Mechanisms of Trichodesmium demise within the New 1 Caledonian during the lagoon VAHINE mesocosm 2 experiment 3 4 D. Spungin¹, U. Pfreundt², H. Berthelot³, S. Bonnet^{3,4}, D. AlRoumi⁵, F. Natale⁵, 5 W.R. Hess², K.D. Bidle⁵, I. Berman-Frank¹ 6 7 [1] {The Mina and Everard Goodman Faculty of Life Sciences, Bar-Ilan University, Ramat-8 9 Gan, Israel} [2] {University of Freiburg, Faculty of Biology, Schänzlestr. 1, D-79104 Freiburg, Germany} 10 [3] {Aix Marseille Université, CNRS/INSU, Université de Toulon, IRD, Mediterranean 11 Institute of Oceanography (MIO) UM 110, 13288, Marseille, France} 12 [4] {Institut de Recherche pour le Développement (IRD), AMU/CNRS/INSU, Université de 13 Toulon, Mediterranean Institute of Oceanography (MIO) UM 110, 13288, Marseille-Noumea, 14 France-New Caledonia} 15 16 [5] {Department of Marine and Coastal Sciences, Rutgers University, New Brunswick, NJ, USA} 17 18 Correspondence to: I. Berman-Frank (ilana.berman-frank@biu.ac.il) 19 20

- 21
- 22
- 23

24 Abstract

The globally important marine diazotrophic cyanobacterium Trichodesmium is seasonally 25 26 abundant in the New Caledonian lagoon (in the Southwestern Pacific ocean) during austral spring/summer. We investigated the cellular processes mediating Trichodesmium mortality 27 from large surface accumulations (blooms) of Trichodesmium in the lagoon. Trichodesmium 28 cells (and associated microbiota) were collected at the time of surface accumulation, enclosed 29 under simulated ambient conditions, and sampled over time to elucidate the stressors and 30 subcellular underpinning of rapid (~ 24 h) biomass demise (> 90 % biomass) and 31 32 disappearance occurring in these conditions. Metatranscriptomic profiling of Trichodesmium biomass 8 h and 22 h after bottle incubation of surface accumulations demonstrated 33 34 upregulation expression of genes required to increase phosphorus (P) and iron (Fe) availability and transport. In contrast, genes responsible for nutrient storage were downregulated. Total 35 36 viral abundance, oscillated throughout the experiment and showed no significant relationship with the development or demise of the Trichodesmium biomass. Enhanced caspase-specific 37 38 activity and upregulated expression of a suite of metacaspase genes as the Trichodesmium biomass crashed implicated autocatalytic programmed cell death (PCD) as the mechanistic 39 40 cause. Concurrently, genes associated with buoyancy and gas-vesicle production were 41 strongly downregulated concomitant with increased production and high concentrations of transparent exopolymeric particles (TEP). The rapid, PCD-mediated, decline of the 42 Trichodesmium biomass, as we observed from our incubations, parallels mortality rates 43 reported from Trichodesmium blooms in situ. Our results suggest that, whatever the ultimate 44 factor, PCD-mediated death in Trichodesmium can rapidly terminate blooms, facilitate 45 aggregation, and expedite vertical flux to depth. 46

- 47
- 48
- 49
- 50
- 51
- 52

55 **1 Introduction**

The filamentous N₂-fixing (diazotrophic) cyanobacteria Trichodesmium spp. are important 56 contributors to marine N2 fixation as they form massive blooms (surface accumulations with 57 high biomass density) throughout the oligotrophic marine sub-tropical and tropical oceans 58 (Capone et al., 2004; Capone and Carpenter, 1982; Capone et al., 1997). These surface blooms 59 with densities of 3000 to > 10,000 trichomes L⁻¹ and chlorophyll a (Chl a) concentrations 60 ranging from 1-5 mg L^{-1} develop swiftly and are characterized by high rates of CO₂ and N₂ 61 fixation (Capone et al., 1998; Luo et al., 2012; Rodier and Le Borgne, 2008; Rodier and Le 62 63 Borgne, 2010). Trichodesmium blooms also occur frequently during austral summer between November and March over large areas of the New Caledonian lagoon in the Southwest Pacific 64 65 Ocean (Dandonneau and Gohin, 1984; Dupouy et al., 2011).

Trichodesmium has been extensively investigated [reviewed in Capone et al. (1997); and 66 Bergman et al. (2012)]. Yet, relatively few publications have examined the mortality and fate 67 68 of these blooms that often collapse abruptly with mortality rates paralleling growth rates and 69 biomass declines > 50 % occurring within 24 h from peak abundance (Bergman et al., 2012; Rodier and Le Borgne, 2008; Rodier and Le Borgne, 2010). Cell mortality can occur due to 70 grazing of Trichodesmium by pelagic harpacticoid copepods (O'Neil, 1998) or by viral lysis 71 (Hewson et al., 2004; Ohki, 1999). Both iron (Fe) and phosphorus (P) availability regulate N₂ 72 fixation and production of Trichodesmium populations, causing a variety of stress responses 73 when these nutrients are limited (Berman-Frank et al., 2001). Fe depletion as well as oxidative 74 75 stress can also induce in Trichodesmium a genetically controlled programmed cell death 76 (PCD) that occurs in both laboratory cultures and in natural populations (Bar-Zeev et al., 2013; Berman-Frank et al., 2004; Berman-Frank et al., 2007). Mortality of Trichodesmium via 77 PCD is morphologically and physiologically distinct from necrotic death and triggers rapid 78 79 sinking of biomass that could enhance carbon export in oligotrophic environments (Bar-Zeev 80 et al., 2013). Sinking is due to concomitant internal cellular degradation, vacuole loss, and the increased production of extracellular polysaccharide aggregates, operationally defined as 81 transparent exopolymeric particles (TEP) (Bar-Zeev et al., 2013; Berman-Frank et al., 2004; 82 Berman-Frank et al., 2007). 83

The VAHINE project investigated the fate of newly fixed N by diazotrophs and aimed to test changes in organic matter export, following diazotroph development and mortality. For this, large (50 m³) mesocosms were deployed in the in the New Caledonian lagoon and followed

87 over the course of 23 days (Bonnet et al., 2016a). Our objective during the VAHINE project was to study the involvement of PCD in the fate of natural Trichodesmium blooms induced in 88 these mesocosms. While *Trichodesmium* was initially present, and conditions in the 89 mesocosms appeared favorable, no Trichodesmium blooms developed within the mesocosms, 90 91 yet UCYN-C did increase, allowing to meet the scientific objectives of the project (Berthelot et al., 2015; Bonnet et al., 2016a; Turk-Kubo et al., 2015). However, Trichodesmium 92 93 developed at different phases of the experimental period outside the mesocosms (Turk-Kubo et al., 2015). Here, we investigated mortality processes in a short-lived Trichodesmium bloom 94 95 that developed and crashed in the lagoon waters at the end of the VAHINE experiment. Using a series of microcosm incubations with collected Trichodesmium biomass, we elucidated the 96 stressors and subcellular underpinning of rapid (~ 24 h) biomass demise and disappearance. 97 Here we present, for the first time, physiological, biochemical, and metatranscriptomic 98 99 evidence for nutrient-stress induced PCD in natural populations that lead to Trichodesmium mortality including concomitant downregulation of gas vesicle synthesis and enhanced TEP 100 production. Such mechanisms would lead to enhanced export flux in natural blooms that also 101 crash within 1-2 days. 102

103

104 2 Methods

2.1. Sampling site and sampling conditions during pre-bloom periods

Our study was performed during the VAHINE mesocosm project set 28 km off the coast of 106 New Caledonia from 13 January 2013 (day 1) to 6 February 2013 in the New Caledonian 107 oligotrophic lagoon (22°29.10' S, 166° 26.90' E). The 25 m deep sandy-bottom lagoon is 108 109 generally protected from the dominant trade winds yet the waters of the lagoon are influenced by the oligotrophic oceanic waters coming into the lagoon via the Boulari Pass (Bonnet et al., 110 2016a). Detailed descriptions of the site selection and sampling strategy are provided 111 elsewhere (Bonnet et al., 2016a). The lagoon water outside the mesocosms was sampled daily 112 during the experiment and serve as 'pre-bloom' data. Large volume samples (50 L) were 113 collected from 1, 6, and 12 m depths at 07:00 using a Teflon® PFA pump and PVC tubing. 114 Samples were immediately transferred back to laboratories aboard the R/V Alis and 115 subsampled for a suite of parameters [as described below and in Bonnet et al. (2016a)]. On 116 day 23 at 12:00 h, we observed a large surface accumulation of *Trichodesmium* in the lagoon 117 close to the enclosed mesocosms. This biomass accumulation (hereafter called - "bloom") 118

served as the source for experiments 1 and 2 to examine the fate of *Trichodesmium* (section2.2, Fig. S1).

121

122 2.2. Short-term incubations to assess bloom decline

123 **Experiment 1** – *Trichodesmium* filaments and colonies were collected from the dense surface bloom (day 23, 12:00 h; designated T₀. Fig. 2a-c) using a plankton net (mesh size, 80 µm) 124 125 towed through different patches of the bloom from the surface water. The total contents of the net were combined and resuspended in filtered seawater (FSW) (0.2 µm pore size), split 126 between six identical 4.5 L Nalgene polycarbonate bottles (Fig. 2d-e), and incubated as 127 detailed below. Based on previous experience (Berman-Frank et al., 2004), resuspension of 128 *Trichodesmium* cells in the extremely high densities of the surface blooms (> 1 mg L^{-1} Chl *a*; 129 Fig. 2a-c) would cause an almost immediate crash of the biomass. Consequently, we 130 resuspended the collected biomass in FSW at ~ 1000 fold lower cell densities (150 μ g L⁻¹) that 131 resemble the cellular abundance at the edges of the slicks (Fig. 2). Experiment 2 – Seawater 132 133 from the surface bloom was collected 5 h after the initial surface bloom was sighted (day 23, 17:00) by using a Teflon® PFA pump and PVC tubing directly filling nine 20 L polyethylene 134 carboys gently to avoid destroying biomass. Bottles from experiments 1 and 2 were placed in 135 136 on-deck incubators, filled with running seawater to maintain ambient surface temperature (~ 26 °C), and covered with neutral screening at 50 % surface irradiance levels. Water from 137 138 experiment 1 was sampled every 2-4 h until the biomass collapsed (after ~ 22 h) for: Chl a concentration, caspase activity, 16S rRNA gene sequencing, and metatranscriptomics. Water 139 from experiment 2 was sampled for PON, POC, NH₄⁺, N₂ fixation rates, TEP production, and 140 virus abundance (days 23-25) (Fig. S1). Prior to incubations, all incubation bottles and 141 carboys were washed with 10 % HCl overnight and rinsed 3 times with ambient seawater. 142

143

2.3. Chlorophyll a concentrations

Samples for the determination of Chl *a* concentrations during pre-bloom days were collected
by filtering 550 mL of seawater on GF/F filters. Filters were directly stored in liquid nitrogen.
Chl *a* was extracted in methanol and measured fluorometrically (Herbland et al., 1985).
During short-term experiment 1, samples for Chl *a* were collected by filtering 200 mL on
GF/F filters (Whatman, Kent, UK). Chl *a* was extracted in methanol and measured

spectrophotometrically (664 and 750 nm; CARY100, Varian, Santa Clara, CA, USA)according to Tandeau de Marsac and Houmard (1988).

151

152

153 **2.4. Particulate organic carbon (POC) and nitrogen (PON)**

154 Detailed POC and PON analyses are described in Berthelot et al. (2015). POC samples were 155 collected by filtering 2.3 L of seawater through pre-combusted (450 °C, 4 h) GF/F filter and 156 determined using the combustion method (Strickland and Parsons, 1972) on an EA 2400 CHN 157 analyzer. Samples for PON concentrations were collected by filtering 1.2 L of water on pre-158 combusted (450 °C, 4 h) and acid washed (HCl, 10 %) GF/F filters and analyzed according to 159 the wet oxidation protocol described in Pujo-Pay and Raimbault (1994) with a precision of 160 $0.06 \mu mol L^{-1}$.

161 **2.5.** N₂ fixation rates and NH₄⁺ concentrations

N₂-fixation rate measurements used in experiment 2 are described in detail in (Berthelot et al., 162 2015). Samples were collected at 17:00 in 4.5 L polycarbonate bottles and amended with ¹⁵N₂-163 enriched seawater, within an hour of biomass collection, according to the protocol developed 164 by Mohr et al. (2010) and Rahav et al. (2013). Briefly, seawater was degassed through a 165 degassing membrane (Membrana, Minimodule®, flow rate fixed at 450 mL min⁻¹) connected 166 to a vacuum pump. Degassed seawater was amended with 1 mL of ${}^{15}N_2$ (98.9 % atom ${}^{15}N$, 167 Cambridge Isotopes) per 100 mL. The bottle was shaken vigorously and incubated overnight 168 at 3 bars to promote ¹⁵N₂ dissolution. Incubation bottles were amended with 1:20 (vol:vol) of 169 ¹⁵N₂-enriched seawater, closed without headspace with silicone septum caps, and incubated 170 171 for 24 h under in situ-simulated conditions in on-deck incubators (described above). 2.2 L from each experimental bottle was filtered under low vacuum pressure (< 100 mm Hg) onto a 172 pre-combusted (450 °C, 4 h) GF/F filter (25 mm diameter, 0.7 µm nominal porosity). The 173 filters were stored at -20 °C and dried for 24 h at 60 °C before mass spectrometric analysis. 174 PON content and PON ¹⁵N enrichments were determined using a Delta plus Thermo Fisher 175 Scientific isotope ratio mass spectrometer (Bremen, Germany) coupled with an elemental 176 177 analyzer (Flash EA, Thermo Fisher Scientific). N₂-fixation rates were calculated according to the equations detailed in Montoya et al. (1996). We assumed significant rates when the ¹⁵N 178 enrichment of the PON was higher than three times the standard deviation obtained from T_0 179

180 samples. The ¹⁵N batch did not indicate that our results were overestimated by contamination
181 of the spike solution (Berthelot et al. 2015).

182 Samples for NH_4^+ were collected in 40 mL glass vials and analyzed by the fluorescence 183 method according to Holmes et al. (1999), using a Trilogy fluorometer (Turner Design).

184 **2.6.** Transparent exopolymeric particles (TEP)

Water samples (100 mL) were gently (< 150 mbar) filtered through 0.45 µm polycarbonate 185 filter (GE Water & Process Technologies). Filters were then stained with a solution of 0.02 % 186 Alcian blue (AB), 0.06 % acetic acid (pH of 2.5), and the excess dye was removed by a quick 187 188 deionized water rinse. Filters were then immersed in sulfuric acid (80 %) for 2 h, and the absorbance (787 nm) was measured spectrophotometrically (CARY 100, Varian). AB was 189 calibrated using a purified polysaccharide gum xanthan (GX) (Passow and Alldredge, 1995). 190 TEP concentrations (μg GX equivalents L⁻¹) were measured according to (Passow and 191 Alldredge, 1995). 192

193

194 2.7. Virus abundance

Total seawater (1 mL) was fixed with 0.5 % glutaraldehyde and snap frozen in liquid N₂ until 195 processed. Flow cytometry was conducted using an Influx Model 209S Mariner flow 196 cytometer and high-speed cell sorter equipped with a 488 nm 200 mW blue laser, 4 way sort 197 module, 2 scatter, 2 polarized and 4 fluorescence detectors (BD Biosciences). Viral abundance 198 was determined by staining fixed seawater samples with SYBR Gold (Life Technologies) and 199 measurements of green fluorescence (520 nm, 40 nm band pass). Samples were thawed, 200 201 diluted 25-fold in 0.22 µm-filtered Tris/EDTA (TE) buffer (pH 8), stained with SYBR Gold (0.5 - 1X final concentration), incubated for 10 min at 80°C in the dark, cooled to RT for 5 202 min, and mixed thoroughly by vortexing prior to counting on the Influx (Brussaard, 2003). 203 Viral abundance was analyzed using a pressure differential (between sheath and sample fluid) 204 of 0.7, resulting in a low flow rate for higher event rates of virus like particles counts. 205

206 2.8. Caspase activity

Biomass was collected on 25 mm, 5 µm pore-size polycarbonate filters and resuspended in 0.6-1 mL Lauber buffer [50 mM HEPES (pH 7.3), 100 mM NaCl, 10 % sucrose, 0.1 % 3-(3cholamidopropyl)-dimethylammonio-1-propanesulfonate, and 10 mM dithiothreitol] and sonicated on ice (four cycles of 30 seconds each) using an ultra-cell disruptor (Sonic Dismembrator, Fisher Scientific, Waltham, MA, USA). Cell extracts were centrifuged (10,000 212 x g, 2 min, room temperature) and supernatant was collected for caspase biochemical activity. Caspase-specific activity was determined by measuring the kinetics of cleavage for the 213 canonical fluorogenic caspase substrate (Z-IETD-AFC) at a 50 mM final concentration (using 214 Ex 400 nm and emission 505 nm; Synergy4 BioTek, Winooski, VT, USA), as previously 215 described in Bar-Zeev et al. (2013). Fluorescence was converted to a normalized substrate 216 cleavage rate using an AFC standard (Sigma) and normalized to total protein concentrations 217 obtained from the same samples. Total protein concentrations were determined by PierceTM 218 BCA Protein Assay Kit (Thermo Scientific product #23225). 219

220 2.9.16S rRNA gene sequencing and data analyses

221 Bacterial community diversity was analyzed by deep sequencing of the 16S rRNA gene in samples from two replicate bottles from experiment 1 (see section 1.2) at three time points 222 223 each. Seawater samples were filtered on 25 mm, 5 µm pore-size Supor filters (Pall Gelman Inc., Ann Arbor, Michigan), snap frozen in liquid nitrogen, and stored at -80 °C for later 224 extraction. Community genomic DNA was isolated from the filters using a phenol-chloroform 225 226 extraction method modified according to Massana et al. (1997). The 16S rRNA genes within community genomic DNA were initially amplified with conserved bacterial primers 27F and 227 1100R (Dowd et al., 2008) using a high fidelity polymerase (Phusion DNA polymerase, 228 Thermo Scientific) with an initial denaturation step of 95 °C for 3 min followed by 20 cycles 229 of 95 °C for 30 sec, 55 °C for 30 sec, and 72 °C for 45 sec. A secondary PCR (same 230 conditions) was performed for next-generation sequencing by using customized fusion primers 231 with different tag sequences. The tags were attached to the 27F primer and to the 338R primer 232 233 (Hamady et al., 2008) to obtain 340 bp fragments suitable for IonTorrent analysis. The use of nested PCR was used to minimize inclusion of false sequences into the sequenced material 234 (Dowd et al., 2008). After secondary PCR, all amplicon products were purified using Ampure 235 magnetic purification beads (Agencourt Bio- science Corporation, MA, USA) to exclude 236 primer-dimers. The amplicons were sequenced at the Bar-Ilan Sequencing Center, using an 237 Ion TorrentTM (Life Technologies, USA). 238

The adapter-clipped sequences were processed using tools and scripts from the UPARSE pipeline (Edgar, 2013). Reads from all samples were pooled for OTU calling. Reads were demultiplexed, primers and barcodes stripped using the script *fastq_strip_barcode_relabel.py*, leaving 42747 raw reads altogether for six samples. As suggested for OTU calling from single-end amplicon sequences (Edgar, 2013), sequences (mostly between 280 nt and 300 nt) 244 were trimmed to a fixed length of 280 nt, and shorter sequences were discarded (26740 trimmed raw reads remaining). For OTU clustering, trimmed raw reads were quality filtered 245 using the *-fastq filter* command with a maximum expected error rate (*-fastq maxee*) of 2 246 (21590 reads remaining), clustered into unicals (100 % identity) and the unicals sorted by 247 weight (number of sequences in the cluster). OTU clustering with an identity threshold of 0.98 248 was done using the -cluster_otus command on sorted unicals, with built-in chimera filtering. 249 250 To infer OTU abundances for each individual sample, the trimmed raw reads per sample (after a more relaxed quality filtering with -fastq_maxee 5) were mapped back to these OTUs with -251 usearch_global and a minimum identity of 98 %. For taxonomic classification, OTUs were 252 submitted to https://www.arb-silva.de/ngs/ and classified using the SINA aligner v1.2.10 and 253 database release SSU 123 (Quast et al., 2013). Sequences having a (BLAST alignment 254 coverage + alignment identity)/2 < 93 % were considered as unclassified and assigned to the 255 virtual group "No Relative" (5.58 % of OTUs). 256

257 2.10. RNA extraction and metatranscriptome sequencing

Metatranscriptomic sequencing was performed for three time points: peak surface 258 accumulation of the bloom (T_{0} , 12:00), 8 h (T_{8} 22:00), and 22 h (T_{22} 10:00) after T_{0} . Cells on 259 polycarbonate filters were resuspended in 1 mL PGTX [for 100 mL final volume: phenol 260 (39.6 g), glycerol (6.9 mL), 8-hydroxyquinoline (0.1 g), EDTA (0.58 g), sodium acetate (0.8 261 g), guanidine thiocyanate (9.5 g), guanidine hydrochloride (4.6 g), Triton X-100 (2 mL)] 262 (Pinto et al., 2009), and 250 μ l glass beads (diameter 0.1 – 0.25 mm). and sonicated on a cell 263 disruptor (Precellys, Peqlab, Germany) for 3 x 15 s at 6500 rpm. Tubes were placed on ice 264 between each 15 s interval. RNA was extracted by adding 0.7 mL chloroform and subsequent 265 266 phase separation. RNA was precipitated from the aqueous phase using 3 volumes of isopropanol at -20 °C overnight. Residual DNA was removed using the Turbo DNA-free Kit 267 268 (Ambion) after the manufacturer's instructions, but adding additional 1 μ l of DNase after 30 min of incubation and incubating another 30 min. RNA was purified using Clean & 269 Concentrator 5 columns (C&C 5) (Zymo Research, Freiburg, Germany). The pure RNA was 270 treated with Ribo-Zero rRNA Removal Kit (Bacteria) (Epicentre, Madison, USA) and 271 purified again with C&C 5. DNA contamination was tested and confirmed negative with a 40 272 cycle PCR using cyanobacteria-specific 16S primers. 273

For removal of tRNAs and small fragments, the RNA was purified with the Agencourt
RNAClean XP kit (Beckman Coulter Genomics, Danvers, USA). First-strand cDNA synthesis

276 for T₈ and T₂₂ samples was primed with a N6 randomized primer, after which the cDNAs were fragmented by ultrasound (4 pulses of 30 sec at 4 °C). Illumina TruSeq sequencing adapters 277 were ligated in a strand-specific way to the 5' and 3' ends and the resulting cDNAs were PCR-278 amplified to about 10-20 ng μL^{-1} using a high fidelity DNA polymerase. Randomly-primed 279 cDNA for T₀ samples was prepared using purified RNA without fragmentation followed by 280 ligation of Illumina TruSeq sequencing adapters to the 5' and 3' ends and fragmentation of 281 cDNA > ~700 bp with ultrasound (4 pulses of 30 sec at 4°C; targeting only cDNA > 700 nt). 282 After repairing ends, fragments were dA-tailed and Illumina TruSeq sequencing adapters were 283 284 ligated again to the 5' and 3' ends of the cDNA and re-amplified. Consequently, a small fraction of the T₀ reads was not strand-specific. All cDNAs were purified using the Agencourt 285 AMPure XP kit (Beckman Coulter Genomics, Danvers, USA) and 2 x 150 nt paired-end 286 sequences generated with an Illumina NextSeq500 sequencer by a commercial provider (vertis 287 AG, Freising, Germany). 288

289 **2.11.** Bioinformatics processing and analysis of metatranscriptome data

290 To remove adapters, perform quality trimming, and set a minimal length cutoff, raw fastq reads were processed with Cutadapt version 1.8.1 (Martin, 2011) in paired-end mode with a 291 minimum adapter sequence overlap of 10 nt (-O 10), an allowed error rate of 20 % (-e 0.2) in 292 the adapter sequence alignment, and a minimum base quality of 20. To remove residual 293 ribosomal RNA reads, the fastq files were further processed with SortMeRNA version 1.8 294 (Kopylova et al., 2012) with the accompanying standard databases in paired end mode, 295 resulting in 9,469,339 non-ribosomal reads for T_0 , 22,407,194 for T_8 , and 18,550,250 for T_{22} . 296 297 The fastq files with all non-ribosomal forward-reads were used for mapping against the 298 Trichodesmium erythraeum IMS101 genome with Bowtie2 (Langmead and Salzberg, 2012) in very-sensitive-local mode. This resulted in 51.9 % of T₀, 5.1 % of T₈, and 3.3 % of T₂₂ reads 299 mapped. Reads were counted per CDS feature as annotated in the genome of Trichodesmium 300 301 erythraeum (NC_008312.1) using htseq-count version 0.6.0 (Anders et al., 2014) and a count table generated with all read counts from T_0 , T_8 , and T_{22} . 302

For detection of differentially expressed genes from T_0 to T_8 and T_8 to T_{22} , the count table was processed with the statistical tool "Analysis of Sequence Counts" (ASC) (Wu et al., 2010). This tool is specifically designed to account for missing replicates by employing a model of biological variation of gene expression (Wu et al., 2010). The posterior probabilities (P) of a gene being > 2-fold differentially expressed (user specified threshold) between any two 308 samples is calculated using an empirical Bayesian analysis algorithm and an internal 309 normalization step. Differential expression of genes was defined as significant if P > 0.98.

310

311 3 Results

312 3.1. Setting the scene – *Trichodesmium* bloom development and bloom within 313 the lagoon.

314 Trichodesmium were present as part of the in-situ community in the lagoon at the outset of the VAHINE experiment. (Bonnet et al., 2015; Turk-Kubo et al., 2015). In the lagoon water, 315 temperatures were high (> 25 °C) and typical oligotrophic conditions of austral summer 316 prevailed. For the first 20 days of the experiment low abundance and biomass was measured 317 for primary and secondary production and specifically for diazotrophic populations (Fig. 1). 318 Total PON and POC in the lagoon fluctuated in the first 20 days of the VAHINE experiment 319 with values ranging between 0.6-1.1 μ mol L⁻¹ and 5-11 respectively. On the morning of day 320 23, values were 0.9 and 9.3 μ mol L⁻¹ PON and POC, respectively (Fig. 1c-d). The total Chl a 321 concentrations ranged between 0.18-0.26 μ g L⁻¹ from days 1-20 (Fig. 1a). The increase in Chl 322 323 a concentrations reflect the composite signature of the total phototrophic community [detailed in (Leblanc et al., 2016; Van Wambeke et al., 2015)] and is not specific to Trichodesmium 324 biomass. Low abundances of Trichodesmium were measured in the lagoon waters throughout 325 the first three weeks of the project (Turk-Kubo et al., 2015), with Trichodesmium-associated 326 16S counts ranging from 0.1 to 0.4 % of the total number of 16S tags (Pfreundt et al., 2016). 327 During the first eight days of sampling, *Trichodesmium* abundance as measured by *nifH* gene 328 real-time PCR ranged from 3.4 x 10^2 -6.5 x 10^3 nifH copies L⁻¹. By days 14 and 16, 329 *Trichodesmium* accounted for 15 % of the total diazotroph population (with 1.1-1.5 x 10^4 nifH 330 copies L⁻¹) increasing by day 22 to 42 % of the diazotroph population (1.4 x 10^5 nifH copies L⁻¹) 331 ¹) (Turk-Kubo et al., 2015). By the morning of day 23, Chl *a* increased to 0.39 μ g L⁻¹ in the 332 upper 1 m depth (Fig. 1a), yet Trichodesmium was still not visually observed at this time as a 333 bloom on the sea surface. Phycoerythrin concentrations fluctuated between 0.1-0.4 μ g L⁻¹ 334 during days 1-14 and then increased to a maximal peak of $> 0.8 \ \mu g \ L^{-1}$ on day 21 with values 335 ~ 0.5 μ g L⁻¹on day 23 reflecting both the doubling in *Synechococcus* biomass (days 15-23) as 336 well as increasing Trichodesmium (days 21-23) (Leblanc et al., 2016). N₂ fixation rates in the 337 lagoon waters ranged between 0.09 -1.2 nmol N L⁻¹ h⁻¹ during the pre-bloom period (Fig. 1c) 338 and on the morning of day 23 measured 0.5 nmol $L^{-1} h^{-1}$ (Fig. 1c). 339

Zooplankton populations in the lagoon fluctuated around 5000 individuals m⁻³ and increased 340 from day 9 to 16 to peak at ~ 14000 individuals m^{-3} (Hunt et al., 2016). From day 16 to day 23 341 the total zooplankton population declined to ~ 8000 individuals m^{-3} with harpacticoid 342 copepods including grazers of Trichodesmium (Macrosetella gracilis, Miracia efferata, and 343 344 Oculosetella gracilis) comprising < 1.5 % of total zooplankton community in the lagoon (Hunt et al., 2016). Virus like particles (VLP) ranged from 1-6 x 10^6 mL⁻¹ throughout the first 345 22 days of the VAHINE experiment and displayed a ~ 2-4 day oscillation (i.e., increasing for 346 2 days, then declining for the next 3 days, etc.) with mean values of $3.8 \times 10^6 \text{ mL}^{-1}$ (Fig. 1b). 347 VLP counts in surface waters on day 23 were $1.8 \times 10^6 \text{ mL}^{-1}$ (Fig. 1b), just prior to the 348 appearance of the Trichodesmium surface bloom. VLPs did not show any distinct correlations 349 with total biomass indices such as PON and POC during the pre-bloom sampling (Fig. 1b-d). 350

Depth-averaged dissolved inorganic phosphorus (DIP) concentrations in the lagoon waters 351 were low at 0.039 \pm 0.001 μ M, with a relatively stable DIP turnover time (T_{DIP}) of 1.8 \pm 0.7 d 352 for the first 15 days, that declined to 0.5 ± 0.7 by day 23 (Berthelot et al., 2015). Alkaline 353 phosphatase activity (APA), which hydrolyzes inorganic phosphate from organic phosphorus, 354 increased ~ 3 fold, from 1.8 \pm 0.7 (average of days 1-4) to 5.0 \pm 1.4 nmole L⁻¹ h⁻¹ (average of 355 days 19-23) (Van Wambeke et al., 2015) demonstrating a response in metabolic activity 356 related to P acquisition for the microbial community probably related to the decreasing 357 availability of DIP in the lagoon waters. 358

359 On day 23 (February 4) of the VAHINE measurements, dense surface accumulations of Trichodesmium were observed at midday (12:00 h) (Fig. 2a-c). Ambient air temperatures (~ 360 361 25 °C) increased to over 26 °C and the winds decreased to < 5 knots. These accumulations (hereafter blooms) appeared in the typical "slick" formations of dense biomass in ribbons 362 363 visible on the surface seawater and spread out over tens of meters in the lagoon water outside the mesocosms (Fig. 2a-c). *Trichodesmium* abundance in these patches was extremely variable 364 with Chl *a* concentrations exceeding 5 mg L⁻¹ within dense patches and trichome abundance > 365 10,000 trichomes mL. These surface accumulations were visible and sampled again 5 h later 366 (experiment 2), yet by the next morning, no such slicks or patches of dense biomass were 367 observed or measured in the lagoon. The disappearance of the *Trichodesmium* in the lagoon 368 water whether by drifting away, sinking to depth, or any other factor, prevented further 369 investigation of these populations. 370

371

372 **3.2.** Investigating *Trichodesmium* mortality in experimental microcosms.

373 3.2.1 Changes in *Trichodesmium* biomass and associated microbial 374 communities.

375 The spatially patchy nature of *Trichodesmium* blooms in the lagoon (Fig. 2a-c), and the rapid temporal modifications in water-column abundance of filaments and colonies probably 376 induced (primarily) by physical drivers (turbulence and wind-stress), complicate in-situ 377 sampling when targeting changes in specific biomass. To overcome this, we collected 378 Trichodesmium populations from the surface midday bloom and examined the physiological, 379 biochemical, and genetic changes occurring with time until the biomass crashed ~ 24 h (see 380 methods section 2.2) (Fig. 2 and Fig. 3). In these enclosed microcosms, Trichodesmium 16S 381 copies comprised > 90 % of total copies (Fig. 3) enabling the use Chl a to follow changes in 382 its biomass (Fig. 2f). Maximal Chl *a* concentrations in the incubations (> 150 \pm 80 µg L⁻¹; 383 n=6) were measured at the start of the incubation soon after the biomass collection and 384 resuspension in FSW. These Trichodesmium populations collapsed swiftly over the next day 385 with Chl *a* concentrations declining to 24 μ g L⁻¹ and 11 μ g L⁻¹ Chl *a* after 10 and 22 h, 386 387 respectively (Fig. 2f).

In experiment 1 we characterized the microbial community associated with the 388 Trichodesmium biomass within the microcosms by 16S rRNA gene sequencing from two 389 replicate bottles (experiment 1). At T₀ 94 % and 93 % of the obtained 16S tags in both 390 replicates (Fig. 3) were of the Oscillatoriales order (phylum Cyanobacteria), with 99.9 % of 391 these sequences classified as Trichodesmium spp. (Fig. 3). In both replicates, the temporal 392 decline of Trichodesmium biomass coincided with an increase in Alteromonas 16S tags, but 393 394 this development temporally lagged in replicate 1 compared to replicate 2 (Fig. 3). Six hours (T_6) after the surface bloom was originally sampled (T_0) , over 80 % of 16S tags from replicate 395 1 were characterized as Trichodesmium. 14 h after T₀, Alteromonadales and Vibrionales 396 replaced Trichodesmium now constituting only 9 % of 16S tags (Fig. 3). In replicate 2, 397 Trichodesmium declined by 80 % 6 h after T₀, with Alteromonadales and Flavobacteriales 398 399 comprising the bulk of the biomass 18 hours after the start of incubations (Fig.3).

400 The rate of decline in *Trichodesmium* biomass within the 4.6 L microcosms paralleled that of 401 *Trichodesmium* collected from the surface accumulations at 17:00 and incubated in 20 L 402 carboys under ambient conditions for > 72 h (defined hereafter as experiment 2: Fig. 4). Here, 403 *Trichodesmium* biomass decreased by > 80 % within 24 h of incubations with trichome 404 abundance declining from ~2500 trichomes mL⁻¹ at bloom collection to ~ 495 trichomes mL⁻¹ 405 (Fig. 4a). No direct correlation was observed between the decline of *Trichodesmium* and viral 406 populations. VLP abundance at the time of the surface bloom sampling was at a maximum of 407 8.2 x 10^6 mL⁻¹ (Fig. 4a), decreasing to 5.7 x 10^6 mL⁻¹ in the next 4 h then remaining stable 408 throughout the crash period (within the next 42 h) averaging ~ 5 x $10^6 \pm 0.7$ mL⁻¹ (Fig. 4a).

As Trichodesmium crashed in the experimental incubations, high values of NH_4^+ were 409 measured (Fig 4b). In experiment 2, NH_4^+ increased exponentially from 73 \pm 0.0004 nmol 410 $NH_4^+L^{-1}$ when the surface bloom was collected and placed in the carboys (17:00 h) to 1490 ± 411 686 after 24 h and values > 5000 nmol L^{-1} 42 h after the incubation start (Fig. 3b). The high 412 ammonia declined somewhat by the end of the experiment (after 72 h), yet was still high at 413 3494 ± 834 nmol L⁻¹. Concurrently with the high NH₄⁺ concentrations, and despite the dying 414 Trichodesmium, we measured an increase N2-fixation rates. N2-fixation rose from 1.5 nmol N 415 L^{-1} h⁻¹ at T₀ to 3.5 ± 2.8 nmol N L^{-1} h⁻¹ 8 h after incubations began and 11.7 ± 3.4 nmol N L^{-1} 416 24 h later (Fig 4b). These high values represent other diazotrophs including UCYN-types and 417 diatom-diazotroph associations that flourished after the Trichodesmium biomass had declined 418 419 in the carboys (Bonnet et al. 2016b; Turk-Kubo personal communication). POC and PON, representing the fraction of C and N incorporated into biomass, ranged between 5.2-11.2 µmol 420 C L⁻¹ and 0.6-1.1 µmol N L⁻¹ during pre-bloom periods (Fig. 1b) and 12.6 ± 4.6 µmol C L⁻¹ 421 and $1.3 \pm 0.5 \mu$ mol N L⁻¹ when the surface bloom was sampled (Fig. 4b-c). 24 hours after 422 collection of bloom biomass POC increased ~ 6-fold to $63.2 \pm 15 \mu mol C L^{-1}$ and PON 423 increased 10-fold to $10 \pm 3.3 \mu$ mol N L⁻¹ (Fig. 4b-c). After 72 h, total POC was $62 \pm 4 \mu$ mol C 424 L^{-1} (Fig. 4c) and PON increased to 14.1 ± 6 µmol N L^{-1} (Fig. 4b). 425

426 Organic carbon in the form of TEP is secreted when *Trichodesmium* is stressed and 427 undergoing PCD (Bar-Zeev et al., 2013; Berman-Frank et al., 2004). TEP concentrations in 428 the lagoon waters during the pre-bloom period (first 20 days) fluctuated around ~ 350 µg gum 429 xanthan (GX) L⁻¹ (Fig. 1d) that increased to ~ 500 µg GX L⁻¹ on day 22 (Fig. 1d). During the 430 time of biomass collection from the surface bloom TEP concentration exceeded 700 µg GX L⁻¹ 431 ¹ (Fig. 4c). After biomass enclosure (experiment 2) TEP concentrations declined to 420 ± 35 432 µg GX L⁻¹ and subsequently to $180 \pm 25 \mu g$ GX L⁻¹ 42 h and 72 h after T₀ (Fig. 4c).

433 3.2.2. Genetic responses of stressed Trichodesmium

434 Metatranscriptomic analyses of the *Trichodesmium* biomass were conducted in samples from 435 experiment 1, at T_0 , T_8 , and T_{22} (Fig. S1). We examined differential expression during this 436 period by investigating a manually curated gene suite including specific pathways involved in 437 P and Fe uptake and assimilation, PCD, or gas vesicle synthesis. Genes involved in the acquisition and transport of inorganic and organic P sources were upregulated, concomitant 438 with biomass demise; significantly higher expression levels were evident at T_8 and T_{22} 439 compared to T_0 (Table S1). Abundance of alkaline phosphatase transcripts, encoded by the 440 phoA gene (Orchard et al., 2003), increased significantly (~ 5 fold) from T_0 to T_{22} (Fig. 5a). 441 The transcript abundance of phosphonate transporters and C-P lyase genes (phnC, phnD, 442 *phnE*, *phnH*, *phnI*, *phnL* and *phnM*) increased significantly (5-12 fold) between T_0 and both T_8 443 and T_{22} (Fig. 5a, Table S1). Of the phosphite uptake genes, only *ptxA* involved in the 444 445 phosphite (reduced inorganic phosphorus compound) uptake system, and recently found to operate in Trichodesmium (Martínez et al., 2012; Polyviou et al., 2015) was significantly 446 upregulated at both T_8 and T_{22} compared to T_0 (4.5 and 7 fold change respectively). The two 447 additional genes involved in phosphite uptake, *ptxB* and *ptxC*, did not change significantly, as 448 Trichodesmium biomass crashed (Fig. 5a). 449

450 Fe limitation induces PCD in Trichodesmium (Berman-Frank et al., 2004; Berman-Frank et 451 al., 2007) we therefore examined genetic markers of Fe stress. At the time of surface bloom sampling (experiment 1, T_0), Fe stress was indicated by higher differential expression of 452 453 several genes. The *isiB* gene encodes flavodoxin and serves as a common diagnostic indicator of Fe stress in Trichodesmium, since it may substitute for Fe-S containing ferredoxin (Bar-454 455 Zeev et al., 2013; Chappell and Webb, 2010). Transcripts of isiB were significantly higher at T_0 (3-fold) than at T_8 and T_{22} (Fig. 5b, Table S1). The chlorophyll-binding protein IsiA is 456 induced in cyanobacterial species under Fe or oxidative stress to prevent oxidative damage 457 (Laudenbach and Straus, 1988). Here *isiA* transcripts increased 2- and 3- fold from T_0 to T_8 458 and T₂₂, respectively (Fig. 5b, Table S1). The Fe transporter gene *idiA* showed a transient 459 higher transcript accumulation only at T₈. As the health of Trichodesmium declined, 460 transcripts of the Fe-storage protein ferritin (*Dps*) decreased by > 70 % at T₂₂ (Fig. 5b, Table 461 **S**1) 462

463 3.2.3. PCD-induced demise.

464 Our earlier work demonstrating PCD in *Trichodesmium* allowed us to utilize two independent 465 biomarkers to investigate PCD induction during *Trichodesmium* demise, namely changes in 466 catalytic rates of caspase-specific activity (Berman-Frank et al., 2004; Berman-Frank et al., 467 2007) and levels of metacaspase transcript expression (Bar-Zeev et al., 2013). When the 468 surface bloom was sampled (experiment 1, T₀), protein normalized caspase-specific activity 469 was 0.23 ± 0.2 pmol mg protein⁻¹ min⁻¹ (Fig. 6a). After a slight decline in the first 2 h, caspase 470 activity increased throughout the experiment with 10 fold higher values (2.9 ± 1.5 pmol L⁻¹ 471 mg protein⁻¹ min⁻¹) obtained over the next 22 h as the bloom crashed (Fig. 6a).

472 We followed transcript abundance over the demise period for the 12 identified metacaspase genes in Trichodesmium [(Asplund-Samuelsson et al., 2012; Asplund-Samuelsson, 2015; 473 Berman-Frank et al., 2004)]; TeMC1 (Tery_2077), TeMC2 (Tery_2689), TeMC3 (Tery_3869), 474 TeMC4 (Tery 2471), TeMC5 (Tery 2760), TeMC6 (Tery 2058), TeMC7 (Tery 1841), 475 TeMC8 (Tery_0382), TeMC9 (Tery_4625), TeMC10 (Tery_2624), TeMC11 (Tery_2158), and 476 477 TeMC12 (Tery_2963)] (Fig. 6b, Table S1). A subset of these genes was previously implicated in PCD of Trichodesmium cultures in response to Fe and light stress (Bar-Zeev et al., 2013; 478 479 Berman-Frank et al., 2004; Bidle, 2015). Here, we interrogated the entire suite of metacaspases in natural *Trichodesmium* populations. As the biomass crashed from T_0 to T_{22} , 7 480 out of 12 metacaspases (TeMC1, TeMC3, TeMC4, TeMC7, TeMC8, TeMC9, and TeMC11) 481 482 were significantly upregulated 8 and 22 h after T₀ (Fig. 6b). For these genes, transcript abundance increased 2.3- to 5.3-fold 8 h after T_0 and 3.5-6.2-fold 22 h after T_0 (Fig. 6b, Table 483 S1) TeMC5 and TeMC10 transcripts increased significantly after 22 h by 2.9- and 3.2 fold, 484 respectively. TeMC6 was upregulated 2.9-fold after 8 h. TeMC2 transcripts did not 485 significantly change over time. We did not detect any expression of TeMC12 throughout the 486 487 experiment.

488 Export flux can be enhanced by PCD-induced sinking (Bar-Zeev et al., 2013) as PCD in Trichodesmium results in degradation of internal components, especially gas vesicles that are 489 required for buoyancy (Berman-Frank et al., 2004). Although we did not measure changes in 490 491 buoyancy itself, we observed rapid sinking of the Trichodesmium biomass in the bottles and carboys. The metatranscriptomic analyses demonstrated that, excluding one copy of 492 493 gvpL/gvpF, encoding a gas vesicle synthesis protein, gas vesicle protein (gvp) genes involved in gas-vesicle formation (gvpA, gvpN, gcpK, gvpG and gvpL/gcpF) were all significantly 494 495 downregulated relative to T_0 (Fig. 7, Table S1).

- 496 **4 Discussion**
- 497

4.1. Mortality processes of *Trichodesmium* – incubation results.

498 **4.1.1 Grazer and virus influence.**

Our microcosm incubations allowed us to specifically focus on the loss factors and show theinvolvement of biotic and abiotic stressors in inducing PCD and mechanistically impacting the

demise and fate of a natural *Trichodesmium* bloom. We appreciate that the enclosure of the biomass in bottles and carboys may accelerate the processes occurring in the natural lagoon setting. Yet, the published rates of *Trichodesmium* mortality from field studies (Rodier and Le Borgne, 2010) indicate that these can parallel our loss rates with natural bloom demise occurring 24-48 h after peak of biomass.

506 We focused initially on biotic factors that could impact the incubated *Trichodesmium* biomass.

507 The low number of harpacticoid zooplankton specific to Trichodesmium (O'Neil and Roman,

1994; O'Neil, 1998) in the lagoon (Hunt et al., 2016) and especially those in the bottles
(personal observation) refutes the hypothesis that grazing caused the massive mortality of *Trichodesmium* biomass in our experimental incubations.

Viruses have been increasingly invoked as key agents terminating phytoplankton blooms 511 512 (Brussaard et al., 2005; Jacquet et al., 2002; Lehahn et al., 2014; Tarutani et al., 2000; Vardi et al., 2012). In Trichodesmium, phages have been implicated in bloom crashes, but this 513 514 mechanism has yet to be unequivocally proven (Hewson et al., 2004; Ohki, 1999); indeed, no specific Trichodesmium phage has been isolated or characterized to date (Brown et al., 2013). 515 516 Here, total VLP abundance was highest at the time of sampling from the surface Trichodesmium bloom and at the start of the incubation at ~ 8 x 10^6 VLPs mL⁻¹ it actually 517 declined 2 fold in the first eight hours of incubation before increasing over the next 32 h (Fig. 518 519 4a). While our method of analysis cannot distinguish between phages infecting Trichodesmium from those infecting other marine bacteria, it argues against a massive, phage-520 induced lytic event of *Trichodesmium*. Such an event would have yielded a notable burst of 521 VLPs upon bloom crash, especially considering the high *Trichodesmium* biomass observed. 522 The coincidence between the maximal abundance of VLPs and highest Trichodesmium 523 biomass is counter to viruses serving as the mechanism of mortality in our incubation 524 experiments. Nonetheless, virus infection itself may be a stimulant for community N₂ fixation 525 perhaps by releasing key nutrients (i.e., P or Fe) upon lysis of surrounding microbes (Weitz 526 and Wilhelm, 2012). Although we did not characterize them here, it is indeed possible that 527 Trichodesmium-specific phages were present in our incubation experiments and they may 528 529 have exerted additional physiological stress on resident populations, facilitating PCD induction. Virus infection increases the cellular production of reactive oxygen species (ROS) 530 531 (Evans et al., 2006; Vardi et al., 2012), which in turn can stimulate PCD in algal cells 532 (Berman-Frank et al., 2004; Bidle, 2015; Thamatrakoln et al., 2012). Viral attack can also 533 directly trigger PCD as part of an antiviral defense system activated to limit virus production

and prevent massive viral infection (Bidle and Falkowski, 2004; Bidle, 2015; Georgiou et al.,1998).

536 **4.1.2 Stressors impacting mortality.**

Nutrient stress can be acute or chronic to which organisms may acclimate on different time 537 scales. Thus, for example, the consistently low DIP concentrations measured in the lagoon 538 during the 22 days preceding the *Trichodesmium* surface bloom probably enabled acclimation 539 responses such as induction of APA and other P acquisition systems. Trichodesmium has the 540 ability to obtain P via inorganic and organic sources including methylphosphonate, 541 ethylphosphonate, 2-aminoethylphosphonate (Beversdorf et al., 2010; Dyhrman et al., 2006), 542 543 and via a phosphite uptake system (PtxABC) that accesses P via the reduced inorganic compound phosphite (Martínez et al., 2012; Polyviou et al., 2015). Our metatranscriptomic 544 data demonstrated upregulated expression of genes related to all three of these uptake systems 545 (DIP, phosphonates, phosphites) 8 and 22 h after incubation began, accompanying biomass 546 547 demise (Fig. 5a). This included one gene for phosphite uptake (*ptxA*) and several genes from the phosphonate uptake operon (phnDCEEGHIJKLM) (Hove-Jensen et al., 2014). 548 549 Upregulated expression of phnD, phnC, phnE, phnH, phnI, phnJ, phnK, phnL and phnM 550 occurred as the *Trichodesmium* biomass crashed (Fig. 5a, Table S1), consistent with previous 551 results demonstrating that *phnD* and *phnJ* expression levels increased during DIP depletion 552 (Hove-Jensen et al., 2014). It is likely that during bloom demise, the C-P lyase pathway of remaining living cells was induced when DIP sources were extremely low, while POP and 553 DOP increased along with the decaying organic matter. The ability to use phosphonates or 554 phosphites as a P source can provide a competitive advantage for phytoplankton and bacteria 555 in P-depleted waters (Coleman and Chisholm, 2010; Martinez et al., 2010). Thus, it is 556 557 puzzling why dying cells would upregulate *phn* genes or *phoA* transcripts after 22 h incubation (Fig. 5a). A more detailed temporal resolution of the metatranscriptomic analyses may 558 elucidate the expression dynamics of these genes and their regulating factors. Alternatively, in 559 PCD-induced populations, a small percentage remains viable and resistant as either cysts 560 (Vardi et al., 1999) or hormogonia (Berman-Frank et al., 2004) that can serve as the inoculum 561 562 for future blooms. It is plausible that the observed upregulation signal was attributable to these sub-populations. 563

The concentrations of dissolved and bioavailable Fe were not measured in the lagoon water during the experimental period as Fe is typically replete in the lagoon (Jacquet et al., 2006). However, even in Fe-replete environments such as the New Caledonian lagoon, dense patches 567 of cyanobacterial or algal biomass can deplete available resources and cause limited microenvironments (Shaked, 2002). We obtained evidence for Fe stress using several proxy genes 568 demonstrating that enhanced cellular Fe demand occurred during the bloom crash (Table S1). 569 Trichodesmium's strategies of obtaining and maintaining sufficient Fe involves genes such as 570 isiB. isiB was highly expressed when biomass accumulated on the surface waters, indicative 571 for higher Fe demand at this biomass load (Bar-Zeev et al., 2013; Chappell and Webb, 2010). 572 Transcripts for chlorophyll-binding, Fe-stress-induced protein A (IsiA) increased (albeit not 573 significantly) 3-fold over 22 h of bloom demise (Fig. 5b, Table S1). In many cyanobacteria, 574 575 isiA expression is stimulated under Fe stress (Laudenbach and Straus, 1988) and oxidative stress (Jeanjean et al., 2003) and functions to prevent high-light induced oxidative damage by 576 increasing cyclic electron flow around the photosynthetic reaction center photosystem I 577 (Havaux et al., 2005; Latifi et al., 2005; Michel and Pistorius, 2004). Dense surface blooms of 578 Trichodesmium are exposed to high irradiance (on day 23 average PAR was 3000 µmol 579 photons m⁻² s⁻¹). It is possible that high Fe demand combined with the oxidative stress of the 580 high irradiance induced the higher expression of *isiA* (Fig. 5b). As cell density and associated 581 self-shading of Trichodesmium filaments decreased during bloom crash, light-induced 582 oxidative stress is likely the principal driver for elevated *isiA* expression. 583

584 The gene *idiA* is another environmental Fe stress biomarker that allows acquisition and transfer of Fe through the periplasm into the cytoplasm (Chappell and Webb, 2010). In our 585 incubation, upregulated expression of *idiA* (an ABC Fe⁺³ transporter) was evident after 8 h. 586 This is consistent with increasing Fe-limitation, as Trichodesmium abundance (measured via 587 588 16S rRNA gene sequencing) was still high at T_6 (after 6 h of incubations) (replicate 1). These findings are consistent with proteomics analyses from deplete iron (0 µM Fe) Trichodesmium 589 590 cultures which revealed an increase in IdiA protein expression (Snow et al., 2015). Lastly, our metatranscriptomic data highlighted a reduction in Fe storage and utilization, as the expression 591 of Fe-rich ferritin-like DPS proteins (Castruita et al., 2006), encoded by dpsA, decreased ~ 5 592 fold by the time that most of the biomass had crashed (T_{22}) (Fig. 5b, Table S1). dpsA was also 593 downregulated under Fe-replete conditions in Synechococcus (Mackey et al., 2015), but the 594 595 downregulation observed here is more likely related to Trichodesmium cells dying and downregulating Fe-demanding processes such as photosynthesis and N₂ fixation. 596

597 **4.1.3. Programmed cell death (PCD) and markers for increased export flux.**

598 The physiological and morphological evidence of PCD in *Trichodesmium* has been previously documented in both laboratory (Bar-Zeev et al., 2013; Berman-Frank et al., 2004) and 599 environmental cultures collected from surface waters around New Caledonia (Berman-Frank 600 et al., 2004). Here, we confirmed characteristic features of autocatalytic PCD in 601 602 Trichodesmium such as increased caspase-specific activity (Fig. 6a), globally enhanced metacaspase expression (Fig. 6b), and decreased expression of gas vesicle maintenance (Fig. 603 604 7). Metatranscriptomic snapshots interrogating expression changes in all Trichodesmium 605 metacaspases (Fig. 6b) generally portrayed upregulated expression concomitant with biomass 606 decline. Our results are consistent with previous observations that Fe-depleted PCD-induced laboratory cultures of Trichodesmium IMS101 had higher expression levels of TeMC1 and 607 TeMC9 compared to healthy Fe-replete cultures (Bar-Zeev et al., 2013; Berman-Frank et al., 608 2004). To our knowledge, this is the first study examining expression levels of metacaspases 609 in environmental Trichodesmium samples during a natural bloom. 11 of the 12 annotated 610 metacaspases in Trichodesmium were expressed in all 3 metatranscriptomes from the surface 611 612 bloom. To date, no specific function has been determined for these metacaspases in Trichodesmium other than their association with cellular stress and death. Efforts are 613 underway to elucidate the specific cellular functions, regulation, and protein interactions of 614 615 these Trichodesmium metacaspases (Pfreundt et al., 2014; Spungin et al., In prep).

616 In cultures and isolated natural populations of *Trichodesmium*, high caspase-like specific activity is correlated with the initial induction stages of PCD with activity declining as the 617 biomass crashes (Bar-Zeev et al., 2013; Berman-Frank et al., 2004; Berman-Frank et al., 618 2007). Here, caspase-like activity increased with the crashing populations of Trichodesmium 619 620 (Fig. 5a). Notably, maximal caspase activities were recorded at T₂₃, after which most Trichodesmium biomass had collapsed. The high protein-normalized caspase-specific activity 621 may be a result of a very stressed and dying sub-population of *Trichodesmium* that had not yet 622 succumbed to PCD (Berman-Frank et al., 2004). Alternatively, the high caspase-like activity 623 624 may be attributed to the large population of *Altermomonas* bacteria that were associated with the remaining detrital Trichodesmium biomass. However, currently, we are unaware of any 625 publications demonstrating high cellular caspase-specific activity in clades of y-626 Proteobacteria. 627

Gas vesicles are internal structures essential for maintaining buoyancy of *Trichodesmium* populations in the upper surface waters enabling them to vertically migrate and respond to light and nutrient requirements (Capone et al., 1997; Walsby, 1978). Mortality via PCD causes 631 a decline in the number and size of cellular gas vesicles in *Trichodesmium* (Berman-Frank et al., 2004) and results in an enhanced vertical flux of trichomes and colonies to depth (Bar-632 Zeev et al., 2013). Our metatranscriptomic data supported the subcellular divestment from gas 633 vesicle production during bloom decline, as the expression of vesicle-related genes was 634 downregulated (Fig. 7). In parallel, TEP production and concentration increased to $> 800 \ \mu g$ 635 GX L⁻¹, a 2-fold increase from pre-bloom periods (Fig. 1d and Fig. 4c). When nutrient uptake 636 is limited, but CO₂ and light are sufficient, uncoupling occurs between photosynthesis and 637 growth (Berman-Frank and Dubinsky, 1999), leading to increased production of excess 638 639 polysaccharides, such as TEP, and corresponding with high TEP found in bloom decline phases rather than during the increase in population density (Engel, 2000; Smetacek, 1985). In 640 earlier studies we demonstrated that PCD-induced demise in Trichodesmium is characterized 641 by an increase in excreted TEP, (Berman-Frank et al., 2007) and enhanced sinking of 642 particulate organic matter (Bar-Zeev et al., 2013). TEP itself may be positively buoyant 643 (Azetsu-Scott and Passow, 2004), yet its stickiness causes aggregation and clumping of cells 644 and detritus, ultimately enhancing sinking rates of large aggregates including dying 645 Trichodesmium (Bar-Zeev et al., 2013). 646

647 **4.1.4.** Changes in microbial community with *Trichodesmium* decline.

In the incubations, other diazotrophic populations succeeded the declining Trichodesmium 648 649 biomass as indicated by increasing N₂ fixation rates, POC, and PON (Fig. 4b). In experiment 2, based on qPCR of targeted diazotrophic phylotypes, the diazotroph community composition 650 shifted from being dominated by Trichodesmium spp. and unicellular groups UCYN-A1, 651 UCYN-A2, and UCYN-B (T0), to one dominated by diatom-diazotroph associations Het-1 652 and Het-2 (T₇₂) (Bonnet et al. 2016b; Turk-Kubo, personal communication). In experiment 1 653 654 heterotrophic bacteria thrived and increased in abundance as the Trichodesmium biomass crashed (Fig. 3). 655

656 Trichodesmium colonies host a wide diversity of microorganisms including specific epibionts, viruses, bacteria, eukaryotic microorganisms and metazoans (Hewson et al., 2009; Hmelo et 657 658 al., 2012; Ohki, 1999; Paerl et al., 1989; Sheridan et al., 2002; Siddiqui et al., 1992; Zehr, 1995). Associated epibiont bacterial abundance in dilute and exponentially growing laboratory 659 660 cultures of Trichodesmium is relatively limited (Spungin et al., 2014) compared to bloom conditions (Hewson et al., 2009; Hmelo et al., 2012). Proliferation of Alteromonas and other 661 662 γ -Proteobacteria during biomass collapse (Fig. 3) confirms their reputation as opportunistic microorganisms (Allers et al., 2008; Hewson et al., 2009; Frydenborg et al., 2014; Pichon et 663

664 al., 2013). Such organisms can thrive on the influx of organic nutrient sources from the decaying Trichodesmium as we observed (Fig. 3). Furthermore, the increase of organic matter 665 including TEP produced by the stressed Trichodesmium (Fig. 1d and Fig. 4c) probably 666 stimulated growth of these copiotrophs. Moreover, as the Trichodesmium biomass declined in 667 the carboys, the high concentrations of NH_4^+ (> 5000 nmol L⁻¹) (Fig. 4b) sustained both 668 autotrophic and heterotrophic organisms (Berthelot et al., 2015; Bonnet et al., 2015; Bonnet 669 et., 2016b). Thus, the increase in volumetric N₂ fixation and PON that was measured in the 670 incubation bottles right after the Trichodesmium crash in experiment 2 (Fig 4b) probably 671 672 reflects both the enhanced activity of other diazotrophs (see above and Bonnet et al. 2016b) and resistant residual Trichodesmium trichomes (Berman-Frank et al. 2004) with increased 673 cell specific N₂ fixation. This scenario is consistent with the hypothesis that PCD induction 674 and death of a fraction of the population confers favorable conditions for survival and growth 675 of individual cells (Bidle and Falkowski, 2004). 676

677

4.2. Implications for the lagoon system and export flux.

Phytoplankton blooms and their dense surface accumulations occur under favorable physical 678 679 properties of the upper ocean (e.g. temperature, mixed-layer depth, stratification) and specifically when division rates exceed loss rates derived from grazing, viral attack, and 680 sinking or export from the mixed layer to depth (Behrenfeld, 2014). Although physical drivers 681 682 such as turbulence and mixing may scatter and dilute these dense accumulations, the rapid disappearance of biomass in large sea-surface *Trichodesmium* blooms (within 1-2 d in the 683 lagoon waters) (Rodier and Le Bourne 2010) suggests loss of biomass by other mechanisms. 684 The lack of Trichodesmium developing within the VAHINE mesocosms and the spatial-685 temporal variability of the surface bloom in the lagoon prohibited *in-situ* sampling of the same 686 biomass for several days and prevented conclusions regarding in-situ mortality rates and 687 export flux. Furthermore, within these dense surface populations as well as in the microcosm 688 and carboy experiments, nutrient availability was probably extremely limited due to high 689 690 demand and competition (Shaked 2002). PCD induced by Fe-depletion experiments with laboratory cultures and natural populations results in rapid biomass demise typically beginning 691 692 after 24 h with > 90 % of the biomass crashing 3 to 5 days after induction (Bar-Zeev et al., 2013; Berman-Frank et al., 2004; Berman-Frank et al., 2007). In similar experiments with P-693 694 depletion, Trichodesmium biomass did not crash rapidly. Rather, limitation induced colony 695 formation and elongation of trichomes (Spungin et al., 2014) and the cultures could be 696 sustained for another couple of weeks before biomass declined significantly (unpublished 697 data). The responses we quantified from the dying *Trichodesmium* in the carboys and bottles (Fig. 3-7) were similar to those obtained from controlled laboratory experiments where the 698 nutrient stressors P and Fe were validated individually. However, the rapid response here 699 probably reflects an exacerbated reaction due to the simultaneous combination of different 700 701 stressors and the presence of biotic components that can compete for and utilize the organic resources (carbon, nitrogen, phosphorus) generated by the dying Trichodesmium. In the 702 703 lagoon, production of TEP by stressed biomass combined with the degradation of gas vesicles and enhanced aggregation will cause such surface accumulations or blooms to collapse 704 705 leading to rapid vertical export of newly fixed nitrogen and carbon in the ocean.

706

707 **5 Conclusions and implications**

708 We demonstrate that the rapid demise of a Trichodesmium surface bloom in New Caledonia, with the disappearance of > 90 % of the biomass within 24 h in 4.5 L bottle incubations, 709 710 displayed cellular responses to P and Fe stress and was mediated by a suite of PCD genes. Virus infection and lysis did not appear to directly cause the massive biomass decline. 711 712 Although virus infection may have modulated the cellular and genetic responses to enhance 713 PCD-driven loss processes. Quorum sensing among epibionts (Hmelo et al., 2012; Van Mooy 714 et al., 2012), allelopathic interactions, and the production of toxins by Trichodesmium (Guo 715 and Tester, 1994; Kerbrat et al., 2010) are additional factors that could be important for a concerted response of the Trichodesmium population, yet we did not examine them here. 716 Collectively, they would facilitate rapid collapse and loss of Trichodesmium populations, and 717 possibly lead to enhanced vertical fluxes and export production, as previously demonstrated in 718 PCD-induced laboratory cultures of Trichodesmium (Bar-Zeev et al., 2013). We posit that 719 PCD induced demise, in response to concurrent cellular stressors, and facilitated by concerted 720 gene regulation, is typical in natural Trichodesmium blooms and leads to a high export 721 722 production rather than regeneration and recycling of biomass in the upper photic layers.

723

724

725 Author contributions

IBF, DS, and SB conceived and planned the study. DS, UP, HB, SB, WRH, KB and IBF all
participated in the experimental sampling. DS, UP, WRH, HB, FN, DAR, KB, and IBF
analyzed the samples and resulting data. IBF and DS wrote the manuscript with further
contributions to the manuscript by UP, WRH, SB, and KB.

730

731 Acknowledgments

Funding was obtained for IBF through a collaborative grant from MOST Israel and the High 732 Council for Science and Technology (HCST)-France, and a United States-Israel Binational 733 Science Foundation (BSF) grant (No: 2008048) to IBF and KB. This research was partially 734 735 funded by the Gordon and Betty Moore Foundation through Grant GBMF3789 to KDB. The 736 participation of IBF, DS, UP, and WRH in the VAHINE experiment was supported by the German-Israeli Research Foundation (GIF), project number 1133-13.8/2011 to IBF and 737 WRH, and the metatranscriptome analysis by the EU project MaCuMBA (Marine 738 Microorganisms: Cultivation Methods for Improving their Biotechnological Applications; 739 grant agreement no: 311975) to WRH. Funding for VAHINE Experimental project was 740 741 provided by the Agence Nationale de la Recherche (ANR starting grant VAHINE ANR-13-JS06-0002), INSU-LEFE-CYBER program, GOPS, IRD and M.I.O. The authors thank the 742 captain and crew of the R/V Alis. We acknowledge the SEOH divers service from the IRD 743 research center of Noumea (E. Folcher, B. Bourgeois and A. Renaud) and from the 744 Observatoire Océanologique de Villefranche-sur-mer (OOV, J.M. Grisoni) as well as the 745 technical service of the IRD research center of Noumea for their helpful technical support. 746 Thanks especially to E. Rahav for his assistance throughout the New Caledonia experiment 747 and to H. Elifantz for technical assistance with the 16S sequencing and data analysis. This 748 749 work is in partial fulfillment of the requirements for a PhD thesis for D. Spungin at Bar-Ilan 750 University. We thank the three reviewers whose comments help improve the manuscript 751 substantially

752

753

754

755 References

Allers, E., Niesner, C., Wild, C., and Pernthaler, J.: Microbes enriched in seawater after addition of coral mucus, Applied and Environmental Microbiology, 74, 3274-3278, 2008.

Anders, S., Pyl, P. T., and Huber, W.: HTSeq–A Python framework to work with highthroughput sequencing data, Bioinformatics, btu638, 2014.

- Asplund-Samuelsson, J., Bergman, B., and Larsson, J.: Prokaryotic caspase homologs:
 phylogenetic patterns and functional characteristics reveal considerable diversity, PLOS One,
 7, e49888, 2012.
- Asplund-Samuelsson, J.: The art of destruction: revealing the proteolytic capacity of bacterialcaspase homologs, Molecular Microbiology, 98, 1-6, 2015.
- Azetsu-Scott, K., and Passow, U.: Ascending marine particles: Significance of transparent
 exopolymer particles (TEP) in the upper ocean, Limnology and Oceanography, 49, 741-748,
 2004.
- Bar-Zeev, E., Avishay, I., Bidle, K. D., and Berman-Frank, I.: Programmed cell death in the
 marine cyanobacterium *Trichodesmium* mediates carbon and nitrogen export, The ISME
 Journal, 7, 2340-2348, 2013.
- Behrenfeld, M. J.: Climate-mediated dance of the plankton, Nature Climate Change, 4, 880-887, 2014.
- Bergman, B., Sandh, G., Lin, S., Larsson, J., and Carpenter, E. J.: *Trichodesmium* a
 widespread marine cyanobacterium with unusual nitrogen fixation properties, FEMS
 Microbiology Reviews, 1-17, 10.1111/j.1574-6976.2012.00352.x., 2012.
- Berman-Frank, I., and Dubinsky, Z.: Balanced growth in aquatic plants: Myth or reality?
 Phytoplankton use the imbalance between carbon assimilation and biomass production to their
 strategic advantage, Bioscience, 49, 29-37, 1999.
- Berman-Frank, I., Cullen, J. T., Shaked, Y., Sherrell, R. M., and Falkowski, P. G.: Iron
 availability, cellular iron quotas, and nitrogen fixation in *Trichodesmium*, Limnology and
 Oceanography, 46, 1249-1260, 2001.
- Berman-Frank, I., Bidle, K., Haramaty, L., and Falkowski, P. G.: The demise of the marine
 cyanobacterium, *Trichodesmium* spp., via an autocatalyzed cell death pathway, Limnology
 and Oceanography, 49, 997-1005, 2004.
- Berman-Frank, I., Rosenberg, G., Levitan, O., Haramaty, L., and Mari, X.: Coupling between
 autocatalytic cell death and transparent exopolymeric particle production in the marine
 cyanobacterium *Trichodesmium*, Environmental Microbiology, 9, 1415-1422, 10.1111/j.14622920.2007.01257.x, 2007.
- Berthelot, H., Moutin, T., L'Helguen, S., Leblanc, K., Hélias, S., Grosso, O., Leblond, N.,
 Charrière, B., and Bonnet, S.: Dinitrogen fixation and dissolved organic nitrogen fueled
 primary production and particulate export during the VAHINE mesocosm experiment (New
 Caledonia lagoon), Biogeosciences, 12, 4099-4112, 10.5194/bg-12-4099-2015, 2015.
- Beversdorf, L., White, A., Björkman, K., Letelier, R., and Karl, D.: Phosphonate metabolism
 by *Trichodesmium* IMS101 and the production of greenhouse gases, Limnology and
 Oceanography, 55, 1768-1778, 2010.
- Bidle, K. D., and Falkowski, P. G.: Cell death in planktonic, photosynthetic microorganisms,
 Nature Reviews Microbiology, 2, 643-655, 2004.

Bidle, K. D.: The molecular ecophysiology of programmed cell death in marinephytoplankton, Annual Review Marine Science, 7, 341-375, 2015.

800 Bonnet, S., Berthelot, H., Turk-Kubo, K., Fawcett, S., Rahav, E., l'Helguen, S., and Berman-801 Frank, I.: Dynamics of N_2 fixation and fate of diazotroph-derived nitrogen in a low nutrient 802 low chlorophyll ecosystem: results from the VAHINE mesocosm experiment (New 803 Caledonia), Biogeosciences, 12, 19579-19626, doi:10.5194/bgd-12-19579-2015, 2015.

Bonnet, S., Moutin, T., Rodier, M., Grisoni, J. M., Louis, F., Folcher, E., Bourgeois, B., Boré,
J. M., and Renaud, A.: Introduction to the project VAHINE: Variability of vertical and trophic
transfer of diazotroph derived N in the South West Pacific, Biogeosciences, doi:10.5194/bg2015-615, 2016a.

- Bonnet, S., Berthelot, H., Turk-Kubo, K., Cornet-Barthaux, V., Fawcett, S., Berman-Frank, I.,
 Barani, A., Dekeazemacker, J., Benavides, M., Charrière, B., and Capone, D.: *Trichodesmium*blooms support diatom growth in the Southwest Pacific Ocean, Limnology and
- 811 Oceanography, 2016b. In Press.
- 812

Brown, J. M., LaBarre, B. A., and Hewson, I.: Characterization of *Trichodesmium*-associated
viral communities in the eastern Gulf of Mexico, FEMS Microbiology Ecology, 84, 603-613,
2013.

- Brussaard, C. P. D., Mari, X., Van Bleijswijk, J. D. L., and Veldhuis, M. J. W.: A mesocosm
 study of Phaeocystis globosa (Prymnesiophyceae) population dynamics II. Significance for
 the microbial community, Harmful Algae, 4, 875-893, 2005.
- Brussaard, C. R. D.: Optimization of procedures for counting viruses by flow cytometry, App.
 Environmental Microbiology, 70, 1506-1513, 2003.

Capone, D., Burns, J., Montoya, J., Michaels, A., Subramaniam, A., and Carpenter, E.: New
nitrogen input to the tropical North Atlantic Ocean by nitrogen fixation by the
cyanobacterium, *Trichodesmium* spp, Global Biogeochemical Cycles, 19, 2004.

- Capone, D. G., and Carpenter, E. J.: Nitrogen fixation in the marine environment, Science,
 217, 1140-1142, 1982.
- Capone, D. G., Zehr, J. P., Paerl, H. W., Bergman, B., and Carpenter, E. J.: *Trichodesmium*, a
 globally significant marine cyanobacterium, Science, 276, 1221-1229, 1997.

Capone, D. G., Subramaniam, A., Montoya, J. P., Voss, M., Humborg, C., Johansen, A. M.,
Siefert, R. L., and Carpenter, E. J.: An extensive bloom of the N₂-fixing cyanobacterium *Trichodesmium erythraeum* in the central Arabian Sea, Marine Ecology Progress Series, 172,
281-292, 1998.

- Castruita, M., Saito, M., Schottel, P., Elmegreen, L., Myneni, S., Stiefel, E., and Morel, F. M.:
 Overexpression and characterization of an iron storage and DNA-binding Dps protein from
- *Trichodesmium erythraeum*, Applied and Environmental Microbiology, 72, 2918-2924, 2006.
- Chappell, P. D., and Webb, E. A.: A molecular assessment of the iron stress response in the
 two phylogenetic clades of *Trichodesmium*, Environmental Microbiology, 12, 13-27,
 10.1111/j.1462-2920.2009.02026.x, 2010.

- Coleman, M. L., and Chisholm, S. W.: Ecosystem-specific selection pressures revealed
 through comparative population genomics, Proceedings of the National Academy of Sciences,
 107, 18634-18639, 2010.
- B41 Dandonneau, Y., and Gohin, F.: Meridional and seasonal variations of the sea surface
 chlorophyll concentration in the southwestern tropical Pacific (14 to 32 S, 160 to 175 E), Deep
 Sea Research Part A. Oceanographic Research Papers, 31, 1377-1393, 1984.
- B44 Dowd, S. E., Callaway, T. R., Wolcott, R. D., Sun, Y., McKeehan, T., Hagevoort, R. G., and
 Edrington, T. S.: Evaluation of the bacterial diversity in the feces of cattle using 16S rDNA
 bacterial tag-encoded FLX amplicon pyrosequencing (bTEFAP), BMC microbiology, 8, 125,
 2008.
- Bupouy, C., Benielli-Gary, D., Neveux, J., Dandonneau, Y., and Westberry, T. K.: An
 algorithm for detecting *Trichodesmium* surface blooms in the South Western Tropical Pacific,
 Biogeosciences, 8, 3631-3647, 10.5194/bg-8-3631-2011, 2011.
- Dyhrman, S. T., Chappell, P. D., Haley, S. T., Moffett, J. W., Orchard, E. D., Waterbury, J. B.,
 and Webb, E. A.: Phosphonate utilization by the globally important marine diazotroph *Trichodesmium*, Nature, 439, 68-71, 2006.
- Edgar, R. C.: UPARSE: highly accurate OTU sequences from microbial amplicon reads,
 Nature Methods, 10, 996-998, 2013.
- Engel, A.: The role of transparent exopolymer particles (TEP) in the increase in apparent
 particle stickiness (alpha) during the decline of a diatom bloom, Journal of Plankton Research,
 22, 485-497, 2000.
- Evans, C., Malin, G., Mills, G. P., and Wilson, W. H.: Viral infection of *Emiliania huxleyi*(prymnesiophyceae) leads to elevated production of reactive oxygen species, Journal of
 Phycology, 42, 1040-1047, 2006.
- Frydenborg, B. R., Krediet, C. J., Teplitski, M., and Ritchie, K. B.: Temperature-dependent
 inhibition of opportunistic vibrio pathogens by native coral commensal bacteria, Microbial
 Ecology, 67, 392-401, 2014.
- Georgiou, T., Yu, Y.-T., Ekunwe, S., Buttner, M., Zuurmond, A.-M., Kraal, B., Kleanthous,
 C., and Snyder, L.: Specific peptide-activated proteolytic cleavage of *Escherichia coli*elongation factor Tu, Proceedings of the National Academy of Sciences, 95, 2891-2895, 1998.
- Guo, C., and Tester, P. A.: Toxic effect of the bloom-forming *Trichodesmium* sp.
 (Cyanophyta) to the copepod Acartia tonsa, Natural Toxins, 2, 222-227, 1994.
- Hamady, M., Walker, J. J., Harris, J. K., Gold, N. J., and Knight, R.: Error-correcting
 barcoded primers for pyrosequencing hundreds of samples in multiplex, Nature Methods, 5,
 235-237, 2008.
- Havaux, M., Guedeney, G., Hagemann, M., Yeremenko, N., Matthijs, H. C., and Jeanjean, R.:
 The chlorophyll-binding protein IsiA is inducible by high light and protects the
 cyanobacterium *Synechocystis* PCC6803 from photooxidative stress, FEBS Letters, 579,
 2289-2293, 2005.

- 877 Herbland, A., Le Bouteiller, A., and Raimbault, P.: Size structure of phytoplankton biomass in
- the equatorial Atlantic Ocean, Deep Sea Research Part A. Oceanographic Research Papers, 32,
- 879 819-836, 1985.

Hewson, I., Govil, S. R., Capone, D. G., Carpenter, E. J., and Fuhrman, J. A.: Evidence of
 Trichodesmium viral lysis and potential significance for biogeochemical cycling in the
 oligotrophic ocean, Aquatic Microbial Ecology, 36, 1-8, 2004.

Hewson, I., Poretsky, R. S., Dyhrman, S. T., Zielinski, B., White, A. E., Tripp, H. J., Montoya,
J. P., and Zehr, J. P.: Microbial community gene expression within colonies of the diazotroph, *Trichodesmium*, from the Southwest Pacific Ocean, ISME Journal, 3, 1286-1300,
10.1038/ismej.2009.75, 2009.

Hmelo, L. R., Van Mooy, B. A. S., and Mincer, T. J.: Characterization of bacterial epibionts
on the cyanobacterium *Trichodesmium*, Aquatic Microbial Ecology, 67, 1-U119,
10.3354/ame01571, 2012.

Holmes, R. M., Aminot, A., Kérouel, R., Hooker, B. A., and Peterson, B. J.: A simple and
precise method for measuring ammonium in marine and freshwater ecosystems, Canadian
Journal of Fisheries and Aquatic Sciences, 56, 1801-1808, 10.1139/f99-128, 1999.

Hove-Jensen, B., Zechel, D. L., and Jochimsen, B.: Utilization of Glyphosate as Phosphate
Source: Biochemistry and Genetics of Bacterial Carbon-Phosphorus Lyase, Microbiology and
Molecular Biology Reviews, 78, 176-197, 2014.

Hunt, B. P. V., Bonnet, S., Berthelot, H., Conroy, B. J., Foster, R., and Pagano, M.:
Contribution and pathways of diazotroph derived nitrogen to zooplankton during the VAHINE
mesocosm experiment in the oligotrophic New Caledonia lagoon, Biogeosciences
Discussions, doi:10.5194/bg-2015-614, 2016.

Ivars-Martinez, E., Martin-Cuadrado, A.-B., D'Auria, G., Mira, A., Ferriera, S., Johnson, J.,
 Friedman, R., and Rodriguez-Valera, F.: Comparative genomics of two ecotypes of the marine
 planktonic copiotroph *Alteromonas macleodii* suggests alternative lifestyles associated with
 different kinds of particulate organic matter, The ISME Journal, 2, 1194-1212, 2008.

Jacquet, S., Heldal, M., Iglesias-Rodriguez, D., Larsen, A., Wilson, W., and Bratbak, G.: Flow
cytometric analysis of an *Emiliana huxleyi* bloom terminated by viral infection, Aquatic
Microbial Ecology, 27, 111-124, 2002.

Jacquet, S., Delesalle, B., Torréton, J.-P., and Blanchot, J.: Response of phytoplankton
communities to increased anthropogenic influences (southwestern lagoon, New Caledonia),
Marine Ecology Progress Series, 320, 65-78, 2006.

Jeanjean, R., Zuther, E., Yeremenko, N., Havaux, M., Matthijs, H. C., and Hagemann, M.: A
photosystem 1 *psaFJ*-null mutant of the cyanobacterium *Synechocystis* PCC 6803 expresses
the *isiAB* operon under iron replete conditions, FEBS letters, 549, 52-56, 2003.

Kerbrat, A.-S., Darius, H. T., Pauillac, S., Chinain, M., and Laurent, D.: Detection of
ciguatoxin-like and paralysing toxins in *Trichodesmium* spp. from New Caledonia lagoon,
Marine Pollution Bulletin, 61, 360-366, 2010.

- Kopylova, E., Noé, L., and Touzet, H.: SortMeRNA: fast and accurate filtering of ribosomal
 RNAs in metatranscriptomic data, Bioinformatics, 28, 3211-3217, 2012.
- Langmead, B., and Salzberg, S. L.: Fast gapped-read alignment with Bowtie 2, Nature
 Methods, 9, 357-359, 2012.
- Latifi, A., Jeanjean, R., Lemeille, S., Havaux, M., and Zhang, C.-C.: Iron starvation leads to
 oxidative stress in *Anabaena* sp. strain PCC 7120, Journal of Bacteriology, 187, 6596-6598,
 2005.
- Laudenbach, D. E., and Straus, N. A.: Characterization of a cyanobacterial iron stress-induced
 gene similar to psbC, Journal of Bacteriology, 170, 5018-5026, 1988.
- Leblanc, K., Cornet, V., Caffin, M., Rodier, M., Desnues, A., Berthelot, H., Turk-Kubo, K.,
 and Heliou, J.: Phytoplankton community structure in the VAHINE mesocosm experiment,
 Biogeosciences Discussions., doi:10.5194/bg-2015-605, 2016.
- Lehahn, Y., Koren, I., Schatz, D., Frada, M., Sheyn, U., Boss, E., Efrati, S., Rudich, Y.,
 Trainic, M., and Sharoni, S.: Decoupling physical from biological processes to assess the
 impact of viruses on a mesoscale algal bloom, Current Biology, 24, 2041-2046, 2014.
- Luo, Y.-W., Doney, S., Anderson, L., Benavides, M., Berman-Frank, I., Bode, A., Bonnet, S.,
 Boström, K., Böttjer, D., and Capone, D.: Database of diazotrophs in global ocean:
 abundance, biomass and nitrogen fixation rates, Earth System Science Data, 4, 47-73, 2012.
- Mackey, K. R., Post, A. F., McIlvin, M. R., Cutter, G. A., John, S. G., and Saito, M. A.:
 Divergent responses of Atlantic coastal and oceanic *Synechococcus* to iron limitation,
 Proceedings of the National Academy of Sciences, 112, 9944-9949, 2015.
- Martin, M.: Cutadapt removes adapter sequences from high-throughput sequencing reads,EMBnet. Journal, 17, pp. 10-12, 2011.
- Martinez, A., Tyson, G. W., and DeLong, E. F.: Widespread known and novel phosphonate
 utilization pathways in marine bacteria revealed by functional screening and metagenomic
 analyses, Environmental Microbiology, 12, 222-238, 10.1111/j.1462-2920.2009.02062.x,
 2010.
- Martínez, A., Osburne, M. S., Sharma, A. K., DeLong, E. F., and Chisholm, S. W.: Phosphite
 utilization by the marine picocyanobacterium *Prochlorococcus* MIT9301, Environmental
 Microbiology, 14, 1363-1377, 2012.
- Massana, R., Murray, A. E., Preston, C. M., and DeLong, E. F.: Vertical distribution and
 phylogenetic characterization of marine planktonic Archaea in the Santa Barbara Channel,
 Applied and Environmental Microbiology, 63, 50-56, 1997.
- Michel, K. P., and Pistorius, E. K.: Adaptation of the photosynthetic electron transport chain
 in cyanobacteria to iron deficiency: the function of IdiA and IsiA, Physiologia Plantarum, 120,
 36-50, 2004.
- Mohr, W., Grosskopf, T., Wallace, D. W., and LaRoche, J.: Methodological underestimation
 of oceanic nitrogen fixation rates, PLOS One, 5, e12583, 2010.

- Montoya, J. P., Voss, M., Kahler, P., and Capone, D. G.: A simple, high-precision, highsensitivity tracer assay for N_2 fixation, Applied and Environmental Microbiology, 62, 986-956 993, 1996.
- Mulholland, M. R.: The fate of nitrogen fixed by diazotrophs in the ocean, Biogeosciences, 4,37-51, 2007.
- O'Neil, J. M., and Roman, M. R.: Ingestion of the Cyanobacterium *Trichodesmium* spp by
 Pelagic Harpacticoid Copepods *Macrosetella*, *Miracia* and *Oculostella*, Hydrobiologia, 293,
 235-240, 1994.
- 962 O'Neil, J. M.: The colonial cyanobacterium *Trichodesmium* as a physical and nutritional
 963 substrate for the harpacticoid copepod *Macrosetella gracilis*, Journal of Plankton Research,
 964 20, 43-59, 1998.
- Ohki, K.: A possible role of temperate phage in the regulation of *Trichodesmium* biomass,
 Bulletin de l'institute oceanographique, Monaco, 19, 287-291, 1999.
- 967 Orchard, E., Webb, E., and Dyhrman, S.: Characterization of phosphorus-regulated genes in
 968 *Trichodesmium* spp., The Biological Bulletin, 205, 230-231, 2003.
- Paerl, H. W., Priscu, J. C., and Brawner, D. L.: Immunochemical localization of nitrogenase in marine *Trichodesmium* aggregates: Relationship to N_2 fixation potential, Applied and Environmental Microbiology, 55, 2965-2975, 1989.
- Passow, U., and Alldredge, A. L.: A dye binding assay for the spectrophotometeric
 measurement of transparent exopolymer particles (TEP), Limnology and Oceanography, 40,
 1326-1335, 1995.
- Pfreundt, U., Kopf, M., Belkin, N., Berman-Frank, I., and Hess, W. R.: The primary
 transcriptome of the marine diazotroph *Trichodesmium erythraeum* IMS101, Scientific
 Reports, 4, 2014.
- Pfreundt, U., Van Wambeke, F., Caffin, M., Bonnet, S., and Hess, W. R.: Succession within
 the prokaryotic communities during the VAHINE mesocosms experiment in the New
 Caledonia lagoon, Biogeosciences, 13, 2319-2337, doi:10.5194/bg-13-2319-2016, 2016.
- Pichon, D., Cudennec, B., Huchette, S., Djediat, C., Renault, T., Paillard, C., and AuzouxBordenave, S.: Characterization of abalone *Haliotis tuberculata–Vibrio harveyi* interactions in
 gill primary cultures, Cytotechnology, 65, 759-772, 2013.
- Pinto, F. L., Thapper, A., Sontheim, W., and Lindblad, P.: Analysis of current and alternative
 phenol based RNA extraction methodologies for cyanobacteria, BMC Molecular Biology, 10,
 1, 2009.
- Polyviou, D., Hitchcock, A., Baylay, A. J., Moore, C. M., and Bibby, T. S.: Phosphite
 utilization by the globally important marine diazotroph *Trichodesmium*, Environmental
 Microbiology Reports, 7, 824-830, 2015.
- Pujo-Pay, M., and Raimbault, P.: Improvement of the wet-oxidation procedure for
 simultaneous determination of particulate organic nitrogen and phosphorus collected on filters,
 Marine Ecology-Progress Series, 105, 203–207, 10.3354/meps105203, 1994.

- Quast, C., Pruesse, E., Yilmaz, P., Gerken, J., Schweer, T., Yarza, P., Peplies, J., and
 Glöckner, F. O.: The SILVA ribosomal RNA gene database project: improved data processing
 and web-based tools, Nucleic Acids Research, 41, D590-D596, 10.1093/nar/gks1219, 2013.
- Rahav, E., Herut, B., Levi, A., Mulholland, M., and Berman-Frank, I.: Springtime contribution
 of dinitrogen fixation to primary production across the Mediterranean Sea, Ocean Science, 9,
 489-498, 2013.
- Rodier, M., and Le Borgne, R.: Population dynamics and environmental conditions affecting *Trichodesmium* spp. (filamentous cyanobacteria) blooms in the south-west lagoon of New
 Caledonia, Journal of Experimental Marine Biology and Ecology, 358, 20-32,
 10.1016/j.jembe.2008.01.016, 2008.
- Rodier, M., and Le Borgne, R.: Population and trophic dynamics of *Trichodesmium thiebautii*in the SE lagoon of New Caledonia. Comparison with *T. erythraeum* in the SW lagoon,
 Marine Pollution Bulletin, 61, 349-359, 2010.
- Shaked, Y.: Iron redox dynamics and biogeochemical cycling in the epilimnion of LakeKinneret, PhD thesis, Hebrew University of Jerusalem, 2002.
- Sheridan, C. C., Steinberg, D. K., and Kling, G. W.: The microbial and metazoan community
 associated with colonies of *Trichodesmium* spp.: a quantitative survey, Journal of Plankton
 Research, 24, 913-922, 2002.
- Siddiqui, P. J., Bergman, B., Bjorkman, P. O., and Carpenter, E. J.: Ultrastructural and
 chemical assessment of poly-beta-hydroxybutyric acid in the marine cyanobacterium *Trichodesmium thiebautii*, FEMS Microbiology Letters, 73, 143-148, 1992.
- 1014 Smetacek, V.: Role of sinking in diatom life-history cycles: ecological, evolutionary and 1015 geological significance, Marine Biology, 84, 239-251, 1985.
- Snow, J. T., Polyviou, D., Skipp, P., Chrismas, N. A., Hitchcock, A., Geider, R., Moore, C.
 M., and Bibby, T. S.: Quantifying Integrated Proteomic Responses to Iron Stress in the
 Globally Important Marine Diazotroph *Trichodesmium*, PLOS One, 10, e0142626, 2015.
- Spungin, D., Berman-Frank, I., and Levitan, O.: *Trichodesmium's* strategies to alleviate
 phosphorus limitation in the future acidified oceans, Environmental Microbiology, 16, 19351947, 2014.
- Spungin, D., Rosenberg, G., Bidle, K. D., and Berman-Frank, I.: Metacaspases and bloom
 demise in the marine cyanobacterium *Trichodesmium*, In Prep.
- Strickland, J. D. H., and Parsons, T. R.: A Practical Handbook of Seawater Analysis, FisheriesResearch Board of Canada, Ottawa, 1972.
- Tandeau de Marsac, N., and Houmard, J.: Complementary chromatic adaptation:
 Physiological conditions and action spectra, in: Methods in Enzymology, Academic Press,
 318-328, 1988.
- Tarutani, K., Nagasaki, K., and Yamaguchi, M.: Viral impacts on total abundance and clonal
 composition of the harmful bloom-forming phytoplankton heterosigma akashiwo, Applied and
 Environmental Microbiology, 66, 4916-4920, 2000.

Thamatrakoln, K., Korenovska, O., Niheu, A. K., and Bidle, K. D.: Whole-genome expression
analysis reveals a role for death-related genes in stress acclimation of the diatom *Thalassiosira pseudonana*, Environmental Microbiology, 14, 67-81, 2012.

Turk-Kubo, K., Frank, I., Hogan, M., Desnues, A., Bonnet, S., and Zehr, J.: Diazotroph community succession during the VAHINE mesocosms experiment (New Caledonia Lagoon),
Biogeosciences 12, 7435-7452, doi:10.5194/bg-12-7435-2015, 2015.

Van Mooy, B. A., Hmelo, L. R., Sofen, L. E., Campagna, S. R., May, A. L., Dyhrman, S. T.,
Heithoff, A., Webb, E. A., Momper, L., and Mincer, T. J.: Quorum sensing control of
phosphorus acquisition in *Trichodesmium* consortia, The ISME Journal, 6, 422-429, 2012.

Van Wambeke, F., Pfreundt, U., Barani, A., Berthelot, H., Moutin, T., Rodier, M., Hess, W.
R., and Bonnet, S.: Heterotrophic bacterial production and metabolic balance during the
VAHINE mesocosm experiment in the New Caledonia lagoon, Biogeosciences Discussions,
12, 19861-19900, doi:10.5194/bgd-12-19861-2015, 2015.

1045 Vardi, A., Berman-Frank, I., Rozenberg, T., Hadas, O., Kaplan, A., and Levine, A.:
1046 Programmed cell death of the dinoflagellate *Peridinium gatunense* is mediated by CO₂
1047 limitation and oxidative stress, Current Biology: CB, 9, 1061-1064, 1999.

Vardi, A., Haramaty, L., Van Mooy, B. A., Fredricks, H. F., Kimmance, S. A., Larsen, A., and
Bidle, K. D.: Host–virus dynamics and subcellular controls of cell fate in a natural
coccolithophore population, Proceedings of the National Academy of Sciences, 109, 1932719332, 2012.

Walsby, A. F.: The properties and bouyancy providing role of gas vacuoles in *Trichodesmium*,British Phycological Journal, 13, 103-116, 1978.

Weitz, J. S., and Wilhelm, S. W.: Ocean viruses and their effects on microbial communitiesand biogeochemical cycles, F1000 Biology Reports, 4, 17, 2012.

Wu, Z., Jenkins, B. D., Rynearson, T. A., Dyhrman, S. T., Saito, M. A., Mercier, M., and
Whitney, L. P.: Empirical bayes analysis of sequencing-based transcriptional profiling without
replicates, BMC Bioinformatics, 11, 564, 2010.

Zehr, J. P.: Nitrogen fixation in the Sea: Why Only *Trichodesmium*, in: Molecular Ecology of
Aquatic Microbes, edited by: Joint, I., NATO ASI Series, Springer-Verlag, Heidelberg, 335363, 1995.

1062

1063

1064 **Figure legends**

Figure 1. Temporal dynamics of pre-bloom measurements in the lagoon waters (a) Chl *a* concentrations (μ g L⁻¹), (b) Virus like particles (VLP, mL⁻¹ x 10⁶), (c) N₂ fixation rates (nmol L⁻¹ h⁻¹) and particulate organic nitrogen (PON, μ mol L⁻¹). (d) Changes in the concentrations of transparent exopolymeric particles (TEP, $\mu g \text{ GX } L^{-1}$) and particulate organic carbon (POC, µmol L⁻¹). Water was sampled from in the lagoon outside the VAHINE mesocosms, at 1 m depth (surface) throughout the experimental period from day 2 to 23 (n=3). For VLP, the standard error for technical replicates (n=3) was < 1 %, which is smaller than symbol size.

1072 Figure 2. (a-c) Dense surface blooms of Trichodesmium observed outside the mesocosms in 1073 the lagoon waters on day 23 at 12:00. Photos illustrate the spatial heterogeneity of the surface 1074 accumulations and the high density of the biomass. (d-e) To examine the mechanistic of demise (Experiment 1), Trichodesmium filaments and colonies were collected by plankton net 1075 1076 (mesh size, 80 μ m) from the dense surface bloom (day 23, 12:00 h; designated T₀) and 1077 resuspended in 0.2 µm pore-size filtered seawater (FSW) in six 4.5 L bottles. Bottles were incubated on-deck in running-seawater pools with ambient surface temperature (~ 26 °C) at 50 1078 % of the surface irradiance. Bottles were sampled every 2-4 h for different parameters until 1079 1080 the biomass crashed. (f) Temporal changes in Chl a concentrations in the bottles from the time of biomass collection and resuspension in the bottles until the Trichodesmium biomass crashed 1081 1082 ~ 24 h after the experiment began (n=3-6). Photo c. courtesy of A. Renaud.

1083 Figure 3. Dynamics of microbial community abundance and diversity during Trichodesmium surface bloom as obtained by 16S rRNA gene sequencing for samples collected from the 1084 surface waters outside the mesocosms during *Trichodesmium* surface accumulation (bloom) 1085 (short-term experiment 1). Pie charts show the changes in dominant groups during the 1086 Trichodesmium bloom and crash from two replicate incubation bottles (please note, 1087 1088 Oscillatoriales consisted only of Trichodesmium in this experiment). The graphs below show the respective temporal dynamics of Trichodesmium (gray circles) and Alteromonas (white 1089 1090 triangles), the dominant bacterial species during the incubation expreriment.

Figure 4. Short-term experiment 2 - measurements from the lagoon waters following 1091 Trichodemsium bloom on day 23. (a) Virus like particles (VLP, $mL^{-1} \times 10^6$) and 1092 Trichodesmium abundance (trichomes L⁻¹) derived from qPCR-based abundances of 1093 Trichodesmium nifH gene copies (Bonnet et al. 2016b) based on the assumption of 100 gene-1094 copies per trichome (b) N₂ fixation rates (nmol L⁻¹ h⁻¹), particulate organic nitrogen (PON, 1095 μ mol L⁻¹) and ammonium concentrations (NH₄⁺, μ mol L⁻¹). (c) Changes in the concentrations 1096 of transparent exopolymeric particles (TEP, µg GX L⁻¹) and particulate organic carbon (POC, 1097 μ mol L⁻¹). For experiment 2, seawater from the surface bloom was collected 5 h after the 1098 1099 initial surface bloom was sighted (day 23, 17:00) by directly filling 20 L polyethylene carboys

1100 gently to avoid destroying biomass. Bottles were placed in on-deck incubators filled with 1101 running seawater to maintain ambient surface temperature (~ 26 °C) and covered with neutral 1102 screening at 50 % surface irradiance levels. For all parameters, replicates were n=3. For VLP, 1103 the standard error for technical replicates (n=3) was < 1 %, which is smaller than symbol size.

1104 Figure 5. (a) Expression of alkaline phosphatase associated genes *phoA* and *phoX* (Tery_3467 and Tery 3845), phosphite utilization genes *ptxA*, *ptxB* and *ptxC* (Tery 0365- Tery 0367), 1105 and phosphonate utilization genes (phn genes, Tery_4993, Tery_4994, Tery_4995, 1106 Tery_4996*, Tery_4997, Tery_4998, Tery_4999, Tery_5000, Tery_5001 Tery_5002 and 1107 1108 Tery_5003). Asterisks near locus tag numbers indicate gene duplicates. (b) Iron-related genes, isiB (Tery_1666), isiA (Tery_1667), idiA (Tery_3377), and ferritin DPS gene dpsA 1109 (Tery_4282). Bars represent log2 fold changes of corresponding genes at T_8 (8 hours after T_0) 1110 and T_{22} (22 hours after T_0) in comparison to T_0 . Significant expression was tested with ASC 1111 1112 (Wu et al., 2010) and marked with an asterisk. Black asterisks represent significant change from T_0 . A gene was called differentially expressed if P > 0.98 (posterior probability). 1113

Figure 6. (a) Dynamics of caspase-specific activity rates (pmol L^{-1} min⁻¹) of *Trichodesmium* 1114 1115 in the New Caledonian lagoon during bloom accumulation and bloom demise, sampled during experiment 1. Samples (n=6) collected from the bloom (day 23, 12:00 T₀), were incubated on-1116 1117 deck in an incubator fitted with running seawater to maintain ambient surface temperature (~ 26 °C). (b) Transcript accumulation of metacaspase genes in the Trichodesmium bloom during 1118 the short-term incubation experiment. Metacaspase genes are TeMC1 (Tery_2077), TeMC2 1119 (Tery_2689), TeMC3 (Tery_3869), TeMC4 (Tery_2471), TeMC5 (Tery_2760), TeMC6 1120 (Tery_2058), TeMC7 (Tery_1841), TeMC8 (Tery_0382), TeMC9 (Tery_4625), TeMC10 1121 (Tery_2624), TeMC11 (Tery_2158) and TeMC12 (Tery_2963). Bars represent log2 fold 1122 1123 changes at T_8 (8 hours after T_0) and T_{22} (22 hours since T_0) in comparison to T_0 . Significant expression was tested with ASC (Wu et al., 2010) and marked with an asterisk. Black asterisks 1124 1125 represent significant change from T_0 . A gene was called differentially expressed if P > 0.981126 (posterior probability).

Figure 7. Change in gas vesicle protein (*gvp*) genes as obtained from metatranscriptomic analyses of the *Trichodesmium* bloom from peak to collapse (experiment 1). *gvpA* genes (Tery_2330 and Tery_2335*) encode the main constituent of the gas vesicles that forms the essential core of the structure; *gvpN* (Tery_2329 and Tery_2334) *gvpK* (Tery_2322), *gvpG* (Tery_2338) and *gvpL/gvpF* (Tery_2339 and Tery_2340*) encode vesicle synthesis proteins. 1132 Bars represent log2 fold changes at T_8 (8 hours after T_0) and T_{22} (22 hours since T_0) in 1133 comparison to T_0 . Significant expression was tested with ASC (Wu et al., 2010) and marked 1134 with an asterisk. Black asterisks represent significant change from T_0 . A gene was called 1135 differentially expressed if P > 0.98 (posterior probability).

- 115/





1163		
1164		
1165		
1166		
1167		
1168		
1169		
1170		
1171		
1172		
1173		
1174		
1175		
1176		
1177		

1180 Figure 3











1	2	n	2
т	Z	υ	3







Supplementary information

Figure S1: Time line of sampling during pre-bloom days (yellow circles, days 1-23), and during experiment 1 (blue circles, days 23-24) and experiment 2 (green circles, days 23-26). For experiment 1, *Trichodesmium* filaments and colonies were collected from the dense surface bloom using a plankton net (mesh size, 80 μ m) from surface water (day 23, 12:00 h; designated T₀). *Trichodesmium* was resuspended in filtered seawater (FSW) that was split to six 4.5 L bottles that were incubate (see below) and sampled every 2-4 h until the biomass crashed. For experiment 2, seawater from the surface bloom was collected 5 h after the initial surface bloom was sighted (day 23, 17:00) by directly filling 20 L polyethylene carboys using a Teflon® PFA pump. Carboys were sampled every 4 h during the first 23 hours and then after 42 h and 72 h since the beginning of the experiment. Bottles and carboys from both experiments were placed in on-deck incubators filled with running seawater to maintain ambient surface temperature (~ 26 °C) and covered with neutral screening at 50 % surface irradiance levels.

Figure S1



Table S1. Metatransciptomic changes of selected marker genes in samples from *Trichodesmium* bloom and crash. Metatranscriptomic samples were obtained from the *Trichodesmium* bloom on day 23 at three successive time points [(T_0 , T_8 and T_{22} , representing 0, 8, and 22 h from surface bloom]. Table data provides the respective locus_tag in the *T. erythraeum* genome, gene name and annotation, expected fold-change calculated from the results of ASC (see methods section), log2 fold change, posterior probabilities for these fold-changes as calculated by ASC (significant if P > 0.98, in bold), and the reads per million reads (RPM) for each time point.

				Fold-change (from ASC)		log2(Fold-change)			Posterior probabilities (P) from ASC						Reads per Million reads			
Locus_tag	Symbol	Annotation	(T8/T0)	(T22/T8)	(T22/T0)	(T8/T0)	(T22/T8)	(T22/T0)	P(T0/T8>2)	P(T8/T0>2)	P(T8/T22>2)	P(T22/T8>2)	P(T22/T0>2)	P(T0/T22>2)	Т0	Т8	T22	
	Tery_2322	gvpA	Gas vesicle synthesis protein- GvpA	0.53	NA	0.27	-0.92	NA	-1.88	0.43	0.00	NA	NA	0.00	0.81	8	0	0
Tery_23	Tery_2323	gvpA	Gas vesicle synthesis protein- GvpA	0.41	NA	0.20	-1.29	NA	-2.32	0.73	0.00	NA	NA	0.00	0.93	15	0	0
	Tery_2324	gvpA	Gas vesicle synthesis protein- GvpA	0.12	4.09	0.48	-3.07	2.03	-1.06	1.00	0.00	0.00	1.00	0.00	0.97	5042	603	3009
	Tery_2325	gvpA	Gas vesicle synthesis protein- GvpA	0.12	2.12	0.26	-3.01	1.08	-1.94	1.00	0.00	0.00	0.79	0.00	1.00	4022	497	1285
Gas Vesicle	Tery_2329	gvpN	Gas vesicle protein- GvpN	0.58	0.45	0.26	-0.78	-1.14	-1.93	0.00	0.00	0.91	0.00	0.00	1.00	1964	1198	653
Protein (Gyn)	Tery_2330	gvpA	Gas vesicle synthesis protein- GvpA	0.08	2.54	0.13	-3.65	1.35	-2.90	1.00	0.00	0.00	0.78	0.00	1.00	444	19	71
formation	Tery_2332	gvpK	Gas vesicle protein- GvpK	0.21	0.70	0.15	-2.22	-0.52	-2.78	1.00	0.00	0.01	0.00	0.00	1.00	1408	306	257
Tery_2334 Tery_2335	Tery_2334	gvpN	Gas vesicle protein- GvpN	1.37	0.04	0.05	0.46	-4.57	-4.45	0.00	0.00	1.00	0.00	0.00	1.00	464	674	19
	Tery_2335	gvpA	Gas vesicle synthesis protein- GvpA	0.67	0.41	0.27	-0.58	-1.27	-1.88	0.00	0.00	0.90	0.00	0.00	1.00	479	335	164
	Tery_2338	gvpG	Gas vesicle protein- GvpG	1.11	0.26	0.28	0.15	-1.95	-1.84	0.00	0.00	1.00	0.00	0.00	1.00	431	505	152
	Tery_2339	gvpL/gvpF	Gas vesicle synthesis protein- GvpLGvpF	2.05	0.33	0.67	1.04	-1.01	-0.58	0.00	0.85	1.00	0.00	0.00	0.00	1608	3473	1371
	Tery_2340	gvp∟/gvp⊢	Gas vesicle synthesis protein- GvpLGvpF	0.24	2.17	0.39	-2.08	1.12	-1.30	1.00	0.00	0.00	0.59	0.00	0.85	102	15	50
Phosphite	Tery_0365	ptxA	Phosphonate transport	3.29	1.18	4.59	1.72	0.24	2.20	0.00	1.00	0.00	0.01	1.00	0.00	16	12	105
Utalization genes	Tery_0366	ptxB	Phosphonate transport	1.99	0.27	0.53	0.99	-1.88	-0.92	0.00	0.48	1.00	0.00	0.00	0.36	193	412	130
J J	Tery 0367	ptxC	Phosphonate transport	0.62	1.93	1.18	-0.69	0.95	0.24	0.06	0.00	0.00	0.42	0.00	0.00	126	79	192
	Tery 4993	phhD	Phosphonate transport	0.87	7.00	7.71	-0.20	2.08	2.95	0.00	0.01	0.00	1.00	0.23	0.00	34	30	344
	Tery 4005	philo	Dheenhonate transport	1 70	1.30	0.24	0.20	2.00	2.00	0.00	0.00	0.00	1.00	1.00	0.00	25	54	210
	Tery 4995	phnE	Phosphonate transport	5.18	0.40	9.34 2.19	2.37	-1.31	1.13	0.00	1.00	0.00	0.00	0.67	0.00	25	166	77
Phosphonate	Tery_4997	phnG	Membrane associated C-P lyase	0.39	2.97	0.68	-1.37	1.57	-0.55	0.79	0.00	0.00	0.75	0.00	0.22	17	0	15
transporters and	Tery_4998	phnH	Phosphonate metabolism	1.16	2.70	4.04	0.21	1.43	2.01	0.00	0.03	0.00	0.82	0.99	0.00	10	14	59
metabolism	Tery_4999	phnl	Phosphonate metabolism	2.06	1.77	4.58	1.05	0.82	2.20	0.00	0.56	0.00	0.33	1.00	0.00	12	35	80
	Tery_5000	phnJ	Membrane associated C-P lyase	2.93	0.91	3.24	1.55	-0.13	1.70	0.00	0.96	0.02	0.00	0.95	0.00	11	47	53
	Tery_5001	phnK	C-P lyase system	2.50	0.97	2.92	1.32	-0.05	1.54	0.00	0.83	0.02	0.01	0.89	0.00	11	39	46
	Tery_5002	phnL	C-P lyase system	3.36	2.09	9.25	1.75	1.06	3.21	0.00	0.99	0.00	0.58	1.00	0.00	11	55	149
	Tery_5003	phnM	Membrane associated C-P lyase	0.84	3.94	3.87	-0.25	1.98	1.95	0.04	0.00	0.00	0.98	0.99	0.00	13	11	74
	Tery_3534	sphX	Phosphate bindig protein (regulated by P supply)	2.57	0.57	1.47	1.36	-0.82	0.55	0.00	0.99	0.22	0.00	0.02	0.00	71	202	136
Inorganic P	Tery_3537	pstS	Phosphate bindig protein	0.39	4.97	1.77	-1.34	2.31	0.83	0.87	0.00	0.00	1.00	0.23	0.00	62	19	146
transporters	Tery_3539	pstS	Phosphate transport system permease protein 2	0.68	1.01	0.66	-0.56	0.01	-0.61	0.04	0.00	0.01	0.00	0.00	0.12	69	47	59
aanoportoro	Tery_3540	pstB	Inorganic P transporter	0.57	0.69	0.38	-0.80	-0.54	-1.40	0.12	0.00	0.08	0.00	0.00	0.92	162	94	77
	Tery_3583	pstC	Inorganic P transporter	2.17	1.34	3.25	1.12	0.42	1.70	0.00	0.68	0.00	0.03	0.99	0.00	25	66	111
Degradation of	Tery_3467	phoA	Aikaiine phosphalase	1.47	2.58	3.80	0.55	1.37	1.95	0.00	0.00	0.00	1.00	1.00	0.00	179	279	660
organic P sources	Tery_3845	phoX	Alkaline phosphatase	0.79	0.87	0.68	-0.34	-0.20	-0.55	0.00	0.00	0.00	0.00	0.00	0.01	223	184	195
	Tery_1560	tonB	TonB family protein	2.38	0.37	0.88	1.25	-1.44	-0.19	0.00	1.00	1.00	0.00	0.00	0.00	979	2459	1096
	1 ery_1666	fldA	Flavodoxin FldA	0.39	1.02	0.40	-1.34	0.02	-1.33	1.00	0.00	0.00	0.00	0.00	1.00	1535	629	777
	Tery_1667	ISIA	Iron-stress induced protein A, photosystem antenna protein-like, IsiA	1.72	1.15	2.06	0.78	0.20	1.04	0.00	0.13	0.00	0.00	0.57	0.00	55	105	149
_	Tery_1953	fur	Ferric uptake regulator	1.81	0.86	1.67	0.86	-0.22	0.74	0.00	0.31	0.03	0.00	0.27	0.00	20	44 50	46
iricnodesmium	Tery_1950	flavodovin	Elavodovia	0.42	2.07	0.09	-1.20	-1.94	1 20	1.00	0.00	0.95	0.00	0.00	0.00	541	60	9
Fe acquisition,	Tery 2787	dns	Ferritin and Dns	2.12	0.24	0.30	1.09	-2.06	-1.30	0.00	0.00	0.00	0.00	0.00	0.39	7	25	203
storage,	Tery 2878	feoB	ferrous iron transport protein B	1.09	1,121	1.23	0.12	0.17	0.30	0.00	0.00	0.00	0.00	0,00	0.00	81	94	130
regulation ans	Terv 2879	feoA	ferrous iron transport protein A	0.38	NA	0.19	-1.38	NA	-2.43	0.79	0.00	NA	NA	0.00	0.95	17	0	0
quota reduction	Tery_3222	futC	iron(III) ABC transporter, ATP-binding protein	1.08	1.79	1.96	0.11	0.84	0.97	0.00	0.00	0.00	0.18	0.41	0.00	152	174	384
genes (Gono lict	Tery_3223	futB	iron(III) ABC transporter, permease protein	2.70	1.70	5.02	1.43	0.77	2.33	0.00	0.99	0.00	0.15	1.00	0.00	38	120	254
from Channell	Tery_3377	idiA	idiA	3.27	0.37	1.23	1.71	-1.42	0.29	0.00	1.00	1.00	0.00	0.00	0.00	413	1434	647
from Chappen	Tery_3404	fur	Ferric uptake regulator	0.28	1.01	0.26	-1.82	0.02	-1.94	1.00	0.00	0.00	0.00	0.00	1.00	221	57	71
and Webb 2009)	Tery_3943	Periplasmic binding protein	Periplasmic binding protein	0.82	0.88	0.71	-0.29	-0.18	-0.50	0.00	0.00	0.01	0.00	0.00	0.04	88	75	80
	Tery_4282	dpsA	Ferritin and Dps	1.21	0.22	0.26	0.27	-2.20	-1.96	0.00	0.00	1.00	0.00	0.00	1.00	491	625	158
	Tery 4448	motA/exbB	MotA/TolQ/ExbB proton channel	0.70	1.89	1.31	-0.51	0.92	0.39	0.03	0.00	0.00	0.40	0.01	0.00	57	40	99
Tery_4	Tery_4449	exbD	Biopolymer transport protein ExbD/ToIR	2.02	1.37	3.39	1.02	0.45	1.76	0.00	0.52	0.00	0.09	0.97	0.00	12	33	59
Tery_0383 Tery_184 Tery_2963 Tery_2053 Tery_2054 Tery_2054	Tery_0382	TeMC8	Putative Chase2 sensor protein, metacaspase	3.25	1.04	3.49	1.70	0.06	1.81	0.00	1.00	0.00	0.00	1.00	0.00	90	322	409
	Tery_1841	TeMC7	Peptidase C14,C caspase catalytic subunit p20, metacaspase	2.60	1.11	2.92	1.38	0.16	1.54	0.00	1.00	0.00	0.00	1.00	0.00	442	1219	1647
	Tery_2963	TeMC12	Hypothetical protein	0.64	NA	0.41	-0.63	NA	-1.30	0.25	0.00	NA	NA	0.01	0.63	5	0	0
	Tery_2058	TeMC6	Peptidase C14,C caspase catalytic subunit p20, metacaspase	2.91	0.44	1.28	1.54	-1.19	0.35	0.00	1.00	0.76	0.00	0.01	0.00	61	199	102
	1 ery_2077	TeMC1	Hypothetical protein, metacaspase	5.31	1.08	5.91	2.41	0.11	2.56	0.00	1.00	0.00	0.00	1.00	0.00	129	747	981
	Tery_2158	TeMC11	Hypotnetical+protein	2.60	1.39	3.89	1.38	0.47	1.96	0.00	0.98	0.00	0.01	1.00	0.00	41	124	214
	Tery_2471	TeMC4	Pepudase C14,C caspase catalytic subunit p20, metacaspase	2.38	1./8	4.30	1.25	0.83	2.10	0.00	1.00	0.00	0.04	1.00	0.00	266	6/5	1461
	Tery_2624		Hupothotical protoin metacophone	1.40	2.07	3.19	0.00	1.05	1.0/	0.00	0.01	0.00	0.00	1.00	0.00	58 119	94	245 247
Tery_2689 Tery_2760 Tery_3869	Tery 2760		Pentidase C14 C caspase catalytic subunit n20 metacaspase	1.98	1.12	2.21	0.99	0.17	1.18	0.00	0.40	0.00	0.00	0.88	0.00	118 <u>4</u> 0	203	180
	Tery 3869	TeMC3	WD-40 repeat containing protein metacaspase	2 79	2 21	6.24	1 48	1 14	2.64	0.00	1.00	0.00	0.97	1.00	0.00	300	889	2390
	Tery 4625	TeMC9	WD-40 repeat-containing protein, metacaspase	2.82	1.54	4.36	1.50	0.62	2.13	0.00	1.00	0.00	0.00	1.00	0.00	507	1516	2826
			g retter, metaeopuee														•	