Appendix C

Chemostat conditions
We used a nutrient-rich mineral medium to avoid nutrient limitation (Stomp et al. 2004). NaCl was added to obtain a salinity of ~12 g/L, resembling the brackish waters of the Baltic Sea. The chemostat vessels were bubbled with compressed air enriched with CO₂ to avoid carbon limitation. The CO₂ inflow was controlled by mass flow controllers (BROOKS Smart Mass Flow) maintaining the pH between 8 and 8.5. The chemostat vessels were maintained at a constant temperature of 21 °C by water jackets placed between the light sources and the chemostat vessel, and connected to a Coloura thermocryostat.

Background turbidity
The absorption spectrum of the background turbidity, \(K_{bg}(\lambda)\), was calculated from the spectrum of the incident light intensity, \(I_{in}(\lambda)\), and the spectrum of light penetrating through the chemostat vessels, \(I_{out}(\lambda)\), when filled with mineral medium only:

\[
K_{bg}(\lambda) = \frac{\ln(I_{in}(\lambda)) - \ln(I_{out}(\lambda))}{z_M}
\]  
(C1)

The background absorption in our shallow chemostat vessels was largely independent of wavelength, and thus could be treated as a wavelength-independent parameter \(K_{bg}\).

Population densities
Samples were taken from the chemostat experiments daily, and were fixed with a solution of 10% gluteraldehyde and 1% formaldehyde (Tsuji & Yanagita 1981). The population densities of the species were counted in duplicate with a MoFlow flowcytometer (Dako Cytomation, Fort Collins, CO, USA), equipped with a blue laser (488 nm) and a red laser (633 nm). The flow cytometer could distinguish between red and green cyanobacteria based upon their different fluorescence signals (Jonker et al. 1995; Stomp et al. 2004). Cells rich in phycoerythrin emitted yellow-orange light (550–620 nm) when excited by the blue laser, whereas cells rich in phycocyanin emitted far red light (> 670 nm) when excited by the red laser. The flow cytometer distinguished between the single-celled picocyanobacteria and the much larger filaments of *Pseudanabaena* by their size (using side scattering). Since the picocyanobacteria were much smaller than the *Pseudanabaena* filaments, the counts obtained by flow cytometry were not a good indicator of the biomasses of the three species. For comparison, therefore,
the population densities of the species were converted to the total light absorption (per cm) by each species, $A_i$, which can be expressed as:

$$A_i = \frac{1}{z_m} \ln \left( \frac{\int_{700}^{400} I_m(\lambda) d\lambda}{\int_{400}^{700} I_m(\lambda) e^{-k(\lambda) N z_m} d\lambda} \right)$$

The fluorescence signals of the *Pseudanabaena* filaments obtained from the blue and red laser provided information on the relative content of phycoerythrin and phycocyanin in *Pseudanabaena*. Using monoculture experiments of *Pseudanabaena* grown under different red-light and green-light conditions, we calibrated these fluorescence signals with absorption spectra measured with the AMINCO DW-2000 double-beam spectrophotometer. This enabled estimation of the fraction phycoerythrin (i.e., the parameter $x_i$ in Equation 6.4 of the main text), and hence the absorption spectrum of *Pseudanabaena* during competition could be reconstructed from the flow-cytometer measurements.