Supporting information

Thermocline depth and euphotic zone thickness regulate the abundance of diazotrophic cyanobacteria in Lake Tanganyika

5 Benedikt Ehrenfels^{1,2}, Maciej Bartosiewicz³, Athanasio S. Mbonde⁴, Kathrin B.L. Baumann^{1,2}, Christian Dinkel¹, Julian Junker^{5,6}, Tumaini Kamulali⁴, Ismael A. Kimirei^{4,7}, Daniel Odermatt¹, Francesco Pomati⁸, Emmanuel A. Sweke^{4,9}, Bernhard Wehrli^{1,2}

1 Eawag, Swiss Federal Institute of Aquatic Science and Technology, Department Surface Waters – Research and Management, Kastanienbaum, Switzerland

- ² 10 ETH Zurich, Institute of Biogeochemistry and Pollutant Dynamics, Zurich, Switzerland 3 University of Basel, Department of Environmental Sciences, Basel, Switzerland 4 TAFIRI, Tanzania Fisheries Research Institute, Kigoma, Tanzania 5 Eawag, Swiss Federal Institute of Aquatic Science and Technology, Department Fish Ecology and Evolution, Kastanienbaum, Switzerland
- 15 ⁶University of Bern, Institute of Ecology & Evolution, Bern, Switzerland 7 TAFIRI, Tanzania Fisheries Research Institute, Kigoma, Tanzania 8 Eawag, Swiss Federal Institute of Aquatic Science and Technology, Department Aquatic Ecology, Dübendorf, Switzerland 9 DSFA, Deep Sea Fishing Authority, Zanzibar, Tanzania

Correspondence to: Benedikt Ehrenfels [\(benedikt.ehrenfels@eawag.ch\)](mailto:benedikt.ehrenfels@eawag.ch)

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1 Sampling

1.1 Station information

Table S1: Station dates and coordinates of the two expeditions at the end of the dry season (Septemper/October 2017) and the end of the 35 rainy season (April/May 2018).

1.2 Phytoplankton community analyses

Table S2: List of taxa identified by microscopic phytoplankton community analyses.

40 **2 Additional information on the thermal structure of the water column**

With the data at hand, we cannot identify the processes responsible for the formation of a deep, secondary thermocline in Sep/Oct (Table S3 & Fig. S1), but splitting of the thermocline may have resulted from surface and subsurface water currents flowing in opposite directions during this time of the year (Verburg et al., 2011). The surface thermocline in Apr/May might be 'seasonal' thermocline (Hecky et al., 1996) or biologically induced (see section 5).

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Table S3: Depth and strength, measured as buoyancy frequency (N^2) , of the thermocline and euphotic depth (z_{eu}) during the end of the dry season (September/October 2017) and the end of rainy season (April/May 2018). At some stations, secondary thermoclines had formed. The primary thermoclines separated nitrate-depleted surface water masses from underlying nitrate-rich waters as used in Fig. 1 & 3. The temperature profile of station 9 exhibited no thermocline.

		secondary surface thermocline		primary thermocline		secondary deep thermocline		
	station	depth [m]	N^2 [s ⁻²]	depth [m]	N^2 [s ⁻²]	depth [m]	N^2 [s ⁻²]	Zeu [m]
Sep/Oct 2017	1			34	2.13E-04	55	2.14E-04	
	\overline{c}			38.5	2.31E-04	64	2.40E-04	
	$\ensuremath{\mathsf{3}}$			36.5	1.66E-04	54.5	1.91E-04	
	4			50	1.78E-04	82.5	1.17E-04	
	$\mathbf 5$			43	1.96E-04	86	2.40E-04	
	$\,6\,$			53.5	2.68E-04	100	1.50E-04	
	7			38	1.89E-04	82.5	1.82E-04	
	8			17.5	1.81E-04			
	$\boldsymbol{9}$			$12*$	1.50E-04*			
Apr/May 2018	$\mathbf{1}$			44.5	2.41E-04	62.5	2.02E-04	54.8
	\overline{c}			63.5	2.50E-04			46.8
	$\ensuremath{\mathsf{3}}$	27.5	2.47E-04	58.5	4.20E-04			40.8
	4	30	1.11E-04	63.0	2.54E-04			51.8
	$\mathbf 5$			55.5	5.03E-04			46.8
	$\,6$			48.5	3.38E-04			46.8
	7			55.5	2.83E-04			51.3
	8			34.0	3.82E-04			51.3
	$\boldsymbol{9}$			23.0	3.80E-04			35.8

50 $*$ for station 9 (Sep/Oct), we indicated the depth and value of the N² maximum, even though there was no clear thermocline. These values were used for the quantitative analyses depicted in Fig. 4 & S3.

Figure S2: Temperature profiles of all 9 sampling stations during September/October 2017 **(a)** and April/May 2018 **(b)** including the 55 approximate depths of the primary and secondary thermocline. The temperature profiles pertaining to April/May 2018 are also presented by C. Callbeck, B. Ehrenfels, K.B.L. Baumann, B. Wehrli, and C.J. Schubert (manuscript in review at *Nat. Comms.*).

Figure S2: Contour plot showing the distribution of the nitrogen (N) deficit in Lake Tanganyika in a North-South transect (whereby 60 station 1 is the northernmost and station 9 the southernmost station) during **(a)** the end of the dry season (September/October 2017) and **(b)** the end of the more stratified rainy season (April/May 2018). Dots indicate samples.

Figure S3: Depth integrated concentrations of the cyanobacterial pigments phycocyanin **(a,b)** and phycoerythrin **(c,d)** for the end of the dry season (September/October 2017) and the end of the rainy season (April/May 2018). We calculated the integrals from the water surface to 125 m depth, to include only photosynthetically active organisms from the oxygenated epi- and metalimnion.

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5 River influence and surface thermocline

The presence of high NO₃ concentrations during sampling at station 3 (one observation at 35 m) may have coincided with the presence of the Malagarasi river inflow. However, our biogeochemical profiles do not support this possibility as neither pH nor conductivity or dissolved organic carbon values (Fig. S4) in this water mass are indicative of the riverine inputs

75 (Athuman and Nkotagu, 2013). We also expect that under severe N depletion in the upper water column any inflowing NO₃ is consumed rapidly after entering surface waters.

Instead, we interpret this NO₃ accumulation as regenerated DIN originating from the surface diazotroph-diatom-chlorophyte community. The high surface cell densities might have created a surface mixed layer that is marked by a near-surface

80 thermocline just above the local NO₃ peak at stations 3 and 4. The strengthening of thermal stratification by the enhanced light absorption of high cell densities is a well-known phenomenon (Edwards et al., 2004; Zhai et al., 2011). This "thermal shielding" (Bartosiewicz et al., 2019) in combination with bioconvetion (Nguyen-Quang and Guichard, 2010) of buoyant cyanobacteria may have created the surface mixed layer with constantly high temperatures of \sim 27 °C in contrast to the steady temperature gradients at the surrounding sites (Fig. S5), and might be responsible for the relatively shallow euphotic depth 85 (40.8 m).

Figure S4: Vertical profiles of physical, biogeochemical and biological data from station 3 at the end of the rainy season (April/May 2018). The left panel shows temperature (T) and euphotic depth (z_{eu}) as well as the nitrate (NO₃'), phosphate (PO₄³), chlorophyll (Chl), 90 and phycocyanin (PC) concentrations, whereas the right panel exhibits conductivity, pH, turbidity and the dissolved organic carbon (DOC) concentrations.

Figure S5: Contour plot of the temperature profiles in a North-South transect at the end of the rainy season (April/May 2018). White lines 95 mark CTD profiles.

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