

1 **Competition between a toxic and a non-toxic *Microcystis* strain**
2 **under constant and pulsed nitrogen and phosphorus supply**

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17 **Running head:** Genotype dynamics in experimental *Microcystis* populations

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19 *Microcystis aeuruginosa*, plankton dynamics, resource pulses, quantitative PCR

20 Abstract

21 The toxicity of a harmful algal bloom is strongly determined by the relative abundance of
22 non-toxic and toxic genotypes, and might therefore be regulated by competition for growth-
23 limiting resources. Here, we studied how the toxic *Microcystis aeruginosa* strain PCC 7806
24 and a non-toxic mutant compete for nitrogen and phosphorus under constant and pulsed
25 nutrient supply. Our monoculture and competition experiments show that these closely
26 related genotypes have distinct nutrient physiologies and that they differ in their ability to
27 compete for nitrogen and phosphorus. The toxic wild type won the competition under
28 nitrogen limitation, while the non-toxic mutant dominated under phosphorus limitation.
29 Pulses of both nitrogen and phosphorus increased the dominance of the toxic genotype, which
30 lead to an even faster competitive exclusion of the non-toxic genotype under nitrogen pulses
31 and to coexistence of both genotypes under phosphorus pulses. Our findings indicate that the
32 genotype level dynamics driven by resource competition can be an important factor in
33 determining cyanobacterial bloom toxicity.

34

35 **Introduction**

36 The harmful blooms of microalgae and cyanobacteria (harmful algal blooms, HABs) that
37 often form in eutrophic freshwater and estuaries are among the most eye-catching and
38 problematic microbial populations found in nature (Paerl and Otten, 2013; Brooks et al.,
39 2015). These microbial mass occurrences are of great societal relevance because they
40 threaten the production of drinking water and the recreational use of water bodies. It has been
41 predicted that global change will further increase the risk and intensity of HABs due to
42 human-induced eutrophication (O'Neil et al., 2012; Mantzouki et al., 2015), global warming
43 (Paerl and Huisman, 2008, 2009), increased water column stratification (Jöhnk et al., 2008),
44 and rising CO₂ levels (Verspagen et al., 2014).

45 Although HABs are often dominated by a single species, they harbour substantial
46 genetic diversity and can vary considerably in toxicity. It has been shown that the toxicity of
47 a bloom depends to a large extent on the relative proportions of toxic versus non-toxic or less
48 toxic strains or genotypes of the same species (e.g. Park et al., 1998; Chorus et al., 2001;
49 Briand et al., 2002; Wiedner et al. 2002; Welker et al., 2003; Huisman et al., 2005). The
50 difference in toxin production level between genotypes can cover several orders of magnitude
51 and exceeds by far the within-genotype variability in toxin production caused by
52 environmental factors (Sivonen and Jones, 1999; Rohrlack et al., 2001; Carrillo et al., 2003).
53 Several studies have indicated that the temporal and spatial distribution of toxic and non-
54 toxic genotypes within natural blooms could be explained as the result of resource
55 competition. For instance, Kardinaal et al. (2007) demonstrated in laboratory experiments
56 that a toxic genotype of the harmful cyanobacterium *Microcystis aeruginosa* was displaced
57 by a non-toxic type under light limitation. The authors suggested that the transition from
58 toxic to non-toxic *Microcystis* genotypes during the duration of a bloom could be explained
59 by the development of high biomass concentrations toward the end of the bloom, which

60 increases self-shading and enhances competition for light. Furthermore, van de Waal et al.
61 (2011) demonstrated that the same non-toxic strain of *Microcystis* outcompeted the toxic
62 strain in CO₂-limited conditions. Yet, HABs are also known to be affected by limitation of
63 nutrients, in particular, nitrogen and phosphorus. If toxic and non-toxic genotypes differ in
64 their competitive ability for these two nutrients, the type of nutrient limitation that arises
65 might strongly influence the toxicity of a bloom. Nitrogen and phosphorus limitation may
66 occur towards the end of blooms, but these have also been observed during earlier phases
67 (Moisander et al., 2003; Walve and Larsson, 2007; Davis et al., 2009). Earlier periods of
68 nutrient limitation may be interrupted by pulses of nutrient supply, for example, when wind-
69 induced mixing of the water column enhances the diffusive inflow of nutrients from deeper
70 water layers (Kononen et al., 1996; Stal et al., 1999; Davis et al., 2009). In fact, it has been
71 argued that in natural systems such fluctuations in nutrient availability are the rule rather than
72 the exception (e.g. Grover, 1988, 1991; Padisák, et al., 1993; Sommer 2002). Thus, if toxic
73 and non-toxic genotypes also differ in their abilities to exploit nutrient pulses, the intensity of
74 nutrient fluctuations might represent an additional important factor that influences how the
75 toxin content of the bloom develops.

76 In this study, we used controlled microcosm experiments to investigate the
77 competition between a toxic and a non-toxic strain (hereafter interchangeably also referred to
78 as genotypes) of the harmful cyanobacterium *Microcystis aeruginosa* for nitrogen and
79 phosphorus under both constant and pulsed supply. First, we measured several growth
80 parameters for each genotype in replicated monocultures. Second, we investigated the
81 competition between both genotypes under constant nutrient supply and under different kinds
82 of nutrient pulses in replicated continuous and semi-continuous cultures. We also applied a
83 parameterized Droop model to study the competition for phosphorus under constant and
84 pulsed conditions in more detail with the help of numerical computer simulations.

85 **Methods**

86 ***Microcystis strains and culture maintenance:***

87 We investigated the competition between two axenic strains (hereafter genotypes) of *M.*
88 *aeruginosa*: the toxic (microcystin-producing) wild type (WT) PCC 7806, obtained from the
89 Pasteur Culture Collection in Paris, and the non-toxic mutant genotype (M), provided by
90 Prof. E. Dittmann. The mutant genotype was originally generated from the PCC 7806 by
91 inserting a chloramphenicol resistance cartridge into the *mcyB* gene involved in microcystin
92 biosynthesis (Dittmann et al., 1997). Axenic pre-cultures of both genotypes were grown at 25
93 $\pm 1^\circ\text{C}$ in a 250 mL Erlenmeyer flask containing 40 mL of Z8 growth medium (Kotai, 1972).
94 Z8 growth medium was prepared with analytical grade water (ELIX[®], Merck Millipore,
95 Billerica, MA, USA). Stock cultures of the mutant genotype also contained 5 μg
96 chloramphenicol mL^{-1} (Dittmann et al., 1997).

97

98 ***Total Microcystis cell densities:***

99 The *Microcystis* populations in our study exhibited a linear relationship between cell density
100 and optical density (OD), as shown previously (Briand et al., 2012). We therefore established
101 a standard conversion formula to estimate cell density as a function of OD. To achieve this,
102 cell numbers of live samples from a chemostat culture at steady state were counted with a
103 hemacytometer counting chamber using a compound microscope (Zeiss Axioskop 2 plus,
104 Oberkochen, Germany). OD was measured at the wavelength 750 nm (UV-1800
105 spectrophotometer, Shimadzu, Japan). In the range of OD values that we observed in our
106 experiments (0.01–0.25), the cell density of *Microcystis* (cells mL^{-1}) could be reliably
107 estimated according to the following linear regression equation: $\text{density} (\text{cells mL}^{-1}) = \text{OD}_{750}$
108 $\times 4 \times 10^7$ ($N = 7$; $R^2 = 0.949$).

109

110 ***Relative genotype proportions:***

111 We sampled our microcosms every 1-3 days, depending on the experiment, with sterile
112 syringe (chemostats) or pipette (batch cultures). Note that not all the samples were analyzed
113 in the end. We used quantitative real-time PCR (qPCR) for quantifying genotype proportions
114 in our experimental samples. We followed previously published general protocols for the
115 *Microcystis* system (Briand et al. 2009, 2012; Kurmayer and Kutzenberger, 2003). In our
116 study, DNA was extracted using the DNeasy Plant Mini Kit (Qiagen). Extractions were
117 diluted 10-fold for qPCR, which was used to estimate the proportion of the mutant genotype.
118 All cells in the culture were quantified with previously described primers and a probe for an
119 intergenic spacer in the phycocyanin (PC) gene (Briand et al., 2009, 2012; Kurmayer and
120 Kutzenberger, 2003). The primers and the probe used to quantify the mutant strain targeted
121 the inserted chloramphenicol resistance cassette in the *mcyB* gene (Briand et al., 2012). The
122 probes were labeled with the fluorescent reporter dye FAM (6-carboxyfluorescein) at the 5'
123 end and the amidite Black Hole Quencher-1 (BHQ-1) at the 3' end. qPCR was performed with
124 Applied Biosystems 7300 RealTime PCR Systems (Life Technologies). The PCR reactions
125 contained 10 μ l of the PerfeCta FastMix[®] II supermix with ROX[™] (Quanta Biosciences), 200
126 nM of forward and reverse primers, 100 nM of probe and 5 μ l of template DNA in a total
127 volume of 20 μ l. The PCR was performed with a 2-step PCR program. The PCR program
128 included initial denaturation for 3 min at 95 °C, followed by 40 cycles of 15 s at 95 °C and 1
129 minute at 60 °C. All PCR reactions were performed in triplicates, including negative
130 reactions not containing DNA template and a positive control containing 50% mutant cells,
131 which was used to normalize results. The fluorescence threshold value was manually set to
132 0.15 in all qPCR runs while baseline values were determined automatically. The proportion
133 of mutant strain related to all cells was determined using the Δ Ct method first developed by
134 Briand et al. (2008) for *Planktothrix agardhii* and subsequently used for *M. aeruginosa*

135 (Briand et al., 2009, 2012; Sabart et al., 2010). Further details of the qPCR protocol can be
136 found from the supplementary information file.

137

138 ***Monoculture parameters:***

139 As a first step, we used monoculture experiments to determine several growth parameters for
140 each genotype. These parameters allow to make predictions about the outcome of
141 competition in mixed culture, which can then be compared to the results from the competition
142 experiments. The monoculture parameters will also be used to parameterize a variable-
143 internal-stores model of competition.

144

145 *Realized maximum specific growth rate:* For each genotype, the maximum specific growth
146 rate, μ_{max} , was determined in semi-continuous cultures ($n = 6$) on N-limited and P-limited
147 growth medium, respectively. Cultures of 100 mL were inoculated with very low cell density
148 and grown in 250 mL Erlenmeyer flasks for three days. Every day, 10% of the culture was
149 replaced with fresh growth medium and the cell density, N_t , at time point t was determined.
150 The maximum specific growth rates in N-limited and P-limited growth medium, $\mu_{max,N}$ and
151 $\mu_{max,P}$, were calculated as the mean of the two growth rates calculated between the two time
152 intervals as $\ln(N_{t+1}/N_t)/(t) + d$, where d represents the dilution rate of 0.1 day^{-1} .

153

154 *Critical nutrient concentrations:* In order to measure the critical resource concentration for
155 nitrogen, R_N^* , and phosphorus, R_P^* , each genotype was grown separately in chemostats ($n =$
156 3) for approximately three weeks until it reached a steady state. The R_N^* and R_P^* values of
157 each genotype were then determined as the residual nutrient concentration in the nitrogen-
158 and phosphorus-limited steady state, respectively. Nutrient analysis was based on triplicate
159 samples of 50 mL that were taken from the steady state cultures and centrifuged at $8000 \times g$

160 and 4°C for 5 minutes in order to remove *Microcystis* cells. The supernatant was then filtered
161 through a 0.22 µm filter with *Supor*[®] membrane (Merck Millipore, Billerica, MA, USA) and
162 stored at –20 °C. Analyses of N and P were conducted at Lammi Biological Station
163 (University of Helsinki) with the following methodology: total dissolved nitrogen and
164 phosphorus were analyzed by flow injection analysis (FIA) colorimetry (Quikchem8000,
165 Lachat Instruments, Hach Company, Loveland, CO, USA) using the alkaline persulfate
166 oxidation method (QuikChem[®]METHOD 10-107-04-1-I, Lachat Instruments, Hach
167 Company, Loveland, CO, USA) for nitrogen concentrations and the alkaline persulfate
168 digestion method (QuikChem[®]METHOD 10-115-01-1-S, Lachat Instruments, Hach
169 Company, Loveland, CO, USA) for phosphorus concentrations, respectively.

170

171 *Minimum nutrient quota:* The minimum quotas for nitrogen and phosphorus, $Q_{min,N}$ and
172 $Q_{min,P}$, of each genotype was determined as the intracellular nutrient concentration measured
173 from cells obtained from each chemostat on two separate days, one week apart, when the
174 culture had reached steady state. 50 mL of culture was collected and centrifuged at 10 000 ×
175 g and 4 °C for 5 minutes. The supernatant was removed from the cells and 3 mL of type-2
176 analytical grade water (ELIX[®], Merck Millipore, Billerica, MA, USA) was used to
177 resuspend the cells. A volume of 1.5 mL of the cell solution was collected into a 2 mL tube
178 and stored at –20 °C. Nutrients were analysed at a later stage as described in the paragraph
179 above on critical nutrient concentrations.

180

181 *Maximum nutrient uptake rates:* To measure the maximum phosphorus uptake rate, 50 mL of
182 a P-depleted culture was added in 500 ml of growth medium containing 25 µmol L⁻¹ of
183 phosphate. The reduction in dissolved phosphorus concentration was then monitored after 0,
184 2, 4, 6 and 24 hours. Plotting the external phosphorus concentrations versus time revealed an

185 initial linear phase so that the maximum P-uptake rate was estimated by linear regression as
186 the absolute value of the slope. Unfortunately, due to technical problems, the experiments to
187 measure the maximum uptake rate of nitrogen failed so that this rate could not be determined.

188

189 *Maximum nutrient quota:* Maximum intracellular nitrogen and phosphorus quotas, $Q_{max,N}$ and
190 $Q_{max,P}$, were determined by culturing cells in 250 mL Erlenmeyer flasks containing Z8 culture
191 medium with nitrogen and phosphorus in non-limiting concentrations (Kotai 1972). Cultures
192 were grown for three days before cells were collected for nutrient measurements. 50 mL of
193 the sample was centrifuged at $8000 \times g$ and $4 \text{ }^\circ\text{C}$ for 5 minutes. The supernatant was removed
194 as precisely as possible, and cells were washed three times with 50 mL of type-2 analytical
195 grade water and the sample was stored frozen in $-20 \text{ }^\circ\text{C}$. Nitrogen and phosphorus
196 concentrations were analysed at a later stage as described for critical nutrient concentrations.
197 Uptake rate and nutrient quota parameters are based on values obtained from a single
198 microcosm.

199

200 *Half-saturation constant:* The half-saturation constant for phosphorus uptake for each
201 genotype, K_i , was estimated by optimizing the numerical solution to the model described in
202 the methods section further below (eq. 1–3). The model was first solved by using the function
203 lsoda of the R-package deSolve, which automatically switches between stiff and non-stiff
204 integration methods (Soetaert et al., 2010). The model predictions were then fit to the data
205 from the monoculture experiments by minimizing the residual sum of squares with the help
206 of the Broyden-Fletcher-Goldfarb-Shanno algorithm of the function optim in R (version
207 3.0.3). Half-saturation constants for nitrogen uptake could not be determined because of the
208 missing estimates of the maximum nitrogen uptake rate.

209

210 ***Design of competition experiments:***

211 We investigated the competition between the toxic and non-toxic strain under N- and P-
212 limitation and three different nutrient supply regimes: **i)** Competition experiments under
213 constant supply were conducted in chemostats (continuous cultures) of 470 mL culture
214 volume with a constant dilution rate of 0.1 day^{-1} (“continuous supply”). The set up and
215 maintenance of the chemostats followed the same methodology as described previously
216 (Fussman et al., 2000; Hiltunen et al., 2013), and all containers used during the experimental
217 procedures were washed with 2% HCl. Chemostat cultures were started as follows: both
218 genotypes were grown in monocultures into steady state, cell densities were enumerated and
219 then both genotypes were mixed in equal proportion. **ii)** Competition experiments under
220 pulsed nutrient supply were run as semi-continuous cultures, using 1000 mL Erlenmeyer
221 flasks that contained 500 mL of culture. For the experiments with short pulse intervals, 10%
222 of the culture was replaced every third day with fresh growth medium (“3-day-interval-
223 pulse”). **iii)** For the experiments with a single large pulse, 10% of the mixed culture was
224 replaced on the first day with fresh growth medium (“single-pulse”). Subsequent
225 replacements occurred every third day, where 10% of the mixed culture was replaced with
226 fresh growth medium devoid of nitrate or phosphate, respectively. Experiments were carried
227 out in Z8 culture medium without chloramphenicol and with varying nitrogen (N) and
228 phosphorus (P) concentrations, depending on the type of nutrient limitation and supply
229 regime. An overview of the different types of experiments and their corresponding nutrient
230 supply concentrations and supply regime is given in Table 1. In the two different pulsed
231 experiments, the total amount of limiting nutrient provided during the course of the
232 experiment was the same. All experiments were replicated at least three times and were
233 initially inoculated at a 1:1 ratio of toxic versus non-toxic strain cells. Light was
234 unidirectional, with $90\text{-}100 \mu\text{mol photons m}^{-2} \text{ s}^{-1}$ intensity. Furthermore, in all experiments,

235 the light intensity measured at the backside of the culture vessels was $35 \mu\text{mol photons m}^{-2}$
236 s^{-1} or higher, which indicates that light availability within the culture was growth-saturating
237 (Hesse et al., 2001; Huisman et al., 2002). Light intensity was measured with a model LI-250
238 light meter (LI-COR, Lincoln, NE, USA).

239 ***Statistics:***

240 We used Student's *t*-test to investigate the effect of genotype identity on the different
241 nitrogen and phosphorus parameters, and we used a repeated measures ANOVA
242 (RMANOVA) to investigate the effects of growth medium and supply regime on *Microcystis*
243 cell density. All analyses were performed with PASW statistics (SPSS Inc. Chicago IL, v.
244 20.0) software.

245

246 ***Mathematical model and numerical simulations:***

247 Many phytoplankton species take up nutrients such as nitrogen and phosphorus in excess
248 when these resources are abundant and store them intracellularly until periods of low nutrient
249 availability arise. Therefore, we use a variable-internal-stores model ("Droop-model") in
250 order to investigate the effect of nutrient fluctuations on the competition between two
251 *Microcystis* genotypes (Droop, 1973). This model has been analyzed previously by Grover
252 (1991) and will therefore be explained here only in brief. The model assumes a spatially
253 homogeneous habitat, such as a well-mixed chemostat. The population density of a specific
254 genotype, N_i , increases with a certain specific growth rate, μ_i , and it decreases with a constant
255 rate, D , the chemostat dilution rate. The specific growth rate of a genotype depends on its
256 intracellular phosphorus pool, Q_i , which increases with the specific uptake rate, v_i , and
257 decreases due to cell division.

258 The phosphorus concentration in the water, R , increases due to a constant inflow of
 259 phosphorus from the supply vessel and decreases due to chemostat dilution and uptake by
 260 phytoplankton cells. The dynamics of N_i , Q_i and R in the chemostat are then given by

261

$$262 \quad \frac{dN_i}{dt} = \mu_i(Q_i)N_i - DN_i \quad (1)$$

$$263 \quad \frac{dQ_i}{dt} = v_i(R, Q_i) - \mu_i(Q_i) Q_i \quad (2)$$

$$264 \quad \frac{dR}{dt} = D(R_{in} - R) - \sum_{i=1}^2 v_i(R, Q_i) N_i \quad (3)$$

265

266 Here, R_{in} defines the supply concentration of phosphorus. The specific growth rate of strain i ,
 267 μ_i , is a saturating function of the internal phosphorus quota, and is given by

268

$$269 \quad \mu_i = \mu_{max,i}' \left(1 - \frac{Q_{min,i}}{Q_i} \right), \quad (4)$$

270

271 where $\mu_{max,i}'$ denotes the maximum specific growth rate of strain i and $Q_{min,i}$ denotes the
 272 persistence quota, i.e. the phosphorus concentration inside the cell of strain i at which growth
 273 ceases (i.e. where $\mu(Q_{min}) = 0$). Note, that the maximum specific growth rate, $\mu_{max,i}'$, is never
 274 reached because phosphorus quotas do not increase infinitely. Instead, the realized maximum
 275 specific growth rate, $\mu_{max,i}$, which is reached at the point where the phosphorus quota is at its
 276 maximum, $Q_{max,i}$, is always lower. The phosphorus uptake rate of strain i , v_i , is given by

277

$$278 \quad v_i = v_{max,i} \left(\frac{R}{K_i + R} \right) \left(\frac{Q_{max,i} - Q_i}{Q_{max,i} - Q_{min,i}} \right), \quad (5)$$

279

280 where K_i describes the half-saturation constant of genotype i for phosphorus uptake. Thus, v
 281 increases with the external phosphorus concentrations as described by the Michaelis-Menten-

282 function and it decreases with increasing intracellular phosphorus quota. For each genotype,
 283 the model parameters were measured in monocultures, except for the half-saturation
 284 constants for phosphorus uptake, K_i , which were estimated by the curve-fitting method as
 285 describe above. The parameterized model was then used to investigate the competition for a
 286 large number of different phosphorus pulse heights and pulse frequencies numerically. In
 287 total, simulations were run for $251 \times 351 = 88101$ different combinations of pulse height and
 288 pulse period in the range of 10^{-2} (= 0.01) and $10^{0.5}$ (= 3.16) $\mu\text{mol P L}^{-1}$ and of 10^{-2} (= 0.01)
 289 and $10^{1.5}$ (= 31.6) days. Note that to be comparable with the experimental results, the model
 290 simulations were not run until the final outcome of competition was observed, but they were
 291 stopped after 40 days. The observed genotype frequencies at this time point were categorized
 292 and the result was drawn on the plane of pulse height and frequency (Fig. 2).

293

294 **Results**

295 ***Monocultures:***

296 *Nitrogen parameters:* The monoculture experiments revealed that the toxic and the non-toxic
 297 *Microcystis* genotype differed in their growth and resource utilization traits. Under nitrogen-
 298 limited conditions, the critical nitrogen concentration, R_N^* , of the toxic genotype measured in
 299 chemostats at steady state was clearly lower than that of the non-toxic strain (t -test: $t(6) =$
 300 7.56 , $p < 0.001$). The toxic genotype also exhibited a higher maximum growth rate, μ_{max} (t -
 301 test: $t(10) = -3.81$, $p = 0.003$). Even though we do not have replication required for statistical
 302 testing, it appears that the WT also had a lower minimum nitrogen quota, $Q_{min,N}$, and a higher
 303 maximum nitrogen quota, $Q_{max,N}$, compared to the non-toxic genotype (Table 2).
 304 Unfortunately, we do not have estimates for the maximum nitrate uptake rate because the
 305 uptake experiments failed due to technical problems. Consequently, it was also not possible

306 to estimate the half-saturation constant for nitrate uptake, K_N , with the help of curve fitting, as
307 the two parameters are interdependent.

308

309 *Phosphorus parameters:* Under phosphorus-limited conditions, the critical phosphorus
310 concentration, R_P^* , of the toxic genotype measured in chemostats at steady state was higher
311 than of the non-toxic genotype (t -test: $t(3) = -5.0$, $p = 0.015$). The toxic genotype also had a
312 higher maximum specific growth rate than the non-toxic genotype (Grover, 1990, 1991; t -
313 test: $t(10) = -2.56$, $p = 0.028$). In contrast, the non-toxic strain had a higher maximum uptake
314 rate for phosphorus, higher maximum phosphorus quota, and a slightly lower minimum
315 phosphorus quota.

316

317 *Competition dynamics:*

318 *Nitrogen-limitation:* The replicated competition experiments under nitrogen-limitation
319 revealed very consistent results. Under constant supply, the total cell density remained
320 constant throughout the experiment (Fig. 1A). Yet, the relative frequency of the toxic
321 genotype gradually increased until it reached 1 after approximately 40 days.
322 Correspondingly, the relative frequency of the non-toxic genotype decreased until it was
323 washed out within the same time period. Thus, under constant nitrogen supply, the toxic
324 genotype clearly outcompeted the non-toxic genotype. When nitrogen was supplied in pulses
325 every three days, the total cell density increased in the beginning of the experiments and
326 reached a constant density after 15 days. The relative frequency of the toxic genotype showed
327 a steep increase within the first six days of the experiment, followed by a more moderate rise
328 until it completely dominated the culture after 24 days. By definition, the relative frequency
329 of the non-toxic genotype exhibited the opposite pattern. When nitrogen was supplied as a
330 single large pulse at the beginning of the experiments, the dynamics of the total cell density

331 showed a hump-shaped curve (Fig. 1C). The relative frequency of the toxic genotype rose
332 steeply in the beginning and reached 100% after 9 days, while the non-toxic genotype was
333 washed out.

334

335 *Phosphorus limitation:* The total cell density in the experiments with constant phosphorus
336 supply remained constant throughout the experiments and did not differ from the experiments
337 with constant nitrogen supply (Fig. 1A,D; average cell density during the experiment, N-
338 limitation: $6.43 \pm 0.49 \times 10^6$ cells mL⁻¹; Fig. 3D; P-limitation $6.31 \pm 0.39 \times 10^6$ cells mL⁻¹;
339 RMANOVA $F_{1,5} = 0.034$, $p = 0.860$). Yet, the relative genotype frequencies showed
340 considerable variation between replicates. Although the critical phosphorus concentrations
341 measured in monocultures suggested that the non-toxic genotype should win the competition,
342 three competition experiments showed that the non-toxic genotype increased in relative
343 frequency or outcompeted the toxic mutant, while two competition experiments showed
344 coexistence of both genotypes at approximately equal frequencies. Yet, the replicate
345 experiments showed consistent results in the pulsed experiments. In the 3-day-interval-pulse
346 experiments, the total cell density remained constant throughout the experiment and did not
347 differ from the constant supply experiment. Both genotypes coexisted at approximately equal
348 densities for 27 days. (Fig. 1E). In the single-pulse experiments, the total cell density showed
349 a hump-shaped curve, and both genotypes coexisted again at approximately equal densities
350 (Fig. 1F).

351

352 ***Rate of competitive exclusion:***

353 As described in the previous section, the nutrient supply regime affected the rate of
354 competitive exclusion, whereby the effect depended on the type of nutrient limitation. We
355 therefore plotted the natural logarithm of the ratio of mutant to wild type relative frequencies

356 versus time, and calculated the rate of competitive exclusion from the slope of the linear
357 regression line (see e.g. van de Waal et al., 2011). In the nitrogen-limited experiments, the
358 rate of competitive exclusion increased with pulse intensity, being slowest under constant
359 supply and fastest when a single large nitrogen pulse was given (Tab. 3; Fig. 1A–C). In the
360 phosphorus limited experiments, competitive exclusion was reversed and was fastest under
361 constant supply, while pulsed phosphorus supply considerably slowed down competitive
362 exclusion (Tab. 3, Fig. 1D–F).

363

364 *Model predictions concerning competition for phosphorus:*

365 Because the phosphorus-limited experiments did not provide very clear results, we also
366 investigated competition under phosphorus limitation with the help of numerical simulations
367 using the growth parameters for each genotype measured in monoculture. The results
368 observed after 40 days show that the non-toxic genotype wins the competition under very
369 short pulse intervals and low pulse height in the lower left corner of Figure 2. Because the
370 results are visualized in a log-log plot, the dominance of the non-toxic genotype is actually
371 confined to a rather small parameter space of nearly-constant phosphorus supply. The
372 dominance region of the non-toxic genotype extends further as a fine, narrowing, diagonal
373 band towards the upper right corner of the graph, but then bends off beforehand and
374 continues rising more slowly with increasing oscillations. To the upper left side of this stripe
375 extends a large region of longer pulse intervals but relatively high pulses, where both
376 genotypes are able to coexist for the period of 40 days. In the lower right corner of the graph,
377 the pulses are too seldom and too low so that both genotypes are washed out before the end of
378 the experiments.

379

380 **Discussion**

381 Our study demonstrates that closely related genotypes of one species can have distinct
382 nutrient physiologies. Our study also shows that these physiological differences can have
383 consequences for the growth and competition between these genotypes, which eventually
384 may have a profound impact on the functioning and the toxicity of blooms in nature. We here
385 discuss the physiological and theoretical background and the ecological implications of these
386 results.

387

388 ***Physiological differences of toxic and non-toxic genotypes***

389 Genetically, the non-toxic mutant genotype used in our study differed from the toxic wild
390 type genotype only due to the deletion of the *mcyB* gene. Transcription of *mcy*-genes has
391 been shown to be stimulated by light (Kaebernick *et al.*, 2000; Tonk *et al.*, 2005) and a
392 number of recent studies suggest that adaptation to high light conditions and oxidative stress
393 protection might be the primary role of microcystin (Zilliges *et al.*, 2011; Makower *et al.*,
394 2015; Meissner, Steinhauser & Dittman, 2015), though also other functions have been
395 suggested (Kaplan *et al.*, 2012). Interestingly, however, our monoculture experiments reveal
396 that the toxic and the non-toxic genotype show significant differences also in other traits,
397 which do not seem to be directly linked to the cellular functions of microcystins that have so
398 far been described. The toxic genotype appeared to have a higher maximum specific growth
399 rate, μ_{max} , than the non-toxic genotype in both types of growth media (see Tab. 1). Yet, this is
400 in disagreement with Hesse *et al.* (2001) and Briand *et al.* (2012), who did not find any
401 difference between the two genotypes. The toxic genotype also had higher maximum quotas
402 for nitrogen, $Q_{max,N}$, and phosphorus $Q_{max,P}$, which suggest that it can store more nutrients and
403 that it can maintain high uptake rates over a longer time period (Tab. 1; Grover, 1991). The
404 non-toxic genotype had a lower critical phosphorus concentration, R_P^* , a lower minimum

405 phosphorus requirement, $Q_{min,P}$, and a higher maximum phosphorus uptake rate, $v_{max,P}$, which
406 suggest that it can deal better with constantly low phosphorus concentrations and that it can
407 better exploit small phosphorus pulses. One possible explanation for this result is that
408 microcystin production is costly in terms of ATP, and thus also in terms of phosphorus,
409 because the corresponding gene cluster and enzyme complex need to be maintained active
410 (Bickel, Lyck & Utkilen, 2000). However, Briand et al. (2106) found in their experiments
411 with the same strains (WT and mutant) under monoculture and co-culture conditions that the
412 mutant which is not able to produce microcystin, produces higher concentrations of the
413 remaining cyanopeptides. In the light of this finding, interpreting the cost of microcystin
414 production is not so straight forward.

415 It is surprising that the toxic genotype also had a lower minimum nitrogen quota, $Q_{min,N}$, and a
416 lower critical nitrogen concentration, R_N^* , than the non-toxic genotype. Yet, this result has to
417 be taken with care, because the very high R_N^* -values compared to other studies (Brussaard &
418 Riegman, 1998; Wilken et al., 2013) indicate that something else might have been limiting
419 growth during these measurements as well. In fact, one would expect the toxic genotype to
420 have a higher nitrogen demand because the production of microcystins is costly in terms of
421 nitrogen (e.g. Long et al 2001; Downing *et al.*, 2005; van de Waal *et al.*, 2009, 2010).

422 We do not yet have a clear explanation for the physiological differences found in our study,
423 Yet, it is known that the deletion of the *mcyB* gene has various consequences for the cell, e.g.
424 for pigmentation and cellular structure (Hesse *et al.*, 2001), but also for the expression of
425 other genes and proteins, which are related to photosynthesis and respiration, energy-
426 metabolism or carbon-nitrogen metabolism, to name a few (Alexova *et al.*, 2011; Zilliges *et*
427 *al.*, 2011; Makower *et al.*, 2015). Thus, the intracellular effects of microcystin production
428 seem manifold, and it is therefore likely that it has side-effects on other phenotypic traits as
429 well.

430

431 ***Competition in pulsed habitats***

432 Our competition experiments show that the distinct nutrient physiologies of the toxic and the
433 non-toxic genotype discovered in the monoculture experiments also have important
434 consequences for their population dynamics. Our results are largely in agreement with
435 common resource competition models (Tilman, 1982; Grover, 1991, 1997). Under constant
436 nutrient inflow competition was won by the affinity-specialist (sensu Sommer, 1984), i.e. by
437 the genotype which had the lowest R^* -value for the limiting nutrient (Tab. 1; Fig. 1A and D).
438 Interestingly, the toxic and the non-toxic genotype exhibited a trade-off in the competitive
439 ability for nitrogen and phosphorus, which is also commonly observed between
440 phytoplankton species (Edwards, Litchman & Klausmeier, 2011). On the one hand our results
441 are consistent with Vézic *et al.* (2002) and Lei *et al.* (2015) who found higher growth rates of
442 the mutant genotype and a non-toxic strain under phosphorus limitation. On the other hand
443 our results are inconsistent with Vézic *et al.* (2002) and Briand *et al.* (2012) who found that
444 the mutant genotype, rather than the wild type, was growing faster under N-limitation, and
445 Lei *et al.* (2015) did not observe a clear difference. One reason for this discrepancy might be
446 that these studies used batch cultures rather than continuous or semi-continuous cultures,
447 where the dominant species or genotype in the competition does not need to be the affinity-
448 specialist. Another reason might be that all three experiments differed in the degree of N-
449 limitation, and that we cannot assure that N-limitation in our experiments was absolute.

450 Under pulsed nutrient inflow resource competition theory predicts that the growth-specialist
451 (sensu Sommer, 1984), i.e. the species with the highest maximum growth rate, μ_{max} , wins the
452 competition under small (and frequent) nutrient pulses, while the storage specialist (sensu
453 Sommer, 1984), i.e. species with lowest Q_{min} , highest Q_{max} and highest v_{max} , wins the

454 competition under large (and unfrequent) nutrient pulses (Grover 1991, 1997). Our
455 monoculture and competition experiments together suggest that the toxic-genotype was the
456 growth- as well as the storage-specialist for both nitrogen and phosphorus (Tab. 1; Fig.
457 B,C,E,F). Thus, under nitrogen limitation, we did not observe a trade-off in the competitive
458 ability under constant and pulsed nutrient inflow. Under phosphorus limitation, however, this
459 trade-off seemed to be existent. Here, the non-toxic genotype was the superior competitor
460 under constant phosphorus inflow (Tab.1, Fig 1D). Under pulsed phosphorus inflow we did
461 not observed a full reversal of competitive dominance, but the non-toxic genotype no longer
462 outcompeted the toxic genotype. Instead, both genotypes coexisted for the duration of the
463 experiment (Fig. 1E,F). Moreover, our simulation results indicate that the dominance of the
464 toxic genotype would further increase with higher phosphorus pulse intensity (results not
465 shown). This result is in agreement with the general observation, that phytoplankton species
466 show a trade-off between the competitive ability under constant and variable phosphorus
467 limitation (Edwards, Klausmeier & Litchman, 2013).

468 Previous studies with toxic and non-toxic *Microcystis* strains have demonstrated distinct
469 competitive abilities under light carbon, nitrogen and phosphorus limitation (Kardinaal *et al.*,
470 2007; Van de Waal *et al.* 2011; LeBlanc Renaud *et al.*, 2011; Briand *et al.*, 2012; Lei *et al.*,
471 2015). Yet, as Briand *et al.*, (2012) and Lei *et al.* (2015) conducted their studies in batch
472 cultures competition for nitrogen and phosphorus in continuous or semi-continuous culture
473 has not been investigated before. At first sight, our experiments suggest that phosphorus
474 limitation might favor the dominance of non-toxic genotypes, and that microcystin
475 production does not provide a clear benefit under P-limitation. As P-limitation should not
476 affect the reducing power of cells to the same extent as C- or N-limitation (Sternner and Elser,
477 2002), this results is in line with the hypothesis that microcystins are beneficial under
478 oxidative stress (Zilliges *et al.*, 2012, Makower *et al.*, 2015; Meissner *et al.*, 2015). Yet, it has

479 to be pointed out that the competitive dominance of the mutant genotype under constant
480 phosphorus limitation was rather fragile and seemed to be limited to a very low parameter
481 range of nearly constant conditions, as rather small nutrient pulses already enabled the toxic
482 genotype to coexist. (Fig. 1D, Fig. 2). It therefore remains an open question whether such
483 highly constant conditions occur in natural freshwater systems and whether phosphorus
484 limitation can really mitigate bloom toxicity.

485

486 *Nutrient limitation in natural blooms*

487 Our experiments suggest that toxic *Microcystis* genotypes might be better competitors
488 under nitrogen limitation than non-toxic genotypes, while (constant) phosphorus limitation
489 favors non-toxic genotypes. This result is consistent with several field studies showing that
490 low N:P ratios seem to favor toxic *Microcystis* genotypes over their non-toxic counterparts
491 (Davis et al., 2009; Rinta-Kanto et al., 2009, Gobler et al., 2016). Furthermore, higher
492 microcystin concentrations have been linked to low N:P ratios in four reservoirs in the United
493 States (Harris et al., 2016), indirectly indicating higher frequencies of toxic genotypes.
494 However, our results do not explain why the relative frequency of toxic genotypes has been
495 found to increase with increasing N concentrations (Yoshida et al., 2005; Davis et al., 2010).

496

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705 **Legends**

706 **Table 1.** Nutrient supply concentrations and supply regimes of the different competition
 707 experiments. * Please note that these supply rates of the limiting nutrient are only for the
 708 beginning of the experiment. After the start of the experiment we did not provide any of the
 709 limiting nutrient (N or P).

710

711 **Table 2.** Nitrogen and phosphorus parameters of the toxic and the non-toxic *Microcystis*
 712 genotypes. Values marked with ^a were estimated by curve fitting. The maximum nitrogen
 713 uptake rate, $v_{max,N}$, and the corresponding half-saturation constant, K_N , were not determined
 714 due to technical problems.

715

716 **Table 3.** Rates of competitive exclusion in the six different competition experiments. Rates
 717 (cells mL⁻¹ day⁻¹) are calculated as the slope of the linear regression of the relative frequency
 718 of the non-toxic mutant versus time. Significant positive slopes indicate competitive
 719 exclusion of the wild type by the mutant; significant negative slopes indicate competitive
 720 exclusion of the mutant by the wild type. Non-significant slopes indicate coexistence.
 721 Significant slopes values ($p < 0.05$) are marked in bold.

722

723 **Figure 1.** Relative frequencies of the non-toxic wild type strain (red line, filled symbols) and
 724 the non-toxic mutant (blue line, closed symbols) during competition for nitrogen (panels
 725 A–C) and phosphorus (panels D–F) at different levels of nutrient fluctuations: (A, D)
 726 nutrients supplied continuously, (B, E) nutrients pulsed at 3-day-intervals, (C, F) single large
 727 nutrient pulse at the beginning of the experiments. Triangles with green dotted line show total

728 *Microcystis* densities (cells mL⁻¹). Symbols represent means of four replicate microcosms
729 (except in A, $n = 5$), and vertical bars show standard error of the mean.

730 **Figure 2.** Simulation results showing the outcome of competition between the toxic
731 *Microcystis* wild type PCC 7806 and the non-toxic *mcyB*-deficient mutant genotype for
732 different periods and heights of phosphorus pulses. Note that to be comparable with the
733 experimental results, the graph shows the state of competition after 40 days rather than the
734 equilibrium outcome.

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749 **Table 1**

Type of experiment	N-supply concentration, $R_{in,N}$ ($\mu\text{mol/L}$)	P-supply concentration, $R_{in,P}$ ($\mu\text{mol/L}$)	N:P-supply ratio	Pulse height ($\mu\text{mol N or P}$)	Pulse period (days)
Continuous supply (chemostat)					
N-limited	105	26.25	4	-	-
P-limited	192	3	64	-	-
3-day-pulse (semi-continuous culture)					
N-limited	105	26.25	4	10.5	3
P-limited	160	2.5	64	0.25	3
Single pulse (semi-continuous culture)					
N-limited	1050*	26.25	40*	105	28
P-limited	160	25*	6.4*	2.5	28

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760 **Table 2**

Parameter	Description	Unit	WT	M
<i>Nitrogen</i>				
R_N^*	critical resource concentration	$\mu\text{mol L}^{-1}$	32.08	37.08
$\mu_{max,N}$	realized maximum specific growth rate	day^{-1}	0.549 ± 0.039	0.472 ± 0.036
$v_{max,N}$	maximum nitrogen uptake rate	$\text{fmol cell}^{-1} \text{ day}^{-1}$	n.d.	n.d.
K_N	half-saturation constant for nitrogen uptake	$\mu\text{mol L}^{-1}$	n.d.	n.d.
$Q_{min,N}$	minimum intracellular nitrogen content	fmol cell^{-1}	8.019	8.506
$Q_{max,N}$	maximum intracellular nitrogen content	fmol cell^{-1}	18.92	16.36
<i>Phosphorus</i>				
R_P^*	critical resource concentration	$\mu\text{mol L}^{-1}$	0.073 ± 0.016	0.03 ± 0.0
$\mu_{max,P}$	maximum specific growth rate	day^{-1}	0.448 ± 0.040	0.398 ± 0.026
$v_{max,P}$	maximum phosphorus uptake rate	$\text{fmol cell}^{-1} \text{ day}^{-1}$	1.853	2.272
K_P	half-saturation constant for phosphorus uptake	$\mu\text{mol L}^{-1}$	3.004 ^a	3.004 ^a
$Q_{min,P}$	minimum intracellular phosphorus content	fmol cell^{-1}	0.285	0.273
$Q_{max,P}$	maximum intracellular phosphorus content	fmol cell^{-1}	1.880	1.210

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765 **Table 3**

Resource supply regime	Limiting nutrient	n	Slope*	S.E.	<i>t</i> -value	<i>p</i> -value	Outcome of competition	Fig.
Constant	N	5	-0.11	0.001	-12.8	<0.0001	WT wins	1A
Constant	P	4	0.012	0.004	3.14	0.005	MT wins	1B
3-day pulse	N	4	-0.144	0.015	-9.80	<0.0001	WT wins	1C
3-day pulse	P	4	-0.001	0.002	-0.59	0.559	Co-existence	1D
Single pulse	N	4	-0.541	0.038	-14.1	<0.0001	WT wins	1E
Single pulse	P	4	0.004	0.002	1.77	0.087	Co-existence	1F

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