A 150-year record of phytoplankton community succession controlled by hydroclimatic variability in a tropical lake

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Abstract. Climate and human-induced environmental change promote biological regime shifts between alternate stable states, with implications for ecosystem resilience, function, and services. While these effects have been shown for present-day ecosystems, the long-term response of microbial communities has not been investigated in detail. This study assessed the decadal variations in phytoplankton communities in a ca. 150 year long sedimentary archive of Lake Nong Thale Prong (NTP), southern Thailand using a combination of bulk geochemical analysis, quantitative polymerase chain reaction (qPCR) and lipid biomarkers techniques including compound-specific hydrogen isotope analysis as a proxy for precipitation. Relatively drier and by inference warmer conditions from ca. 1857 to 1916 Common Era (CE) coincided with a dominance of the green algae Botryococcus braunii, indicating lower nutrient levels in the oxic lake surface waters, possibly related to lake water stratification. A change to higher silica (Si) input around 1916 CE was linked to increased rainfall and concurs with an abrupt takeover by diatom blooms lasting for 50 years. These were increasingly outcompeted by cyanobacteria from the 1970s onwards, most likely because of increased levels of anthropogenic phosphate and a reduction in rainfall. Our results showcase that the multi-proxy approach applied here provides an efficient way to track centennial-scale limnological, geochemical and microbial change, as influenced by hydroclimatic and anthropogenic forcing.

1 Introduction

Natural and anthropogenically induced climatic change is often cited as the main factor controlling ecosystem dynamics (Scheffer et al., 2001; Malmqvist et al., 2008; Woodward et al., 2010). The resulting environmental effects easily observed today in plant and animal communities (Scheffer et al., 2001) have been discussed as causing regime shifts (Folke et al., 2004) and/or introducing alternative ecosystem states (Dent et al., 2002). Changes in the ecosystem due to climate dynamics, particularly for microbial communities, would likely affect the capture and storage of CO₂ (Zeglin et al., 2013). However, predicting the effects of changing climate on shifts in microbial community structures even on short-time scales is challenging (Lotter and Birks, 1997; Woodward et al., 2010). This is partially due to the difficulty of coupling micro-scale external and internal ecosystem variables to specific microbial communities, would likely affect the capture and storage of CO₂ (Zeglin et al., 2013). Very little is therefore known about how natural microbial ecosystems respond to climatic changes over longer time scales and whether the associated feedbacks due to climate shifts promote regime shifts and alternative stable states. Moreover, the impact of external processes, such as weathering and runoff intensity, on carbon and nitrogen cycles still remain relatively unknown.
Phytoplankton constitutes one of the most ecologically important groups of microorganisms in aquatic ecosystems (Yin et al., 2011) due to their vital position at the base of the food chain (Carpenter et al., 1987; Young et al., 2013). As photoautotrophs, their activities are directly coupled to climatic changes through CO₂ drawdown, organic matter availability and oxygen production. Natural and/or human-induced disturbances may thus result in far-reaching consequences for the nutrient status of lakes, in which phytoplankton is the primary producer (Yin et al., 2011). For instance, the demise of the botryococcene lipid-producing green algae Botryococcus braunii was linked to early eutrophication in a Norwegian fjord (Smittenberg et al., 2005).

Studies have shown that phytoplankton communities are sensitive to environmental stressors (Woodward et al., 2010; Häder and Gao, 2015) where variations in phytoplankton community structure, abundance and function have been related to changes in lake trophic status (Ravasi et al., 2012; Hou et al., 2014). Accordingly, the emerging paradigm suggests that the structuring of phytoplankton communities, characterized by potential successional shifts in population dynamics, may serve as a tracer for the trophic status in lakes (Mackay et al., 2003; De Senerpont Domis et al., 2007). However, reconstructing the trophic status of lakes based on shifts in dominant phytoplankton communities over time are yet to be explored in detail. Also, the broader linkages between different microbial groups and their response to past environmental conditions are poorly understood. This is partly due to the lack of suitable proxies that can capture and distinguish between the diverse parameters impacting microbial ecosystem structure.

Lipid biomarkers specific for various types of microbes provide an important proxy of microbial ecosystem structure and have therefore been employed in the reconstruction of past ecosystems preserved in sedimentary records (Zimmerman and Canuel, 2000; Coolen et al., 2004; Smittenberg et al., 2005). Lipids are an integral part of the cell membranes of both prokaryotes and eukaryotes. They aid as structural support and as storage compounds within various microbial cells (Jungblut et al., 2009), and can be specific for certain groups or species. Moreover, lipid biomarkers can incorporate information of the chemical and physical environment in which they have formed via their relative abundance or isotopic composition. Because lipids are resistant to post-mortem biodegradation, ecological variations through time can be reconstructed (Zimmerman and Canuel, 2000; Coolen et al., 2004; Smittenberg et al., 2005) by tracking their occurrence and abundance over longer time scales. Moreover, when coupled to hydroclimate variables such as hydrogen isotopes of leaf waxes (δDwax), a proxy for precipitation (e.g. Konecky et al., 2013; Niedermeyer et al., 2014), microbial lipids may help underpin the impact of past climates (wetter/drier) on microbial ecosystem changes (see Supplement).

Quantitative polymerase chain reactions (qPCR) of specific DNA of living organisms and well-preserved DNA in lake sediments are excellent tools to assess present and past microbial ecosystem structures (Coolen and Gibson, 2009; Ravasi et al., 2012; Hou et al., 2014). These analyses provide specific proof of recent and past biological processes by targeting specific microbial taxa and key genes involved in various metabolic pathways (Takai and Horikoshi, 2000; Hou et al., 2014).

Here we explore the novel combination of biolipid analysis, δDwax, qPCR, bulk isotopes of carbon and nitrogen, and sedimentary geochemistry to reconstruct phytoplankton community dynamics over a 150-year history of Lake Nong Thale Prong (NTP), Southern Thailand. Together, these proxies allow unravelling how external forcing (hydroclimate and human impact) influences internal abiotic feedback processes, which in turn control phytoplankton regime shifts.

2 Materials and methods

2.1 Study area, fieldwork, sediment sampling, and dating

NTP (17°11’ N, 99°23’ E) is a shallow (< 4 m water depth), small (~210 m²) sinkhole lake located on the Thai–Malay Peninsula in southern Thailand (Fig. 1) at ~60 m above sea level (Snansieng et al., 1976) (see Supplement). Prior to coring, a preliminary assessment of NTP was conducted based on the catchment geology and topography, basin size and water depth. Sounding in different parts of the lake showed that the deepest part (~3.7 m water depth) was in the northeast and that there was little variability in the distribution of sedimentary materials throughout the lake. Two sediment cores (~74 cm length) were retrieved from the deepest part of the lake in January 2012 using an HTH gravity corer (70 mm diameter, 1 m length). The sediment presented a strong sulfidic smell, suggesting anoxic conditions. Visually, the cores were lithologically similar. Therefore further analysis was performed on one core, while the other core was archived. The sediment core was sliced on-site into 1 cm sub-samples, packed in sterile plastics bags and chilled with ice. The samples were kept at temperature of about 4 °C during transportation from Thailand to Stockholm. Such temperatures have been shown to be low enough to inactivate and preserve whole tropical communities in sediments without the need for dramatic freezing because of a narrower temperature range of activity (e.g. Robador et al., 2016). After arrival at the Department of Geological Sciences, Stockholm University the samples were immediately frozen (~18 °C) until further analysis. The sampled sediment sequence was dated using 210Pb (46.51 keV) and 226Rn (295.2 keV) on an EG&G ORTEC® co-axial low energy photo spectrometer (LEPS) fitted with a high-purity germanium crystal (see Supplement).
A total of 15 sub-samples were taken at different core depths for a low-resolution quantification of total organic carbon (TOC), total nitrogen (TN), and carbon and nitrogen isotopes ($\delta^{13}$C$_{\text{bulk}}$ and $\delta^{15}$N$_{\text{bulk}}$). Samples were freeze-dried and homogenised before analysis. For the stable isotope measurements, the samples were pre-treated with HCl to remove carbonate carbon before analysis on a Carlo Erba NC2500 elemental analyzer, coupled to a Finnigan MAT Delta+ mass spectrometer. $\delta^{13}$C$_{\text{bulk}}$ and $\delta^{15}$N$_{\text{bulk}}$ values are reported in parts per mille (‰) relative to the Vienna PeeDee Belemnite (VPDB, for C) and standard air (for N), respectively, with an analytical error of ±0.15 ‰. The working standards used for the analysis were: (1) acetanilide ($\delta^{13}$C = −27.07 ‰; %C = 71.09 ‰; $\delta^{15}$N = 1 ‰; %N = 10.36 ‰) and (2) methionine ($\delta^{13}$C = −26.23 ‰; %C = 40.25 ‰; $\delta^{15}$N = −2.24 ‰; %N = 9.39 ‰). These standards were calibrated against standards from IAEA.

Relative estimates of the chemical composition of the sediments were obtained by elemental mapping using Environmental Scanning Electron Microscopy (FEI, Quanta FEG 650)-Electron Dispersive Spectroscopy (ESEM-EDS). An aliquot of the dried sediment was mounted on aluminium stubs with carbon tape and imaged at 10 kV in low vacuum. Elemental analysis was conducted in low vacuum with EDS at 30 kV. Approximately 75 elemental maps distributed over 15 samples across the entire core were acquired with the AZTech software, at a horizontal field width of 2 mm, 512 pixels and an average frame count of 5 with 100 µs pixel dwell time. The relative elemental abundance acquired was normalised to 100 ‰.

### 2.3 Biomarker analysis

Description of the biomarker analysis follows that of Chawchai et al. (2015) and Yamoah et al. (2016). Freeze-dried and powdered samples were extracted three times with a mixture of dichloromethane and methanol (DCM–MeOH, 9 : 1, v/v) to obtain a combined total lipid extract (TLE), using a microwave system (MILESTONE Ultra Wave Single Chamber Microwave Digestion System) fitted with a LABTECH smart H150-1000 Water Chiller. The TLE from the sediments was dried in a vacuum concentrator (Scanvac MaxiVac Beta, Labogene ApS, Denmark) before being redissolved in DCM and then adsorbed onto a small amount of silica gel. This was evaporated on a warm plate, under a very gentle stream of nitrogen gas, and placed on top of 15 g silica gel (deactivated with 5 % (wt.) H$_2$O) in 6 mL glass SPE tubes. Hydrocarbon (F1), ketone (F2) and polar (F3) fractions were recovered with pure hexane, a hexane and DCM mixture (1 : 1) and DCM–MeOH (1 : 1), respectively. F2 and F3 samples were stored in the freezer for later use. The F1 fraction was analysed on a Shimadzu GCMS-QP2010 Ultra gas chromatography–mass spectrometer (GC–MS), equipped with an AOC- 20i auto sampler and a split-splitless injector operated in splitless mode. A Zebron ZB-5HT Inferno GC column (30 m × 0.25 mm × 0.25 µm) was used for separation.

The GC oven temperature was programmed from 60 to 180 °C at a ramp of 20 °C min$^{-1}$ followed by a ramp of 4 °C min$^{-1}$ until 320 °C where it was held for 20 min. MS operating conditions were set to an ion source temperature of 200 °C and 70 eV ionisation energy. Spectra were collected using GC solution Workstation software (v2). n-alkanes, C$_{25}$ highly branched isoprenoids (HBIs) and botryococcene compounds were identified by retention times and comparison against mass spectra from the literature. Quantification of the n-alkanes, C$_{25}$ HBIs and botryococcene compounds was done with an external standard consisting of a mixture of C$_{20}$–40 n-alkanes of known concentration. Specifics on the mass spectra and retention times of the n-alkanes, HBIs and Botryococcenes, including chromatograms as reference, are included in Supplement (Figs. S1–S6).

### 2.4 δD analysis of leaf waxes

The F1 fraction was further separated into three fractions (F1a, F1b, and F1c) over a pipette column filled with 10 %...
AgNO₃-coated silica gel. F1a, which comprises n-alkanes, was eluted with hexane; F1b, made up of a few unidentified compounds, was eluted with hexadecane-DCM (1:1); and F1c consisting of HBIs and botryococccenes, was eluted with DCM-Acetone (9:1). F1b and F1c were also stored in the freezer for further analysis. F1a was analysed by gas chromatography–isotope ratio monitoring–mass spectrometry (GC-IRMS) using a Thermo Finnigan Delta V Plus mass spectrometer interfaced with a Thermo Trace GC 2000 using a GC Isolink II and Conflo IV system. Helium was used as a carrier gas at constant flow mode. A detailed description of the GC-IRMS program is described in Chawchai et al. (2015). Instrumental performance and calibration of the reference gas (H₂) was achieved by running a standard mixture of n-alkanes with a known isotopic composition (reference mixture A4, provided by Arndt Schimmelmann, Indiana University, USA) several times daily. All analyses performed follow that of Sessions et al. (1999, 2001). Results are reported as the weighted mean of triplicate measurements with an average standard deviation of both standards and samples around 4% (see Supplement).

2.5 DNA extraction and qPCR

Freeze-dried samples were selected according to initial biomarker screening results, in order to estimate the abundance of different groups of organisms related to: (1) the Prokarya, Archaea and bacteria, cyanobacteria, and microorganisms involved in anaerobic methane cycling ( quantification of the mcrA gene, e.g., Hallam et al., 2003, 2004), (2) Eukarya, diatoms, and Botryococcus sp. The samples were analysed in order to specifically reflect the sample conditions used for the biomarker analysis. Freeze-drying was not expected to introduce significant biases, but it enhances cell breakage and the release of intracellular DNA, following the freeze thaw method of DNA extraction (e.g. Tsai and Olson, 1991). This is especially useful for soil and sediment samples (e.g. Tsai and Olson, 1991). Around 0.2 g (from 0.17 to 0.26 g) of freeze-dried sediment was extracted for DNA, using the MoBio PowerSoil® DNA kit (Carlsbad, CA), following the manufacturer’s instructions. About 500 µL of sterile PBS 1X was also added to PowerSoil® Bead tube in order to enhance cell lysis efficiency.

The qPCR amplifications were conducted in 96 well qPCR plates in a CFX96 Touch™ Real-Time PCR Detection System Instrument (C1000 Touch™ Thermal, Cycler, Bio-Rad) and its software. The reactions consisted of a final volume of 25 µL, using the SsoAdvanced™ Universal SYBR® Green Supermix (Bio-Rad) following the manufacturer’s recommendations. Reactions run in 35 cycles contained 5 µL of DNA template and specific primer sets at their appropriate concentrations and annealing temperatures (see Supplement).

Standard curves were calibrated using 10-fold serial dilutions from pure cultures of each representative target group (Supplement). The qPCR detection of 16S rRNA genes, 18S rRNA genes in all of the samples and in 10-fold serial dilutions used to construct the standard curves was run in triplicates. For each qPCR, several negative controls were performed in order to check for laboratory contamination. The efficiencies of the qPCR analyses was up to 90% with a correlation to the standard calibration curve of up to $R^2 = 0.996$ (see Supplement).

A total of 16S rDNA and 18S rDNA gene copy numbers per g of sediment were calculated from the triplicate average of each sample as described by Sylvan et al. (2013). Overall Prokaryotic cell abundance per gram of sediment was estimated by taking into account the average of the 16S rRNA gene per cell equivalent to 1.86 for Archaea, 4.1 for bacteria (Lee et al., 2009) as previously used by Sylvan et al. (2013) and 2.18 for cyanobacteria (after calculation of the average using the data from Schirrmeister et al., 2012). Due to lack of references from lake sediments, one copy per cell of the mcrA gene was used to quantify the population of organisms involved in anaerobic methane cycling.

The raw data of eukarya, diatoms, and Botryococcus sp. were not further quantified into copy numbers per g sediment, for two reasons: (1) there is high variability in 18S rRNA gene copies per cell within the eukarya and diatoms (i.e. from 3 to more than 25000 copies per cell in the plants, Prokopowich et al., 2003; Zhu et al., 2005) and between 61 to 36 896 for the diatoms (Godhe et al., 2008); (2) the paucity of information related to the number of 18S rRNA gene copy number in the Botryococcus sp. genome. Therefore, the results reported here are indicative from the universal eukarya primer and should be considered as relative abundance of the total eukarya due to the tendency of not detecting all eukaryotic groups. Yet, the data are still useful to depict trends in the sediment record (see Supplement for limitations of qPCR methods).

3 Results

The $^{210}$Pb activity shows an exponential decay curve with depth, i.e. it shows a decreasing linear trend when plotted on a log-scale ($\ln (^{210}$Pb$_{\text{unsupported}}$ vs. depth; $r^2 = 0.827$) (Fig. 2). The profile indicates minimal sediment bioturbation and is used to calculate an average sedimentation rate of about 4.7 mm yr$^{-1}$.

Biogeochemical and biolipid screening of the sediment core, discussed further below, demarcates three distinct periods: ca. 2010–1969 CE, ca. 1969–1916 CE and ca. 1916–1857 CE. The sedimentary deposits are highly organic with TOC contents between 30 and 40%. TOC gradually increases from ca. 1857 to 1870 CE and then shows a decrease from ca. 1870 to 2008 CE (Fig. 3a). From ca. 1857 to 1970 CE both the $\delta^{13}$C$_{\text{bulk org}}$ (Fig. 3b) and TN (Fig. 3c) show a gradually increasing trend while $\delta^{15}$N$_{\text{org}}$ values rise steadily between ca. 1857 and 1969 CE and then decreases...
from ca. 1970 to 2010 CE (Fig. 3d). The C/N ratio, on the other hand, decreases gradually from the bottom to the top of the sediment core (Fig. 3e). Si/Ti ratio, a marker of silicon input into the lake from terrestrial sources, shows an increase between ca. 1916 and 1969 CE (Fig. 3f and g). Scanned photos using ESEM show higher abundances and diversity of diatoms between ca. 1916 and 1969 CE (see Supplement; Fig. S6). The P/Ti ratio, which can be used as a proxy for trophic conditions in lakes (Kirilova et al., 2011), show an increase from ca. 1970 to 2010 CE (Fig. 3g).

Biomarkers and qPCR analysis targeting cyanobacteria, diatoms and botryococcus exhibited similar trends, except ca. 1970–2010 CE where botryococcus was not detected although the qPCR analysis detected the presence of Botryococcus sp. (Fig. 4) (see supplement for detailed qPCR results). The botryococcene lipid concentrations show an increase between ca. 1857 and 1916 CE (Fig. 4a), followed by a decreasing trend from ca. 1916 to 2010 CE. The qPCR data set also shows an increase in Botryococcus sp. abundances from ca. 1857 to 1916 CE, a decrease from ca. 1916 to 1969 CE (Fig. 4b), but then again an increase from ca. 1970 to 2010 CE. The HBIs exhibit minimum levels until 1916, after which they show much higher abundance from ca. 1916 to 1969 CE, similar to the diatom abundance as detected by qPCR (Fig. 4c and d), and as seen by ESEM. After 1970 they decrease gradually. The C_{17} n-alkanes concentrations show a close resemblance to the Cyanobacteria sp. detected by qPCR, with rising concentrations and abundances between ca. 1970 and 2010 CE (Fig. 4e and f). The hydrogen isotopic composition of leaf waxes ($\delta D_{wax}$; weighted mean of $\delta D$...
C27–31 n-alkanes) shows a long-term oscillation over the entire 150-year record (Fig. 4g). δDwax values have been used as a proxy for rainfall intensity such that lower and higher values indicate an increase and decrease in rainfall intensity, respectively (e.g., Konecky et al., 2013; Niedermeyer et al., 2014). Reconstructed hydroclimatic conditions for southern Thailand over the last 150 years based on δDwax values show a relatively drier period from ca. 1857 to 1916 CE and ca. 1970–2010 CE and relatively wetter conditions from ca. 1916 to 1969 CE.

4 Discussion

The bulk geochemical trends (Fig. 3) indicate that multiple processes control the organic matter (OM) input into Lake NTP. Decreasing C/N ratios with TOC contents of approximately 30% from the bottom to the top of the core suggests organic matter diagenesis, which indicates an aquatic dominated system (Fig. 3a and b). The observed decreasing trend of the C/N ratio, as well as increasing δ13Corg values, may be explained by a preferential (anoxic) mineralisation of nitrogen-rich OM, leading to residual OM with a higher C/N ratio (Emerson and Hedges, 2003). This explanation would be consistent with the long-term diagenetic OM transformation (Sun et al., 2004). However, successional deposition of different phytoplankton communities, botryococcus (ca. 1857–1916), diatoms (ca. 1916–1969) and cyanobacteria (ca. 1970–2010), with different C/N ratios, δ13Corg, and δ15Norg can also influence the bulk profiles.

An alternate interpretation of the increasing δ13Corg values from the bottom to the top of the core lies in a successional shift of the trophic status of the lake (Brenner et al., 2000; Meyers and Teranes, 2001). Increasing nutrient levels promote primary productivity, thereby depleting the amount of dissolved inorganic carbon. This could lead to lower net fractionation against 13C by the phytoplankton during photosynthesis, thereby increasing the δ13Corg values (Meyers and Teranes, 2001).

Observed variations of dominant phytoplankton community over time can be explained best by changes in the trophic level of the lake. The period from ca. 1857 to 1916 CE is marked by significant increases in both Botryococcus sp. abundance (Fig. 4a) and botryococccene lipids (Fig. 4b), which also corresponds with slightly lower precipitation (Fig. 4g). Botryococcus braunii has been suggested to thrive in relatively oligotrophic conditions (Souza et al., 2008) and can, therefore, be used as a proxy for such conditions within the oxygenated epilimnion (Waldmann et al., 2014). Although the highly organic sediment may suggest a productive lake, at variance with oligotrophic lake water conditions, we argue that the final TOC content in lake sediments, especially in the tropics, is more determined by OM preservation than by the incoming flux. In the tropics, the mean air temperature (MAT) is a direct result of incoming solar radiation, and the relationship between the MAT and the amount of rainfall are typically inversely proportional (Imboden and Wüst, 1995; Boehler and Schulze, 2008). Dry, cloudless and warmer conditions lead to stronger stratification in freshwater lakes (Imboden and Wüst, 1995). We hypothesise that a long-term decrease in rainfall and slightly higher temperatures would result in a thermal stratification of the lake. Under stratified conditions reformed nutrients would remain in the anoxic hypolimnion. As a result, the sur-
face water remained relatively oligotrophic enabling *Botryococcus braunii* to thrive, whereas the bottom of the lake was characterized by bacterial breakdown of organic N. This argument is supported by increasing δ\(^{15}\)N\(_{\text{org}}\) values from ca. 1857 to 1916 CE, which suggests extensive denitrification in the lake sediment, as denitrification typically leads to more depleted δ\(^{15}\)N values in the residual organic matter (Wada, 1980).

A stark difference in the dominant phytoplankton community is observed from ca. 1916 to 1969 CE. This period is marked by significant decrease in *Botryococcus* sp. gene abundance (Fig. 4a) and botryococcene lipid concentrations (Fig. 4b) while diatom abundance (Fig. 4c) and C\(_{25}\) HBIs concentrations (Fig. 4d), which is a useful indicator of diatom-derived OM inputs to sediments (McKirdy et al., 2013) increases. More directly, we observed a marked increase in diatom diversity on the ESEM image scans of the sediments (Supplement; Fig. S6). Diatoms dominate phytoplankton communities as long as there is abundant silica irrespective of changes in environmental conditions and nutrient levels (Egge and Aksnes, 1992). Interestingly, the increase in diatom markers (Fig. 4c and d) coincides with an increase in reconstructed rainfall intensity (Fig. 4g). Moreover, the increase in Si / Ti ratios (Fig. 3f), a run-off signal (Murphy et al., 2000), coincides with high diatom blooms, in particular between ca. 1916 and 1969 CE. Since Ti is a highly immobile element, weathering and transportation of Si is not accompanied by significant Ti delivery to aquatic basins. Therefore, the Si / Ti ratio can serve as a proxy for nutrient dynamics linked to hydrological changes (e.g. Cartapanis et al., 2014) and as an indicator for enhanced diatom production in lakes (Wennrich et al., 2014). Altogether, it appears that the wetter conditions between ca. 1916 and 1969 CE increased catchment runoff into the lake, leading to elevated nutrient and silicate mineral flux to the lake water (e.g. Paerl and Fulton III, 2006). In line with the argument above, it is also well possible that an increase in precipitation was accompanied by cooler temperatures, which then led to a decrease in thermal stratification and an increase in mixing between the epilimnion and hypolimnion, making reformed nutrients from the bottom waters available at the lake water surface.

After ca. 1969 CE, the phytoplankton community structure changed again, with a diminishing role for diatoms as evidenced by lower concentrations of C\(_{25}\) HBIs (Fig. 4c) and the start of a marked increase in cyanobacteria gene numbers (Fig. 4e) and C\(_{17}\) n-alkane concentrations (Fig. 4f). C\(_{17}\) n-alkanes are recognised biomarkers of aquatic algae and photosynthetic bacteria such as cyanobacteria (Meyers, 2003). Indeed, cyanobacteria gene numbers relative to bacterial quantification based on the qPCR data covery strongly with C\(_{17}\) n-alkane concentration, which indicates that the C\(_{17}\) n-alkanes were likely produced by cyanobacteria. Additionally, the decreasing δ\(^{15}\)N\(_{\text{bulk}\_\text{org}}\) values during this period (Fig. 3d) also suggest nitrogen fixation, a process strongly associated with cyanobacteria blooms (Vahtera et al., 2007).

The most recent period (ca. 1969–2010 CE) is also characterized by somewhat lower rainfall amounts (Fig. 4g). The amount of Si indeed decreased, and diatoms became less abundant, allowing non-diatomaceous phytoplankton not dependent on Si to take over (Egge and Aksnes, 1992).

The number of *Botryococcus* genes increased again (Fig. 4a). However, this was not accompanied by the production of botryococcene lipids. This mismatch can be the result of the primer used for the quantification of *Botryococcus braunii* not being specific enough. *Botryococcus braunii* are classified into three races: A, B, and L, and it is only the B race that produces botryococcenes (Eroglu et al., 2011), whereas the A and L races produce long-chain alkadienes and lycopadienes, respectively (Metzger and Largeau, 2005).

A change in race thus may explain the disappearance of botryococcenes from the upper part of the sediment record. Since the qPCR primers for *Botryococcus* were not specific enough, it is also possible that it picked up closely related green algae that took over the ecological niche of *B. braunii* under different trophic conditions. The absence of long-chain alkadienes and lycopadienes in the upper sediment layers supports this argument and indicate that conditions had indeed become unfavourable to *B. braunii*, in a similar way as described by Smittenberg et al. (2005).

Phosphorus bioavailability is one of the most important factors limiting aquatic cyanobacteria blooms (Paerl and Fulton III, 2006; Paerl and Valerie, 2012), and the shift to cyanobacterial prevalence thus suggests eutrophic phosphorus-rich conditions, instead of the oligotrophic-like conditions that occurred a century earlier under otherwise similar climatic conditions. Indeed, higher levels of phosphorus were detected in the upper part of the sediment (Fig. 3g). The source of the elevated phosphorus is unclear but has likely resulted from human activities. Under a land development program in the 1990s, more than 20% of Thailand’s 56,000 villages were located in forest reserves (Gray, 1991; Puri, 2006), which allowed the expansion of land encroachment and agricultural activities. For instance, southern Thailand has seen an increase in the cultivation of rubber trees on small farms at rates above 7 % yr\(^{-1}\) (Leturque and Swigings, 2011). The use of fertilizers during farming activities, untreated wastewater effluents and the use of detergents are likely sources of the elevated phosphorus inputs into the lake (Litke, 1999; Chislock et al., 2013). These accelerated the eutrophic state of the lake beyond the natural rate of nutrient enrichment, which takes centuries to achieve (Litke, 1999).

According to our analysis, photosynthetic primary production is at the base of internal organic matter production in the lake. However, changes in precipitation and anthropogenic nutrient input have produced fluctuations in dominant primary producer communities over the last 150 years, from botryococcene lipid-producing algae to diatoms to a dominance of cyanobacteria. Overall, we suggest that NTP is sensitive to environmental stressors and that multiple processes discussed above control the OM input and carbon storage.
The bulk geochemical analysis was clearly inadequate to disentangle the processes that have been controlling the limnological, ecological and microbial dynamics in NTP. Instead, the combination of bulk geochemistry, lipid biomarkers, and molecular DNA analysis could appropriately constrain many of these processes.

5 Summary and conclusion

Validation of paleoenvironmental and paleoclimatic proxies is necessary to constrain geochemical patterns through time. The coupled lipid biomarker and qPCR data allowed different microbial groups in the lake sediments to be distinguished and quantified, leading to the identification of meaningful biological relationships between the phytoplankton community structure response to either anthropogenic or natural environmental changes over the last 150 years.

Our results show that between ca. 1857 and 1916 CE, relatively drier climate in southern Thailand coincided with relatively oligotrophic surface water conditions in Lake NTP, which allowed Botryococcus braunii species to bloom. From ca. 1916 to 1969 CE, an increase in precipitation resulted in higher Si input into the lake, which led to a rapid takeover by diatoms as primary producers. Since the 1970s, many aspects of the initial limnic state returned upon drier conditions, except that anthropogenic impact led to an increase in phosphorus thereby allowing cyanobacteria to become a major contributor to primary productivity. Although the qPCR method did again detect genetic evidence for the presence of Botryococcus sp. over the last decades, the specific lipid biomarkers for Botryococcus sp. were not found anymore, which can be due to limitations of either proxy.

We conclude that the combination of geochemical and lipid biomarker proxies together with qPCR analyses is a useful approach that has the potential to assist in tracking the effects of changing climate on primary producers and also to assess these effects on the carbon cycle. This multi-proxy approach may help refine the knowledge about the use and shortcomings of the different proxies, which is critical for their interpretation especially when used on a more stand-alone basis.

Author contributions. This study was conceived and led by K. A. Yamoah, E. Chi Fru and R. H. Smittenberg. K. A. Yamoah, N. Callac, A. Wiech and A. Chabangborn carried out laboratory analyses. K. A. Yamoah, N. Callac, E. Chi Fru, B. Wohlfarth and R. H. Smittenberg wrote the manuscript. All authors discussed the results and their implications and commented on the manuscript as it progressed.

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