammonium to nitrogen-starved cells should cause a strong decrease of the cellular 2-OG levels. This leads to the observed accumulation of \( P_{P_{II}} \) to the cytoplasmic membrane, preventing an excess uptake of too much ammonium. After a while, a new equilibrium will be established leading to a partial re-localization of \( P_{P_{II}} \) to its cytoplasmic targets, such as NAGK. Indeed, *Synechocystis* cells recovering from nitrogen chlorosis start to produce cyanophycin already after a few hours (Watzer and Forchhammer, 2018).

**\( P_{P_{II}} \) Regulates Nitrate Uptake by Interacting With NrtD and NrtC Subunits**

Several previous studies documented an involvement of \( P_{P_{II}} \) in the regulation of nitrate/nitrite uptake. Our data provide novel insights regarding the mechanism of \( P_{P_{II}} \)-mediated regulation of the NrtABCD transporter. It appears that the NrtABCD complex is directly regulated by interaction of \( P_{P_{II}} \) with the cytoplasmic ATPase subunits NrtC and NrtD, as indicated by the co-isolation of wild-type \( P_{P_{II}} \) with both of these proteins. The inability of
the Pll(I86N) complemented strain to regulate nitrate utilization as well as the reduced interaction of Pll(I86N) with NrtC and NrtD subunits in BACTH assays could in principle be explained by the altered T-loop conformation or by the different ligand binding properties of this variant, discussed above. As shown by BACTH assays and by recent in vitro studies (Watzer and Forchhammer, unpublished), the Pll(I86N) variant is indeed able to interact with PipX. This interaction requires a vertically extended conformation of the T-loop (Jlacer et al., 2010; Forcada-Nadal et al., 2018). Although Pll-PipX complexes are favored by the ADP-state of Pll, efficient PipX-Pll interaction also occurs with the Pll-ATP complex. As explained above, the Pll(I86N) variant is trapped in the ATP-bound state. The fact that this Pll variant binds PipX indicates, that the T-loop of Pll(I86N) is flexible enough to adopt various conformations. Therefore, it is likely that formation of the Pll-NrtC or NrtD complex requires the ADP-bound state of Pll, which is disabled in the Pll(I86N) variant. As for NAGK and Amt1, the Pll(R9L) variant is unable to interact with any of the NRT subunits, indicating that Pll interaction involves the lower part of the Pll body, where the T-loop emanates.

The phosphomimetic variant Pll(S49D) shows, albeit weaker, interaction with the NRT subunits. This positive interaction explains previous reports, that Synechocystis and Synechococcus strains expressing these phosphomimetic variants, display a wild-type like regulation of nitrate/nitrite utilization (Lee et al., 2000; Kobayashi et al., 2005). Why the diminished interaction measured by BACTH assays does not properly reflect the physiological behavior of the Pll(S49D) expressing strain (full complementation) could be due to different sensitivities of the assays. Kobayashi et al. (2005) used for the complementation of the Synechocystis deficient mutant plasmid-borne Pll constructs, which cause higher expression levels as in the wild-type genotype. Therefore, a weaker interaction of Pll with the NRT subunits could still be sufficient to inhibit NRT. The mechanism of NRT inhibition by Pll still awaits structure-functional explanation. Our BACTH assays indicate that Pll interacts with both ATPase subunits of NRT. Of these, the NrtC subunit seems to play a particular role: Truncation of the regulatory domain of NrtC directly senses the nitrate-state of the cells (Koropatkin et al., 2006) to prepare NRT for inhibition by the Pll-ADP complex, whereas Pll in the 2-OG-Mg-ATP complex (signaling high C/low N state) would not interact and, therefore, not inhibit NRT. This dual regulatory model combines information from the nitrate status with the global C/N and energy state sensing of Pll to tune NRT activity to the actual need. This model also explains the light-response of NRT: Low energy conditions favor the ADP-complex of Pll, explaining why in the dark, complete switch-off of NRT requires Pll.

**Pll Regulates Urea Uptake by Interacting With the UrtE Subunit**

In addition to Amt1 and NRT, our data also identified the ABC-type urea transporter (UrtABCDE) as a novel Pll target, with the UrtE subunit being the direct interaction partner of Pll. No BACTH interaction was observed for UrtD. This implies that the UrtD protein identified in the FLAG-tag pull-down resulted from pull-down of the entire URT complex. As deduced from the urea-utilization phenotype of the different Pll variant strains, as well as from the properties of BACTH interactions, the interaction of Pll with UrtE is distinct from the mode of Amt1 or NrtC and NrtD interaction. Pll interaction with UrtE seems to be T-loop independent, since the phosphomimetic S49D variant as well as the Pll(I86N) and Pll(R9L) variants interacted in BACTH assays like wild-type Pll. The ability of Pll(R9L) to interact with UrtE indicates that the interaction may involve a region of Pll distinct from the usual interaction face used by NAGK, PipX or Amt1.

In agreement with a different binding mode between Pll and UrtE, the Pll(I86N) variant could substitute wild-type Pll regarding the control of urea utilization. This suggests that the Pll-ATP complex is able to interact with UrtE in a non-conventional manner. The control of urea uptake prevents its futile hydrolysis and the consequent release of ammonium. Although the Pll-Venus variant displayed higher urea uptake rates than the Pll deficient mutant, it could prevent concomitant ammonium excretion, indicating that also the Pll-Venus variant successfully regulates urea utilization. Consistently, all the tested Pll variant strains (wild-type, I86N and Pll-Venus) were able to carry out ammonium inhibition of urea uptake. Careful examination of the data shows that the Pll-Venus strain displayed slightly higher urea uptake rates than the other two strains. It is possible that due to the bulky fluorescent proteins fused to Pll the interaction with Pll does not tune down urea uptake as efficient as in the case of the non-tagged Pll variants. Nevertheless, the Pll-Venus construct is functional, which also agrees with the re-localization of Pll-Venus to the plasma membrane upon addition of urea to nitrogen-starved cells. Under certain conditions, the entire multisubunit urease complex appeared in Pll-FLAG-tag pull-down experiments. This observation deserves further investigation. It suggests either a direct interaction of the urease complex with Pll or a possible metabolic channeling between the urea uptake system and the multisubunit urease complex, leading to a co-immunoprecipitation of the entire super-complex. Further studies are required to reveal the molecular mechanism of urea metabolism and the involvement of Pll in this process.

**CONCLUSION**

All together, the present study has expanded the insights in Pll regulatory interactions in *Synechocystis*. According to this study, the Pll regulatory network includes all relevant nitrogen uptake systems in *Synechocystis*. It is highly probable that the same holds true for other cyanobacteria. In *Synechococcus elongatus*, (Chang et al., 2013) genetic and physiological studies suggested that in addition to NRT, Pll is likely involved in the control of a cyanate ABC transporter (CynABC), which is not
present in *Synechocystis*. Localization of PII-YFP fusions to the cytoplasmic membrane could also be seen in fluorescent images from *S. elongatus* (Espinoso et al., 2018). Control of transport proteins appears to be a highly conserved property of the PII family members, as exemplified by the widely distributed GlnK-Amt interaction. Recently, a more distantly related member of the PII superfamily, SbtA, an accessory component of the sodium-dependent bicarbonate transporter SbtA, was shown to be involved in regulation of bicarbonate component of the sodium-dependent bicarbonate transporter II distributed GlnK-Amt interaction. Recently, a more distantly related member of the PII family members are genetically linked to transport and channel proteins. However, except for GlnK-AmtB, no structures are known for PII-protein complexes with membrane channel proteins. Whether the interactions of the PII members with the channels follow a universal mode, or whether different types of interaction exist, remains to be demonstrated. However, the differences observed between the interactions of PII variants with UrtE, NrtC/D and Amt1 suggest that the interactions could as versatile as observed for the various soluble PII interactors (Forcada-Nadal et al., 2018).

**DATA AVAILABILITY**

The datasets generated for this study can be found in the PRIDE ProteomeXchange consortium, PXD013411.

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