The marine bacterium *Phaeobacter inhibens* secures external ammonium by rapid buildup of intracellular nitrogen stocks

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One sentence summary: *Phaeobacter inhibens* DSM 17395 uncouples NH4⁺-utilization from growth to build up a broad array of intracellular nitrogen stocks, paralleled by the secretion of RTX-like proteins and the antibiotic tropodithietic acid.

**ABSTRACT**

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Reduced nitrogen species are key nutrients for biological productivity in the oceans. Ammonium is often present in low and growth-limiting concentrations, albeit peaks occur during collapse of algal blooms or via input from deep sea upwelling and riverine inflow. Autotrophic phytoplankton exploit ammonium peaks by storing nitrogen intracellularly. In contrast, the strategy of heterotrophic bacterioplankton to acquire ammonium is less well understood. This study revealed the marine bacterium *Phaeobacter inhibens* DSM 17395, a Roseobacter group member, to have already depleted the external ammonium when only $\sim \frac{1}{4}$ of the ultimately attained biomass is formed. This was paralleled by a three-fold increase in cellular nitrogen levels and rapid buildup of various nitrogen-containing intracellular metabolites (and enzymes for their biosynthesis) and biopolymers (DNA, RNA, and proteins). Moreover, nitrogen-rich cells secreted potential RTX proteins and the antibiotic tropodithietic acid, perhaps to competitively secure pulses of external ammonium and to protect themselves from predation. This complex response may ensure growing cells and their descendants exclusive provision with internal nitrogen stocks. This nutritional strategy appears prevalent also in other roseobacters from distant geographical provenances and could provide a new perspective on the distribution of reduced nitrogen in marine environments, i.e. temporary accumulation in bacterioplankton cells.

**Keywords:** *Phaeobacter inhibens* DSM 17395; Roseobacter; heterotrophic bacterioplankton; nitrogen storage; RTX proteins; tropodithietic acid

**INTRODUCTION**

A main originator of the elemental cycles in the world’s oceans is the primary production accomplished by CO$_2$ and N$_2$-fixing phytoplankton and cyanobacteria. Their metabolic activity is responsible for about 50% of the global biomass production (Field et al. 1998) and also shapes the chemical composition of the ocean as conceptualized by the Redfield ratio (Redfield 1958). Transformation (to dissolved organic matter) and remineralization of about half of the net primary production is mainly executed by heterotrophic members of the bacterioplankton (microbial loop) in the sunlit euphotic zone (Azam and Malfatti 2007), even though a considerable share of the primary production is also exported via the biological pump to the deep sea (Laws et al. 2000). Nutrients are not homogeneously distributed across the oceanic water bodies, but range in a spatio-temporal manner from very low (e.g. in the open oceans) to high levels, e.g. in coastal, estuarine and upwelling regions (Falkowski, Barber and Smetacek 1998). The collapse of a seasonal phytoplankton bloom provides a regional nutrient pulse, triggering a dynamic succession of heterotrophic bacterioplankton populations with distinct metabolic proficiencies (Teeling et al. 2012).

The bioavailability of reduced nitrogen controls the productivity in the oceans to a large part (Zehr and Kudela 2011). While marine heterotrophic bacteria can utilize a large variety of different nitrogen sources (dissolved inorganic and organic nitrogen), ammonium (NH$_4^+$) is the preferred inorganic nitrogen source as it can be readily incorporated into cell biomass during anabolism. In the euphotic zone, concentrations of NH$_4^+$ are often in the low $\mu$M range, whereas it can be present at much higher concentrations in coastal and estuarine regions. Especially here, NH$_4^+$ availability is expected to increase further owing to rising agricultural deposition (Li et al. 2016). Heterotrophic bacteria consume $\sim$30% of the available NH$_4^+$ and thereby compete with phytoplankton to exploit ephemeral peak supplies (Wheeler and Kirchman 1986; Kirchman 1994). The smaller sized bacteria grow faster and import NH$_4^+$ more rapidly (Goldman and Dennett 2001), implicating a tight coupling of NH$_4^+$ acquisition with growth and carbon source consumption, respectively (e.g. Goldman and Dennett 1991; Beg et al. 2012). In contrast, algae can uncouple growth from NH$_4^+$ uptake due to their long known capacity for transient nitrogen storage (Dortch et al. 1984; Admiraal, Peletier and Laane 1986).

Roseobacters constitute a metabolically diverse group within the alphaproteobacterial Rhodobacterales and can account for $\sim$20% of coastal and $\sim$15% of mixed-layer ocean bacterioplankton communities. They inhabit coastal and open oceans, sea ice and the sea floor and occur planktonic as well as particle-associated (Buchan, Gonzalez and Moran 2005; Wagner-Döbler and Biebl 2006). Notably, roseobacters were also found to associate with marine algae during phytoplankton blooms, thereby playing an important role in the recycling of seasonal peaks of phytoplankton-derived biomass (Teeling et al. 2012; Buchan et al. 2014; Luo and Moran 2014). A recent study examining effective population size as a measure of the adaptive potential suggested roseobacters to stay competitive in changing ocean habitats (Luo et al. 2014). The study organism, *Phaeobacter inhibens* DSM 17395, is a nutritionally versatile representative of roseobacters with a complex metabolism and regulation (Thole et al. 2012; Drüppel et al. 2014), and is apparently specialized for growth on biotic (e.g. algae) and abiotic surfaces in coastal areas (Gram et al. 2015).

A recent growth physiological study covering a wide concentration range of NH$_4^+$ and PO$_4^{3-}$ revealed *P. inhibens* DSM 17395 to be well adapted to fluctuating availability of inorganic N and P, and to grow optimally at N:P supply ratios ($\sim$50–120) markedly above Redfield (Trautwein et al. 2017). Considering the important role of NH$_4^+$ for growth of *P. inhibens* DSM 17395 in particular and the general relevance of this macronutrient for the productivity of marine heterotrophic bacteria, the present study aims at investigating the NH$_4^+$ acquisition and processing strategies of *P. inhibens* DSM 17395 as a well-studied representative of the Roseobacter group.

**MATERIALS AND METHODS**

Media, cultivation, sample retrieval and harvesting of cells

*Phaeobacter inhibens* DSM 17395 (originally deposited as *P. gallaeciensis* DSM 17395) (Buddrus et al. 2013) was obtained from the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ, Braunschweig, Germany) and maintained in our laboratory. Each growth experiment was started from a glycerol stock (stored at $-80^\circ$C) prepared from cultures of *P. inhibens* DSM 17395 grown in marine broth (MB) medium. Revival of stock cultures proceeded in defined mineral medium supplemented with 11 mM glucose and residual glycerol/MB were eliminated by a dilution series (up to $10^{-3}$) (Zech et al. 2009). This was followed...
by two successive passages in the same medium supplemented with different NH₄⁺ concentrations (0.4, 0.8 or 1.8 mM; added from a sterile NH₄Cl stock solution). Each culture was inoculated with cells at the respective half-maximal optical density (OD₆₀₀ nm). Cultivation was routinely performed in Erlenmeyer flasks (250 mL or 1 L), containing 50 or 250 mL of medium, respectively. All cultures were incubated on a rotary shaker (100 rpm) at 28 °C in the dark and growth was monitored by measuring the OD at 600 nm (UVmini-1240; Shimadzu, Duisburg, Germany). All cultivation experiments and subsequent analyses are based on three to six biological replicates. Chemicals used were of analytical grade.

Selected bacterial strains from the marine bacterioplankton (Table S1, Supporting Information, 12 in total) as well as Escherichia coli were obtained from the DSMZ. Each strain was tested for growth under the identical cultivation conditions as described above for P. inhibens DSM 17395 (for further experimental details see Table S1, Supporting Information). The only exception was E. coli that was grown in standard M9 minimal medium (Sambrook and Russell 2001), accordingly supplemented with glucose and ammonium. In addition to four other roseobacters, only R. blandensis and E. coli grew reproducibly enough and sufficiently well in the undisturbed marine media to perform detailed growth experiments.

Samples for the determination of NH₄⁺ and glucose concentrations, as well as for flow cytometric analyses were retrieved from each culture in regular intervals. Each sample (1.8 mL culture broth) was centrifuged (25,245 × g, 15 min, 4 °C) and the supernatant transferred into new tubes. The samples for NH₄⁺ determination were acidified with 10 μL of 0.1 M HCl. All samples were stored at −20 °C until further analysis. For cellular dry weight (CDW), cultures were harvested by centrifugation (11,300 × g, 20 min, 4 °C), washed twice with 50 mM ammonium acetate and then resuspended in 300 μL 50 mM ammonium acetate and transferred into predried and weighed 1.5 mL tubes. The CDW was determined by gravimetric analysis of tubes after drying to constant weight at 60 °C. Strong correlations (R² ≥ 0.96) between OD and CDW were observed. CDW samples were stored at room temperature and used for determination of cellular carbon (C) and nitrogen (N) contents. To analyze the cellular composition (N-containing macromolecules, proteome, metabolome and lipidome, 3-6 replicate cultures (with 250 mL medium each) were harvested per sampling point and analysis type from 14 to 96 h after inoculation in order to cover all growth phases. Reproducible growth of each harvested culture was controlled by continued incubation of a 20 mL subculture transferred to 100 mL Erlenmeyer flasks prior to harvesting. At low ODs, several cultures were pooled to obtain sufficient cell material for chemical and compositional analyses of biomass. Cell harvesting included centrifugation (14,300 × g, 20 min, 4 °C) and washing steps specific for the various analysis types: cell pellets for (i) chemical analysis of biomass ( aliquoted to cell pellets of 1, 2 and 5 mg dry cells) were washed twice with 3.7% (w/v) NaCl, (ii) proteome analysis were washed twice with Tris/HCl buffer (100 mM Tris, 5 mM MgCl₂ × 6 H₂O, adjusted to pH 7.5) and (iii) analysis of the intracellular metabolome or lipidome were washed twice with 3.7 or 0.9% (w/v) NaCl, respectively. Each cell pellet was immediately shock frozen in liquid N₂ and stored at −80 °C until further analysis. Samples (5–96 h after inoculation) for analysis of the exoproteome (200 or 400 mL) were retrieved from the culture supernatant after the first centrifugation step, immediately filtered (0.2 μm CA; Sartorius, Göttingen, Germany), and then directly processed (see below).

Chemical analyses

The concentration of NH₄⁺ in cell-free culture supernatants was determined according to the method reported by Chaney and Marbach (1962). For calibration of the assay, 0–200 μM NH₄Cl was dissolved in defined (NH₄⁺-free) mineral medium. The procedure of the assay was as follows: 800 μL of sample or calibration standard were placed in 1.5 mL reaction tubes and 100 μL reagent A (3 g phenol, 3 mg sodium nitroprusside; both dissolved in 100 mL H₂O, final pH 5.3; stored for up to 2 weeks at 4 °C–8 °C) were added. After mixing, 100 μL of reagent B (2 g NaOH, 0.5 mL 13% (w/v) NaOCl; dissolved in 100 mL H₂O, final pH 13.5; stored at 4 °C–8 °C) were added. Following further mixing, the assay mixture was incubated at room temperature for 1 h in the dark and then centrifuged (20,000 × g, 4 °C, 1 min). The supernatant was transferred into a cuvette and the absorption was measured at 635 nm (UVmini-1240). The limit of quantitation was 5.1 μM.

The assay was adapted to lower sample volumes and higher throughput (96-well plates) by employing a microplate reader (FLUOstar OPTIMA; BMG Labtech GmbH, Ortenberg, Germany) (Ruppersberg et al. 2017). The limit of quantitation in this case was 36 μM NH₄⁺.

The concentration of glucose in cell-free culture supernatants was determined by HPLC analysis. The system consisted of an UltiMate 3000 Rapid Separation LC (ThermoFisher Scientific, Gormanering, Germany) equipped with a Eurokate separation column (8 × 300 mm, 10 μm bead size; Knauer, Berlin, Germany) temperature controlled at 75 °C and a refractive index (RI) detector. The eluent was composed of 5 mM H₂SO₄ and administered at a flow rate of 1.2 mL min⁻¹. Calibration was performed with a glucose standard (retention time at 8 min) diluted in mineral medium ranging from 0.01 to 15 mM.

The cellular C and N content was determined by subjecting samples (1–5 mg dry cells) to high temperature oxidation, sequential heat-dependent release from adsorption columns and thermal conductivity detection using a Vario EL cube (Elementar Analysensysteme GmbH, Hanau, Germany) essentially as described before (Zech et al. 2013a).

The cellular content of DNA or RNA was quantified independently in cell pellets with the ‘Genomic DNA from tissue kit’ or the ‘NucleoSpin RNA kit’ following the manufacturer’s instructions (both from Macherey–Nagel, Düren, Germany). Isolated nucleic acids were measured spectrophotometrically in a tray cell (Hellma, Müllheim, Germany) at 230, 260 and 280 nm (UV-1800; Shimadzu) to determine the concentration and purity of DNA and RNA. The cellular protein content was quantified in cell pellets after alkaline lysis of cells in 0.5 M NaOH and 2% (w/v) SDS for 10 min at 95 °C in a thermoblock mixing at 600 rpm. This was followed by a 10 min incubation in an ultrasonic bath at 60 °C, and centrifugation of the lysate (20 000 × g, 5 min, 20 °C). The protein concentration in the supernatant was determined with the DC Protein Assay Kit II (BioRad, Munich, Germany) according to the manufacturer’s instructions, using bovine serum albumin (BSA; Sigma-Aldrich, Hamburg, Germany) as calibration standard. Each of four biological replicate samples for DNA, RNA and protein was measured three times.

Poly(3-hydroxybutanoate) (PHB) was analyzed in cell pellets (5–10 mg dry weight each) essentially as described (Trautwein et al. 2008). The total amount of PHB was quantified by GC in relation to methyl esters obtained from purchased sodium salt of 3-hydroxybutanoic acid.
The concentration of tropodithietic acid (TDA) was estimated using an indirect spectrophotometric assay that is based on the reported color shift (from transparent to brown) upon binding of TDA to ferric/ferrous iron present in the medium (D’Alvise et al. 2015).

Flow cytometry

The DNA, RNA, protein and PHB content of cells was also determined with a FACSCalibur flow cytometer (Becton Dickinson, Franklin Lakes, NJ, USA). Immediately after sample retrieval, cells were fixed with formaldehyde and after several washing steps, stored at −20 °C (Trautwein et al. 2012). Fixed cells were diluted 1:150 in phosphate-buffered saline (PBS) (145 mM NaCl, 8 mM Na2HPO4, 1.4 mM NaH2PO4, adjusted to pH 7.4) and stained with SYBR green I (for DNA; Molecular Probes, Eugene, OR, USA), SYTO RNASelect green fluorescent cell stain (for RNA; Molecular Probes), Flamingo (for proteins; Bio-Rad) or Nile red (for PHBs; Sigma-Aldrich). Nile red (final concentration 5 μg mL−1) staining was carried out at room temperature in the dark for 30 min. If necessary, samples were further diluted 1:2 with PBS prior to analysis. Measurements (irrespective of the type of staining) were standardized to the fluorescence of suitable fluorescent latex beads (pink or yellow-green; Molecular Probes) added to each sample prior to analysis. A total of 70,000 (40,000 only at very low cell densities) events were recorded per measurement, and each sample (3–4 biological replicates per sampling point) was measured in triplicate. Data acquisition and analysis were performed with CellQuest Pro software (Becton Dickinson). Data analysis was manually controlled for each sample and included ~85%–90% of the total cell events (core population) to reduce the influence of morphological changes of a minor cell fraction during growth on average fluorescence values.

Proteomic analyses

The proteomic strategy targeted intracellular soluble (by 2D DIGE and whole cell shotgun analysis), membrane and extracellular proteins. Protein concentrations in the soluble intra- and extracellular protein fractions were determined with the method described by Bradford (1976), while that introduced by Lowry et al. (1951) was used for the membrane proteins (RC DC Protein Assay Kit II; Bio-Rad). In both cases, BSA (Sigma-Aldrich) was used as calibration standard.

2D DIGE

Cell pellets (from 4–6 independent cultures per sampling point) of P. inhibens DSM 17395 were resuspended in 1 mL lysis buffer (7 M urea, 2 M thiourea, 30 mM Tris, 4% CHAPS; final pH 8.5) and cells were disrupted using the Plus One Sample Grinding Kit (GE Healthcare, Munich, Germany) as described by Gade et al. (2003). The pre-electrophoretic labeling with CyDyes (GE Healthcare) was carried out as described before (Gade et al. 2003), with the exception that 50 μg of each protein sample were labeled with 200 pmol of commercial fluorescent dye (Druppel et al. 2014). In the present study, protein extracts of cells harvested 30 h after inoculation served as reference state and were labeled with Cy5. Protein extracts of cells harvested at all other sampling points served as individual test states and were each labeled with Cy3. The internal standard was composed of equal amounts of all protein extracts and was labeled with Cy2. Isoelectric focusing (IEF) was carried out as previously described (Zech et al. 2009), employing a combination of narrow (3–5.6) and wide (3–11) range non-linear (NL) pH gradients (IPG strips, 24 cm; GE Healthcare) and a protein load of 150 μg protein per IPG strip. Equilibration of IPG strips with buffers containing dithiothreitol (DTT) or iodoacetamide was performed as described by Görg, Weiss and Dunn (2004). Second dimension protein separation was achieved with 12.5% (v/v) acrylamide gels and the EttanDalt II system (GE Healthcare) as described before (Gade et al. 2003). Samples from four independent cultures (biological replicates) were analyzed per sampling point.

2D DIGE gels were digitalized with a Typhoon 9400 laser scanner (GE Healthcare) set to a resolution of 100 μm. Gel images were cropped with the ImageQuant software (version 7.0; GE Healthcare) to obtain good alignment between the pH 5–11 (from pH 3–11 gel) and the pH 3–5.6 regions. Cropped gel images were sorted into two different work packages (pH 5–5.6 or pH 5–11) for image analysis with the DeCyder 2D software (version 7.0; GE Healthcare), which was performed as described previously (Zech et al. 2009). Differentially abundant protein spots had to fulfill the following criteria: average ratio (fold change in protein abundance) of <−1.5 or >1.5, ANOVA p-value of <0.05, t-test value of <10−4 and matched in at least 75% of the gel images in the work package.

Protein spots with changed abundances were excised with the EXQuest spot cutter (Bio-Rad) from separate, preparative 2DE gels (300 μg protein load) stained with colloidal Coomassie Brilliant Blue (cCBB) (Doherty et al. 1998). Each protein spot was excised from two replicate gels and processed independently. Subsequent washing steps, in-gel tryptic digestion, spotting onto Anchorchip steel targets (Bruker Daltonik GmbH, Bremen, Germany) and mass spectrometric analysis with an UltraflexXtreme MALDI-TOF/TOF mass spectrometer (Bruker Daltonik GmbH) was performed as described recently (Koßmehl et al. 2013). Proteins were identified by PMF and PFF mapping, respectively, using an in-house Mascot server (version 2.3; Matrix Science, London, UK) via the ProteinScape platform (version 3.1; Bruker Daltonik GmbH). Acquired MS and MS/MS spectra were searched against the translated genome of P. inhibens DSM 17395 (Thole et al. 2012) as described by Koßmehl et al. (2013).

Whole cell shotgun

Samples from three independent cultures were subjected to shotgun proteomic analysis per sampling point. Essentially, suspending cells in lysis buffer (7 M urea, 2 M thiourea, 30 mM Tris, pH 8.5), cell breakage, removal of cell debris, reduction with DTT, alkylation with iodoacetamide and tryptic in-solution digestion were performed as described previously (Zech et al. 2013a). Total peptide mixtures were separated with an Ultimate 3000 nanoRSLC system (ThermoFisher Scientific) in a trap column setup equipped with an analytical column (C18; pore size, 100 Å; bead size, 2 μm; inner diameter, 75 μm; length, 25 cm; ThermoFisher Scientific), as previously described (Zech et al. 2013a). Continuous analysis of the eluent was performed with an online-coupled electrospray ion source (captive spray; Bruker Daltonik GmbH) ion trap mass spectrometer (amazon ETD; Bruker Daltonik GmbH) operated in positive mode with a capillary current of 1.3 kV and dry gas flow of 3 L min−1 at 150 °C. Per full scan MS, 20 MS/MS spectra of most intense masses were acquired. Active precursor exclusion was set for 0.2 min. Protein identification was performed with ProteinScape as described above, allowing for a mass difference of 0.3 Da and applying a target decoy strategy (false discovery rate <1%).

Membrane proteins

Membrane protein-enriched cell fractions were prepared from two biological replicates per sampling point. Preparation of
membrane proteins involved cell disruption by French Press (Sim-Amino Ltd., Rochester, NY, USA), treatment with ice-cold carbonate and solubilization of membrane proteins with hot SDS as previously described (Trautwein et al. 2012; Kößmehl et al. 2013). Electrophoretic separation of similar amounts of membrane proteins was conducted with 7 cm long, 1 mm thick 12.5% (v/v) SDS-PAGE gels using the Mini-Protein Tetra System (Bio-Rad) (Zech et al. 2013a). Gels were stained with cCBB and following digitalization, each sample lane (from two different gels) was cut into four slices. Each slice was then cut into ∼1 mm² pieces, which were washed, reduced, alkylated and tryptically digested as described (Zech et al. 2013a). Separation of peptide mixtures and mass spectrometric analysis were performed with a nanoLC-ESI-MS/MS system and protein identification by means of the ProteinScape platform as described above for the shotgun analysis of soluble proteins.

Exoproteins

Extracellular proteins were precipitated by addition of trichloroacetic acid (final concentration 10% (v/v)) added to the filtered culture supernatants (200 or 400 mL) directly after centrifugation. After incubation for 12–14 h at 4 °C, the precipitated proteins were collected by centrifugation (17,700 × g, 1 h, 4 °C) and washed six times with 1 mL 96% (v/v) ethanol (20,000 g, 15 min, 4 °C). The protein pellets were dried and immediately resuspended in 400 μL lysis buffer (pH adjusted to 8.0). One final round of centrifugation (20,817 g, 10 min, 12 °C) was carried out to remove insoluble material. Protein extracts were aliquoted, then immediately frozen in liquid N2 and was carried out to remove insoluble material. Protein extracts were dried and stored at −20 °C. Prior to analysis, samples were reconstituted in methanol directly before measurement. The crude lipid extracts were analyzed by a UPLC-qTOF mass spectrometer (Acquity UPLC; Waters, Milford, USA; maXis MS, Bruker Daltonik GmbH) using a Cortecs UPLC C18 column (pore size, 90 Å; particle size, 1.6 μm; length 150 mm; inner diameter, 2.1 mm; Waters) and applying 1.5 min isocratic elution at 32% B, followed by a gradient leading to 97% B in 21 min (solvent A: 60/40 acetonitrile/water, 10 mM ammonium formate, 0.1% formic acid; solvent B: 90/10 n-propanol/acetonitrile, 10 mM ammonium formate, 0.1% formic acid) and a 4 min plateau at 9% B (column temperature 40 °C, sample temperature 5 °C, flow 0.225 mL min⁻¹, injection volume 10 μL). Mass spectra were acquired in positive electrospray mode. Parameters of the qTOF instrument were set for best resolution and mass accuracy in the mass range of approximately 500–900 Da. Analysis of a phospholipids standard solution was included in the analytical workflow as quality control (Boncompain et al. 2014). Genedata Refiner (Genedata AG, Basel, Switzerland) was applied for chromatographic and mass spectrometric calibration and alignment. 2656 lipid clusters were detected, for which MassTRIX annotation (Suhre and Schmitt-Kopplin 2008) (max. error <0.01 Da) was performed. Statistical analysis was done in Genedata Analyst (Genedata), Simca P (Umetrics, Malmö, Sweden) and HCE 3.0 (Human-Computer Interaction Lab, Maryland, MD, USA).

Identification was achieved by MS² experiments using either targeted quadrupole extraction for lipid species containing nitrogen and showing a decreasing abundance over incubation time or in auto MS mode at 40 eV, resulting in fragmentation of the dominant ion species.

Intracellular metabolome

Five to six samples per harvesting time point were prepared for gas chromatography-mass spectrometry (GC-MS) analysis as previously described (Zech et al. 2013a). Essentially, cell pellets were resuspended in 1.5 mL of ribitol-containing ethanol and cell breakage was achieved by ultra-sonication. Following removal of hydrophobic compounds by extraction with chloroform and centrifugation (3900 × g, 5 min, 4 °C), samples were dried and stored at −20 °C. Prior to analysis, samples were derivatized according to the two-step procedure previously described (Zech et al. 2009). Essentially, samples were first treated with methoxyamine hydrochloride and then silylated using N-methyl-N-(trimethylsilyl)-trifluoroacetamide (MSTFA).

Derivatized samples were analyzed using a Leco Pegasus 4d GC × GC TOF mass spectrometer (Leco Instrumente, Mönchengladbach, Germany). The instrument was operated in GC-TOF mode, equipped with a MPS 2 XL autosampler connected to a programmed temperature vaporizing (PTV) injector (Gerstel, Mülheim a.d. Ruhr, Germany) and calibrated with perfluorotributylamine. Samples (1 μL) were injected in splitless mode and vaporized along the following temperature profile: 70 °C for 0.02 min, ramping by 12 °C s⁻¹ to 330 °C, and then constant for 5 min. The GC instrument (7890 Agilent GC, Agilent, Santa Clara, USA) was equipped with a ZB-5MS column (particle size, 25 μm; length, 30 m; inner diameter 0.25 mm; Phenomenex, Aschaffenburg, Germany) operated with helium (1.2 mL min⁻¹) and the following temperature profile: 70 °C for 1 min, ramping by 10 °C s⁻¹ to 330 °C, and then constant for 3 min. The transfer line to the mass spectrometer was set to 275 °C, the ion source to 250 °C and the detector voltage 200 V higher than the output of the automatic tuning. Following a solvent delay of around 300 s, full mass spectra were collected from 45 to 600 m/z at 8 scans s⁻¹ using the ChromaTOF software (version 4.24; Leco Instrumente).

Data analysis was done with the Metabolite Detector software (Hiller et al. 2009) as described before (Zech et al. 2009) and included normalization by the internal standard ribitol and cell mass.

Lipidome

Bacterial cell mass was reconstituted in 1.5 mL methanol; 5 mL of methyl-tert-butyl ether (MTBE) was added and lipid extraction was performed for 1 h in an ultra-sonic bath on ice (Matyash et al. 2008). The organic solvents were evaporated and the samples were reconstituted in methanol directly before measurement.

The crude lipid extracts were analyzed by a UPLC-qTOF mass spectrometer (Acquity UPLC; Waters, Milford, USA; maXis MS, Bruker Daltonik GmbH) using a Cortecs UPLC C18 column (pore size, 90 Å; particle size, 1.6 μm; length 150 mm; inner diameter, 2.1 mm; Waters) and applying 1.5 min isocratic elution at 32% B, followed by a gradient leading to 97% B in 21 min (solvent A: 60/40 acetonitrile/water, 10 mM ammonium formate, 0.1% formic acid; solvent B: 90/10 n-propanol/acetonitrile, 10 mM ammonium formate, 0.1% formic acid) and a 4 min plateau at 9% B (column temperature 40 °C, sample temperature 5 °C, flow 0.225 mL min⁻¹, injection volume 10 μL). Mass spectra were acquired in positive electrospray mode. Parameters of the qTOF instrument were set for best resolution and mass accuracy in the mass range of approximately 500–900 Da. Analysis of a phospholipids standard solution was included in the analytical workflow as quality control (Boncompain et al. 2014). Genedata Refiner (Genedata AG, Basel, Switzerland) was applied for chromatographic and mass spectrometric calibration and alignment. 2656 lipid clusters were detected, for which MassTRIX annotation (Suhre and Schmitt-Kopplin 2008) (max. error <0.01 Da) was performed. Statistical analysis was done in Genedata Analyst (Genedata), Simca P (Umetrics, Malmö, Sweden) and HCE 3.0 (Human-Computer Interaction Lab, Maryland, MD, USA).

Identification was achieved by MS² experiments using either targeted quadrupole extraction for lipid species containing nitrogen and showing a decreasing abundance over incubation time or in auto MS mode at 40 eV, resulting in fragmentation of the dominant ion species.

Modeling

A cell population model was fitted to experimentally determined values for OD, NH₄⁺ and glucose concentrations, as well as to N:P ratios of biomass. The applied model described by Droop (1968) was adopted to include the relevant solute uptake systems for NH₄⁺ and glucose, respectively. NH₄⁺ uptake is described by
Monod kinetics (Monod 1942) as follows

$$\rho^A(A) = \frac{\rho_{\text{max}} A}{\rho_{\text{g}}^2 + A}$$

with NH₄⁺ concentration A, maximum uptake rate \(\rho_{\text{max}}\), and half-saturation constant \(\rho_{\text{g}}^2\). Glucose uptake \(\mu^G(G)\) is described analogously with glucose concentration \(G\), maximum uptake rate \(\rho_{\text{max}}^G\) and half-saturation constant \(\rho_{\text{g}}^G\).

The modified Droop equations (see below) were applied to describe the dynamics of the bacterial population in the batch culture: the population density \(P\), extracellular NH₄⁺ concentration \(A\), extracellular glucose concentration \(G\) and cell quota \(Q\) (reflecting the nitrogen content per cell) vary according to

$$\dot{P} = \left[\mu(Q,G) - m\right]P$$
$$\dot{A} = -\rho^A(A)P$$
$$\dot{G} = -\beta \mu(Q,G)P$$
$$\dot{Q} = \rho^A(A) - \alpha \mu(Q,G)Q$$

where

$$\mu(Q,G) = \min\left\{\mu_A(Q), \mu_G(G)\right\}$$

$$\mu_A(Q) = \mu_{\text{max}}^A \left[1 - \frac{Q_{\text{max}}}{Q}\right]$$

$$\mu_G(G) = \frac{\rho_{\text{max}}^G(G)}{\beta}$$

with conversion coefficients \(\alpha\) and \(\beta\), mortality \(m\), maximal quota-dependent growth \(\mu_{\text{max}}^A\) and minimum quota \(Q_{\text{max}}\). The variable growth rate \(\mu(Q,G)\) implements Liebig’s law of the minimum, selecting either growth rate \(\mu_A(Q)\), which depends on the nitrogen cell quota \(Q\), or glucose-dependent growth rate \(\mu_G(G)\) if smaller.

Model parameters (Table S2, Supporting Information) were determined by minimizing the sum of relative mean deviations between experimental data and model simulations (i.e. OD and \(P\), NH₄⁺ and \(A\), glucose and \(G\), N:P ratio and \(Q\)) using the meta heuristics differential evolution (Storn and Price 1997) and adaptive simulated annealing (Ingber 1989) implemented by Oldenhuys and Vandekerckhove (2009).

**Genomic analysis**

The genome sequence of \(P\). inhibens DSM 17395 (Thole et al. 2012) served as backbone for integrated metabolic reconstruction based on proteomic and metabolomic data. FunPhaeobacter-ctional assignments for potential transport proteins were based on the Transporter Classification Database (TCDB) (Saier et al. 2016). Metabolic pathways were assessed with BRENDA (Chang et al. 2015) and MetaCyc (Caspi et al. 2016).

**RESULTS AND DISCUSSION**

**Phaeobacter inhibens DSM 17395 rapidly depletes external ammonium and increases the cellular N:C ratio**

In growth experiments with marine mineral medium, \(P\). inhibens DSM 17395 rapidly and completely depleted the nitrogen source supplied (0.8 mM NH₄⁺) during early (exponential) growth, i.e. when only low (~0.22) of maximal biomass concentrations were formed (Fig. 1a). This marked uncoupling of NH₄⁺ consumption from bulk (linear) growth suggests that \(P\). inhibens DSM 17395 accumulates reduced nitrogen during early growth (~20 h).

The buildup of intracellular nitrogen reservoirs is evident from cellular N:C ratios (Fig. 1b), increasing from ~0.09 (inoculum) to maximal values of ~0.22 within less than 10 h (~10% NH₄⁺ depleted) and then remained constant for 5–6 h during bulk NH₄⁺ depletion. Subsequently, N:C ratios decreased continuously until a constant minimal level of ~0.06 was reached at the transition into the stationary growth phase (~50 h after inoculation), marking the onset of nitrogen starvation. Varying the concentration of initially supplied NH₄⁺, i.e. more limiting (0.4 mM) or excess (1.8 mM), revealed very similar N:C ratio profiles (Fig. S1, Supporting Information), thus indicating that molecular crowding and solvation/diffusion capacities could constrain elemental homeostasis and therefore the upper limit (N:C ratios of 0.22–0.24). The low N:C ratios (0.06) of \(P\). inhibens DSM 17395 during the stationary growth phase agree with growth limitation by nitrogen and storage of carbon as PHB, the formation of which started upon depletion of external NH₄⁺ (Fig. 1). The nutrient quota (Droop 1968) modeled for nitrogen complies with profiles of cellular N:C ratios in \(P\). inhibens DSM 17395 (Fig. 1b; Fig. S2, Supporting Information). The initially high quota reflects the storage of nitrogen during NH₄⁺ depletion, followed by its decrease due to allocation of internal nitrogen pools to descending cells during cell division.

The marked uncoupling of NH₄⁺ consumption from bulk growth by \(P\). inhibens DSM 17395 is noteworthy, as previous reports showed \(K\). pneumoniae to tightly couple NH₄⁺ consumption with growth under similar cultivation condition, i.e. growth ceased upon NH₄⁺ depletion from the medium (Wanner and Egli 1990). Subsequent slow and slight increases in biomass observed for \(K\). pneumoniae and other bacteria (e.g. \(E\). coli, Aerobacter aerogenes) result from conversion of excess carbon source into storage polymers, rather than from growth per se (Duguid and Wilkinson 1953; Wanner and Egli 1990; Guevara-Martinez et al. 2015). In contrast, it is long known that utilization of phosphate, another macronutrient, can be decoupled from growth in \(K\). pneumoniae and \(E\). coli, with elevated levels of DNA and RNA serving as phosphate storage to sustain successive growth (Horiuchi 1959; Wanner and Egli 1990).

**Rapid nitrogen storage prevalent in other Phaeobacter spp. and Ruegeria pomeroyi**

Recent metagenomic studies revealed that Phaeobacter and Ruegeria spp. constitute a major fraction of the rosebacterias associating with large-scale blooms of the widespread marine coccolithophore Emiliana huxleyi (Segev et al. 2016). Further tested strains of \(P\). inhibens, \(P\). gallaeciensis and \(R\). pomeroyi exploited external NH₄⁺ essentially as observed for \(P\). inhibens DSM 17395 (Fig. 1), i.e. they uncoupled NH₄⁺ depletion from bulk growth and at the same time increased the cellular N:C ratio (Fig. 2; Fig. S3, Supporting Information). Potential eco-
Growth and substrate consumption profiles with 0.8 mM ammonium provided as sole nitrogen source in glucose-containing seawater medium. The provision with different concentrations of NH4+ (Fig. S1, Supporting Information) indicated that P. inhibens DSM 17395 rapidly assimilated the imported NH4+ via the canonical pathways (glutamate dehydrogenase GltBD and several aminotransferases). Nitrogen assimilation resulted in the buildup of manifold nitrogen-containing small metabolites (Fig. 3), with most of them serving as building blocks for the biosynthesis of cellular macromolecules (Fig. 1).

Amino acids and their derivatives represented the largest group of intracellular metabolites formed during depletion of external NH4+ (Fig. 3). Relative abundances of individual amino acids in P. inhibens DSM 17395 (Table S4, Supporting Information) reflected their distribution known to universally occur in living cells. In accord, particularly high abundance increases (up to ∼20-fold) were detected for branched-chain amino acids (Leu, Val, Ile), which was paralleled by the presence of the corresponding enzymes for their biosynthesis (mostly) also at elevated levels (Fig. S4, Supporting Information). Additional strongly elevated N-containing metabolites comprised three nucleotides (C, T, U), six carboxylates (e.g. 4-aminobutanoate), the biogenic amine putrescine, the amino alcohol diethanolamine and the dipeptide glycyl-glycine (Fig. 3). Among cellular lipids, N-containing glycerol phospholipids (phosphatidylcholine, phosphatidylethanolamine) contributed only a minor fraction (Fig. 3), with most of them serving as building blocks for the biosynthesis of cellular macromolecules (Fig. 1).

Chemical identity of intracellular nitrogen species buildup by P. inhibens DSM 17395 during early growth

Uptake of NH4+ by P. inhibens DSM 17395 during early growth is apparently facilitated by the AmtB family transporter PGA1_c29410 (identified; Table S3, Supporting Information), regulated by its cognate PII family protein GlnK (identified) (Thomas, Coutts and Merrick 2000; Conroy et al. 2007). The proteomic and metabolomic datasets (Tables S3 and S4, Supporting Information) indicated that P. inhibens DSM 17395 rapidly assimilated the imported NH4+ via the canonical pathways (glutamate dehydrogenase GluD, glutamine-forming glutamate synthase GltBD and several aminotransferases). Nitrogen assimilation resulted in the buildup of manifold nitrogen-containing small metabolites (Fig. 3), with most of them serving as building blocks for the biosynthesis of cellular macromolecules (Fig. 1).

The determined cellular N:C ratios are within the wide range of values (0.07 to 0.35) observed for marine bacteria (Fagerbakke, Heldal and Norland 1996), reflecting the heterogeneous nitrogen availability in the oceans, as well as species- and growth phase-specific differences in nitrogen requirements. The capacity to store nitrogen internally during transient pulses of NH4+ or nitrate is a typical feature of marine phytoplankton (Dorcht et al. 1984; Admiraal, Peletier and Laane 1986) and was recently also observed in the cyanobacterium Planktothrix agardhii (Van de Waal et al. 2010), but has to our knowledge not been reported for heterotrophic bacterioplankton members.
Figure 2. Rapid nitrogen storage prevalent in other Phaeobacter spp. and Ruegeria pomeroyi. Uncoupling of bulk growth from NH$_4^+$ depletion was detected also in other roseobacters that were cultivable in the same mineral medium as used for P. inhibens DSM 17395 (see Fig. 1). All strains had already depleted NH$_4^+$ completely when only 16.1%–47.7% of the maximal optical densities (OD$_{max}$) were attained. During NH$_4^+$ depletion (at $\sim$50% consumed), all strains buildup intracellular nitrogen stocks, as evidenced by the 2.7–4.5-fold increase in cellular N:C ratios (in comparison to cells at OD$_{max}$). The non-Roseobacter strains R. blandensis and E. coli conducted differently with respect to both, ammonium depletion and increase of cellular N:C ratio. Displayed data (average ± average deviation) are based on 3–4 biological replicates. Strains were originally isolated from distant geographical provenances and disparate habitats. Detailed growth curves including substrate consumption profiles are provided in Fig. S3, Supporting Information.

S3, Supporting Information) may be used in an integrated yet presently unknown manner. The genes encoding these two nitrogen regulatory systems also co-occur in other roseobacters (Smith et al. 2013). In contrast, heterotrophic bacteria such as E. coli employ only the NtrBC/PII system (van Heeswijk, Westerhoff and Boogerd 2013), whereas the oligotrophic marine bacterium ‘Candidatus Pelagibacter ubique’ relies exclusively on NtrXY (Smith et al. 2013).

With the depletion of external NH$_4^+$, levels of most N-containing metabolites rapidly decreased in P. inhibens DSM 17395 (Fig. 3), until they were either strongly reduced in abundance or could not be detected during the stationary growth phase (at ≥50 h). Pronounced changes in cellular macromolecules were observed for DNA, RNA and proteins (Fig. 1), reflecting the elevated levels of the corresponding metabolite precursors (amino acids, nucleotides), as well as exponential growth (during NH$_4^+$ depletion). The high demand for anabolic enzymes during NH$_4^+$ assimilation (Fig. 4; Fig. S4 and Table S3, Supporting Information) may further contribute to the abundance increase of total proteins. The decrease in DNA, RNA and protein levels upon depletion of external NH$_4^+$ paralleled the decreasing cellular N:C ratios (Fig. 1), and suggests degradation of these polymers to recycle polymer-bound nitrogen.

The copiously produced N-containing metabolites should distribute evenly in nitrogen-rich cells of P. inhibens DSM 17395 simply by diffusion, allowing for uniform propagation of cellular nitrogen from mother to daughter cells during cell division. Moreover, they serve as ‘prebuilt’ monomers for instantaneously polymer biosynthesis, especially during bulk growth after depletion of external NH$_4^+$. The broad distribution of assimilated nitrogen into diverse cellular monomers and polymers reveals an intuitive nitrogen storage strategy in P. inhibens DSM 17395. Most cyanobacteria as well as some heterotrophic bacteria respond to surplus nitrogen supply by specifically boosting the biosynthesis of arginine, as major building block for the nitrogen-rich storage polymer cyanophycin (multi-L-arginyl-poly-L-aspartate) generated by cyanophycin synthetase (Frommeyer, Wiefel and Steinbüchel 2016). Lacking genes for the latter, P. inhibens DSM 17395 cannot synthesize cyanophycin. E. coli responds to NH$_4^+$ pulses (10 mM) by increasing the levels of several amino acids for a time span that was by far too short ($\ll$ doubling time) to serve growth-relevant storage functions (Yuan et al. 2009). In contrast, the observed strategy of P. inhibens DSM 17395 to build up surplus internal nitrogen stocks that fuel bulk growth more closely resembles that of eukaryotes. Marine phytoplankton internally stores nitrogen in the form of DNA, RNA, proteins or free amino acids, but also as NH$_4^+$ or nitrate in vacuoles (Dortch et al. 1984; Admiraal, Peletier and Laane 1986).

High cellular nitrogen levels coincide with production of potential exotoxins and antibiotic TDA in P. inhibens DSM 17395

Phytoplankton-associated P. inhibens DSM 17395 is suggested to rely on secreted organic compounds (e.g. exoproteins, secondary metabolites) as important assets for its host-associated lifestyle (e.g. Porsby, Nielsen and Gram 2008; Seyedsayamdost et al. 2011). The exoproteomes of diverse roseobacters (Christie-Oleza et al. 2012), including P. inhibens DSM 17395 (Koßmehl et al. 2013), are dominated by potential RTX (Repeats-in-ToXin) proteins exported via type I secretion systems, also at hand in P. inhibens DSM 17395 (Koßmehl et al. 2013). These toxins are
active against prokaryotic and eukaryotic cells, and have been suggested, amongst others, to form ion-permeable channels, thus dissipating ion gradients and the membrane potential (Linhartová et al. 2010). However, the actual mode of action of RTX-like proteins in roseobacters is currently unknown. In addition, P. inhibens DSM 17395 and related species produce the secondary metabolite TDA (e.g. Berger et al. 2011), which exhibits broad-spectrum antibiotic activity (Wilson et al. 2016). Here, we show that formation and maximal abundance of several potential RTX proteins and of TDA co-occur with high cellular N:C ratios buildup during depletion of external NH$_4^+$ (Fig. 5).

The most abundant of the eight identified potential RTX proteins (PGA1_65p00350) accounted for ~50%–60% of the total SDS PAGE-resolvable exoproteome during NH$_4^+$ consumption and was estimated to reach maximal concentrations of ~600 μg protein L$^{-1}$ in the prepared exoproteome upon complete NH$_4^+$ depletion (Fig. S6, Supporting Information). The peak in production of this protein (~7 μg RTX protein per mg CDW) (Fig. 5a) was observed at highest cellular N:C ratios, which occurred before NH$_4^+$ depletion (Fig. 1b). We estimate that biosynthesis of this single RTX-like protein (molar N:C ratio of 0.32) at the observed concentrations should consume up to ~1.0% (7–8 μM) of the initially supplied NH$_4^+$. With decreasing N:C ratios, the level of PGA1_65p00350 decreased continuously and reached almost zero in late stationary growth phase. The concentration profile of PGA1_65p00350 deviated strongly from that of the total exoproteins after NH$_4^+$ depletion (Fig. 5a), agreeing with the altered composition of the SDS PAGE-resolved total exoproteome (Fig. S6, Supporting Information). The decline in total exoprotein abundance at ~OD$_{max}$ coincided with the appearance of four secreted proteases/peptidases, possibly involved in external proteolysis of the excreted potential RTX and other proteins formed beforehand (Fig. S6, Supporting Information).

Several identified amino acid- and peptide-specific transporters (Table S3, Supporting Information) potentially scavenge the proteolytic products for intracellular recycling of organic nitrogen during nitrogen starvation.

TDA levels in culture supernatants were measured indirectly by photometric detection of the diagnostic brown-colored TDA-iron complexes (Prol García et al. 2014; D’Alvise et al. 2015). As observed for PGA1_65p00350, maximal production of TDA was recorded at highest cellular N:C ratios and decreased after NH$_4^+$ was completely depleted (Fig. 5b). The TDA molecule itself does not contain nitrogen, but its biosynthesis drains N-containing intracellular metabolites (Phe, Cys) and requires specific enzymes (Fig. 4). This is evident from the higher abundance of biosynthetically relevant key metabolites (e.g. O-acetylserine) and enzymes (e.g. Tda proteins) during maximal TDA production (Figs. 4 and 5b). These results suggest that the metabolic status (levels of cellular nitrogen/amino acids) triggers the onset and duration of TDA formation irrespective of the initially supplied NH$_4^+$ concentration (0.4, 0.8 or 1.8 mM) (Fig. 1b; Figs. S1 and S7, Supporting Information). This link was not recognized previously owing to P. inhibens DSM 17395 being routinely cultured with nitrogen-rich complex media, which supported continuous TDA production during active growth (e.g. Berger et al. 2011; Trautwein et al. 2016). Moreover, integration of metabolic and quorum sensing signals (e.g. Berger et al. 2011; Prol García,
Figure 4. Biosynthesis of antibiotic TDA during ammonium depletion in *P. inhibens* DSM 17395 (see Fig. 5b). TDA biosynthesis is closely intertwined with the biosynthesis of amino acids (Phe, Ser, Cys) and with assimilatory sulfate reduction, thus requiring an integrated cross-regulation of carbon, nitrogen and sulfur metabolism in *P. inhibens* DSM 17395. TDA biosynthesis drains phenylpyruvate from Phe anabolism, proceeding via the conventional shikimate-chorismate pathway. Phenylpyruvate-derived phenylacetyl-CoA is then transformed via epoxidation and isomerization (oxepin-CoA forming), followed by ring cleavage to 3-oxo-5,6-dehydrosuberoyl-CoA (Teufel et al. 2011). The latter has been proposed to be converted to the seven-membered tropone carbocycle, ultimately receiving reduced sulfur from cysteine-derived S-thiocysteine (Brock, Nikolay and Dickschat 2014). Boxes indicate the timepoints (15, 20, 25, and 40 h; Fig. 1) for cell harvesting and their different colors reflect (i) relative changes in abundance of identified metabolites and soluble proteins (D; determined by 2D DIGE), as well as (ii) identification metascores for soluble (S; determined by shotgun analysis) and membrane (M) proteins. The complete dataset is presented in Fig. S8, Supporting Information. Compound abbreviations in alphabetic order: 2-OG, 2-oxoglutarate; PEP, phosphoenolpyruvate; 3-PG, 3-phosphoglycerate; TDA, tropodithietic acid. Enzyme names in alphabetic order (identified: blue font color; predicted: grey font color): AroA, 3-phosphoshikimate-1-carboxyvinyltransferase; AroB, 3-hydroquinate synthase; AroC, chorismate synthase; AroE, shikimate dehydrogenase; AroH, phospho-2-dehydro-3-deoxyheptanoate aldolase; CysE, serine acetyltransferase; CysH, phosphoadenosine phosphosulfate reductase; CysK, cysteine synthase; IorAB, indolepyruvate oxidoreductase; PaaACG, proteins involved in phenylacetate degradation; PaaZ2, oxepin-CoA hydrolase/3-oxo-5,6-dehydrosuberoyl-CoA semialdehyde dehydrogenase (NADP⁺); PatB, cystathionine beta-lyase; PGA1c00590, chorismate mutase; PGA1c07640, O-acetylserine sulfhydrylase; PGA1c20760, putative nitrite/sulfite reductase; PGA1c24800, putative bifunctional SAT/APS kinase; PGA1c30200, prephenate dehydrogenase; SerA, 3-phosphoglycerate dehydrogenase; SerB, phosphoserine phosphatase; SerC, phosphoserine aminotransferase; TdaA, transcriptional regulator, LysR family; TdaB, putative beta etherase; TdaC, conserved hypothetical protein, similar to prephenate dehydratase; TdaD, thiosterase superfamily protein; TdaE, acyl-CoA dehydrogenase; TdaF, putative flavoprotein, HFCD family, TyrB, aromatic amino acid aminotransferase.
the abundant potential RTX protein PGA1 during growth with 0.8 mM ammonium (see Fig. 1). Production of TDA was determined indirectly by its absorbance at 398 nm (Abs398). Dis-}

otic secondary metabolite TDA, draining amino acids during its biosynthesis (see B)

exoproteins (for details see Fig. S6, Supporting Information) and of (A) the antibi-}

otic secondary metabolite TDA, draining amino acids during its biosynthesis (see Fig. 4). TDA was determined indirectly by its absorbance at 398 nm (Abs398). Dis-}

played data (average ± average deviation) are based on 3–4 biological replicates.

Figure 5. Nitrogen-dependent secretion of exoproteins and TDA by P. inhibens DSM 17395 during growth with 0.8 mM ammonium (see Fig. 1). Production of (A) the abundant potential RTX protein PGA1,05p00350 in comparison to the total exoproteins (for details see Fig. S6, Supporting Information) and of (B) the antibi-

otic secondary metabolite TDA, draining amino acids during its biosynthesis (see Fig. 4). TDA was determined indirectly by its absorbance at 398 nm (Abs398). Dis-

played data (average ± average deviation) are based on 3–4 biological replicates.

D’Alvise and Gram 2013b) could contribute to the higher maximal levels of TDA during growth with excess of nitrogen (Fig. S7, Supporting Information). High cellular N:C ratios apparently also positively influence toxin production in marine phytoplankton (Camargo and Alonso 2006).

Ecological implications

Heterotrophic bacteria are essential drivers of the nitrogen cycle in the oceans, amongst others by regenerating NH$_4^+$ during the mineralization of biomass and/or competing with phytoplankton for its utilization. The efficient sequestering of NH$_4^+$ appears to be useful for P. inhibens DSM 17395 and related strains (Fig. 2) in several ways: (i) internal (‘private’) reservoirs exclusively ben-

efit their descendants, (ii) competing bacterioplankton (including ammonia-oxidizing archaea and bacteria (Martens-Habbena et al. 2009)) and phytoplankton species could be deprived of this nutrient, and (iii) it could prevent competitors from poten-

tially growing faster. The capacity to rapidly store nitrogen internally possibly reflects the strong influence of NH$_4^+$ availability on the growth of P. inhibens DSM 17395 (Trautwein et al. 2017) and in more general terms an adaptation to fluctuating NH$_4^+$ supply. The latter is expected to occur frequently in the natural habitat of the studied roseobacters: on a larger scale in coastal regions and on a microcosm-scale on marine snow particles representing nutrient-rich hotspots in otherwise oligotrophic water columns (Azam and Long 2001; Gram et al. 2002). To extrapolate these new findings for P. inhibens DSM 17395 and its close relatives from this defined study to the natural environment is, of course, speculative at present, since in situ concentrations of NH$_4^+$ vary considerably, and nutrient supply (incl. various nitrogen sources from DIN and DON) as well as community composition are much more complex.

Investing the costly acquired nitrogen at the right time into massive secretion of potential RTX proteins, as well as into the complex biosynthesis of TDA (Figs. 4 and 5), may be part of a defense mechanism in P. inhibens DSM 17395. This strategy would protect the nitrogen-rich cells of P. inhibens DSM 17395 from predation and aggressive displacement from colonized nutritional hotspots (e.g. algae, rotifers, fish larvae) by compet-

ing species. Restricting maximal production of potential RTX proteins and TDA to nitrogen-rich cells implies that eukaryotic interaction partners could benefit even longer from protection against pathogens (e.g. Vibrio sp.), if they continuously provide P. inhibens DSM 17395 with sufficient amounts of reduced nitrogen. N$_2$-fixing phytoplankton constantly release N-containing metabolites (mainly free amino acids) (Mague et al. 1980; Sarmen
to et al. 2013), which increases even more under inorganic nutrient limitation when internal carbon storage capacities are exceeded (Obernosterer and Herndl 1995; Wilhelm et al. 2006). During growth of P. inhibens DSM 17395 with an excessive supply of nitrogen provided by a complex medium rich in amino acids and resembling the composition of phytoplankton exu-

dates, the organism simultaneously utilized most of the amino acids, sustained a high cellular N:C ratio and continuously pro-

duced potential RTX proteins and TDA until OD$_{max}$ (Zech et al. 2013b; Trautwein et al. 2016). Reduced (in)organic nitrogen could thus play a key role in fostering the proposed mutually beneficial interactions between P. inhibens DSM 17395 and phytoplankton (Seyedsayamdost et al. 2011).

CONCLUSIONS

Phaeobacter inhibens DSM 17395 exploits external NH$_4^+$ in a com-

plex manner throughout the different stages of growth that starts with (i) the transient buildup of diverse intracellular nitrogen reservoirs to facilitate rapid depletion and assimilation of external NH$_4^+$ during early growth (Fig. 6), followed by (ii) prop-

agation and redistribution of intracellular nitrogen during bulk growth accompanied by carbon (PHB) storage (Fig. 1), and finally (iii) breakdown and recycling of DNA, RNA and proteins (inter-

nal and secreted) during nitrogen starvation in the station-

ary growth phase (Supplementary Results and Discussion). The allocation of assimilated nitrogen into monomeric and poly-

meric cellular constituents for transient nitrogen storage (Fig. 6) appears intuitive, but should come with high energetic costs for boosted global production of the many biosynthetic enzymes required, and for biosynthesis itself. This contrasts the metabolically much simpler way of nitrogen storage in bacteria produc-

ing a homogeneous N-reserve polymer such as cyanophycin. Instead, the nitrogen storage strategy of P. inhibens DSM 17395 resembles that of eukaryotic phytoplankton and may reveal another facet of an apparently close relationship between a heterotrophic bacterioplankton member and its eukaryotic inter-

action partners. Nitrogen-dependent secretion of potential RTX proteins and TDA (Fig. 5) could enable P. inhibens DSM 17395 to bequeath the secured nitrogen successfully to subsequent generations. Considering that biosynthesis of the antimicrobial sec-

ondary metabolite TDA occurs at the interface between amino acid biosynthesis and sulfur assimilation (Fig. 4), an integrated cross-regulation of C-, N- and S-metabolism in P. inhibens DSM 17395 seems likely. The rapid storage of nitrogen described for Phaeobacter spp. and R. pomeroyi may add a new facet to the
Figure 6. Schematic model of transitory nitrogen storage in *P. inhibens* DSM 17395. External ammonium, possibly derived from different environmental sources, is rapidly assimilated (via glutamate and glutamine) and utilized for biosynthesis of diverse N-containing metabolites and cellular macromolecules. Investing parts of the assimilated nitrogen in biosynthesis and secretion of potential RTX proteins and antibiotic TDA may ensure competitive nutrient acquisition and protection of nitrogen-rich cells from predation.

marine nitrogen cycle, as heterotrophic bacteria, like phytoplankton, may pursue a ‘luxury goods’ strategy, resulting in a transient intracellular accumulation of reduced nitrogen and conversely its removal from the readily accessible communal seawater pool.

SUPPLEMENTAL DATA
Supplementary data are available at FEMSEC online.

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