Blooms of cyanobacteria in a temperate Australian lagoon system post and prior to European settlement

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Abstract

Blooms of noxious N$_2$ fixing cyanobacteria such as *Nodularia spumigena* are a recurring problem in some estuaries. Here we report the results of a palaeoecological study on a temperate Australian lagoon system (The Gippsland Lakes) where we used stable isotopes and pigment biomarkers in dated cores as proxies for eutrophication and blooms of cyanobacteria. Pigment proxies show a clear signal, with an increase in cyanobacterial pigments (echinenone, canthaxanthin and zeaxanthin) in the period coinciding with recent blooms. Another excursion in these proxies was observed prior to the opening of an artificial entrance to the lakes in 1889, which markedly increased the salinity of the Gippsland Lakes. A coincident increase in the sediment organic carbon content in the period prior to the opening of the artificial entrance suggests the bottom waters of the lakes were increasingly stratified and hypoxic, which would have led to an increase in the recycling of phosphorus. After the opening of the artificial entrance there was a ~ 60 year period with low values for the cyanobacterial proxies as well as a low sediment organic carbon content suggesting a period of low bloom activity associated with the increased salinity of the lakes. During the 1940s, the current period of re-eutrophication commenced as indicated by a steadily increasing sediment organic carbon content and cyanobacterial pigments. We suggest increasing nitrogen inputs from the catchment led to the return of hypoxia and increased phosphorus release from the sediment, which drove the re-emergence of cyanobacterial blooms.

1 Introduction

Harmful algal blooms (HABS) are becoming increasingly prevalent throughout the world. One of the key causes of this is eutrophication of aquatic environments by excessive nutrient inputs (Conley et al., 2009). Climatological and hydrological factors are increasingly recognized as an important contributor to HABS through altered temperature and salinity regimes (Paerl and Paul, 2012). Blooms of toxic cyanobacteria...
such as *Nodularia spumigena* are particularly conspicuous in some estuaries such as the Baltic Sea, the Peel-Harvey Estuary and the Gippsland Lakes (Cook and Holland, 2012; Lukatelich and McComb, 1986; Conley et al., 2009). The most likely reasons for their dominance in these systems are: (1) long water residence time, (2) intermediate salinity (∼ 5–20); and (3) a high supply of phosphorus (both from the catchment and from anoxic bottom waters and sediments). As such, the frequency and occurrence of these blooms is likely to result from a strong interaction between anthropogenic nutrient loading and climatological and hydrological factors.

In the case of the Baltic Sea, cyanobacterial blooms have occurred sporadically since the formation of the Littorina Sea 8000 years BP (Bianchi et al., 2000). The presence of cyanobacteria is most likely controlled by the extent of bottom water hypoxia, which leads to an efficient recycling of phosphorus (Funkey et al., 2014). The extent of hypoxia in the Baltic has been controlled by morphological and hydrological changes, however, the most likely control over hypoxia and cyanobacterial blooms over the past two millennia is the expansion and contraction of human activities (Zillen and Conley, 2010). Similarly, in the Gippsland Lakes and Peel-Harvey Estuary, the frequent and intense blooms are thought to be relatively recent phenomena, with significant blooms only observed after the late 1970s (McComb and Humphries, 1992). In the Peel-Harvey Estuary, the intensity of *N. spumigena* blooms is strongly related to river discharge during the previous winter/spring which delivers significant quantities of phosphorus as well as reducing the salinity of the estuary to a range favourable to *N. spumigena* growth (McComb and Humphries, 1992). The critical importance of salinity in controlling *N. spumigena* blooms is well illustrated in the Peel-Harvey Estuary, where a newly cut channel to the sea increased the salinity of the estuary and virtually eliminated *N. spumigena* blooms (Wildsmith et al., 2009).

In the Gippsland Lakes, Australia, it has been shown that *N. spumigena* bloom size is decoupled from catchment inputs owing to internal recycling of P driven by stratification (Cook and Holland, 2012). As such, *N. spumigena* blooms typically occur during wet years when stratification is highest, however there is no relationship be-
tween catchment nutrient loads and bloom size (Cook and Holland, 2012). Nevertheless, anthropogenic activities are likely to have played a role in the occurrence of recent blooms through increased phosphorus inputs leading to increased sediment phosphorus storage as well as increased nitrogen inputs which are likely to drive increased releases of phosphorus from the sediment through increased anoxia (Cook et al., 2010).

Prior to European settlement, the Gippsland Lakes were connected to the ocean by an ephemeral entrance (Bird, 1978). In 1889, a permanent artificial entrance was opened, which increased the salinity regime of the lakes (Saunders et al., 2008). There are anecdotal accounts of *N. spumigena* blooms prior to the opening of the artificial entrance, but there is no information on the relative frequency or intensity of blooms at this time. The Gippsland Lakes provide an ideal case study of global relevance to investigate the relative importance of salinity and nutrient regimes in driving cyanobacterial blooms. Two alternative hypotheses were tested here. (1) the fresher, less flushed and more stratified environment prior to the opening of the artificial entrance may have been more conducive to anoxia, associated sediment phosphorus release and cyanobacterial blooms than post opening. (2) Alternatively, low nutrient inputs prior to European settlement may have led to a lower incidence of hypoxia and associated cyanobacterial blooms. The aim of this study was to investigate changes in the trophic status and frequency of cyanobacterial blooms in the Gippsland Lakes before the opening of the artificial entrance up to the present day using pigment, organic matter and isotope proxies on dated cores taken from the centre of the lake system. The findings provide an important, longer-term perspective from which to frame modern management regimes within the Gippsland Lakes as well as other systems modified by humans more generally.
2 Materials and methods

2.1 Study site

The Gippsland Lakes are located in South Eastern Australia, and experience a temperate climate with a water temperature range of \( \sim 10–25^\circ \text{C} \) (Fig. 1). The Lakes are fed by 5 river systems including the Latrobe and Avon in the west, and the Mitchell, Tambo and Nicholson in the east. Lake Wellington in the west is a shallow basin with an average depth of \( \sim 2.6 \text{ m} \), and typically has a salinity < 15. Lakes King and Victoria in the east have an average depth of \( \sim 5 \text{ m} \). The lakes cover an area of 356 km\(^2\), making them one of the largest estuarine systems in Australia. Maximum river flows and floods typically occur in the Austral winter–spring, which reduce surface salinities to \( \sim 5–15 \text{, which then increase to } > 25 \text{ over summer in Lakes King and Victoria. Lakes King and Victoria are typically salinity stratified, with bottom water salinities of } \sim 30–35 \text{ and during intense stratification following high river flow, the bottom waters of the Lakes King and Victoria may become anoxic. Winter and spring inflows typically lead to blooms of diatoms and dinoflagellates, and since 1987, periodic blooms of } Nodularia spumigena \text{ have occurred in Lake King during late spring and summer when surface water salinities are } \sim 9–20 \text{ (Cook and Holland, 2012). Previous studies have shown these blooms are phosphorus limited and that they are sustained by high sediment phosphorus release focused in the northern basin of Lake King and that these blooms can fix significant quantities of nitrogen with a } \delta^{15}\text{N of } \sim 0 \text{‰ (Cook and Holland, 2012; Cook et al., 2010; Holland et al., 2012; Woodland and Cook, 2014; Woodland et al., 2013).}

2.2 Sampling

Sediment cores were taken from Lake King North (LKN; 37.875620° S, 147.757280° E, Fig. 1) at a water depth of 7 m on the 15 March 2012. The uppermost core, LKN1 (0–56 cm) was retrieved by a piston corer to collect the recent, unconsolidated sedi-
ments. This core was sectioned in the field at 0.5 cm intervals (contiguous), with a view to attaining a chronology to aid the identification of recent changes in the sediment. Subsamples (1–2 g) from each section were stored in glass vials, with the rest of the sample stored in zip lock bags. All samples were kept on ice and, on returning to the laboratory, the bags were transferred to the refrigerator (4 °C) and the vials frozen until required for stable isotope analysis. Wet samples were kept in darkness in order to reduce light exposure that could change the sediment composition.

A second drive (LKN2) using a Russian peat corer collected a deeper, older sedimentary sequence and consisted of a series of 50 cm (overlapping) drives. All cores were stored in halved PVC pipes, wrapped in cling film and aluminium foil and kept cool until refrigerated in the laboratory. The cores were sectioned into 1 cm layers using a blade and spatula, and stored as per LKN1. The two sequences were correlated based on the calculated field depths and this was validated by dating across the two sequences as described in the next section.

2.3 Dating

The sediment core was dated at the Australian Nuclear Science and Technology Organisation (ANSTO) Institute for Environmental Research using the lead-210 ($^{210}$Pb) dating method (Appleby, 2001). Samples were chemically processed following the methods described in Atahan et al. (2015) and analysed by alpha spectrometry to determine unsupported $^{210}$Pb activities on thirteen subsamples from core LKN1 (0–51.5 cm) and the 42–92 cm LKN2 sequence. A modified CIC (Constant Initial Concentration) model was used to calculate the ages of the sediment samples (Appleby, 2001). The $^{210}$Pb chronology was validated with the presence of a subsurface peak of caesium-137 ($^{137}$Cs), which identifies the year of 1964, due to global atmospheric nuclear weapons tests (Leslie and Hancock, 2008). $^{137}$Cs activities in 8 subsamples were determined by gamma spectrometry.
2.4 Charcoal

Wet sediment (1 mL) from the LKN2 sequence was subsampled into a 50 mL Falcon tube. 25 mL of 10% tetra sodium pyrophosphate (Na$_4$P$_2$O$_7$) was added to the tube, the contents shaken vigorously and left to sit. After 30 min, 25 mL of 12.5% sodium hypochlorite (NaOCl) was added, and the tube was again shaken vigorously and then left to sit for 14–18 h. The samples were then sieved through 250 µm and then 125 µm mesh, rinsed and placed on a water filled petri dish where the total number of charcoal and grass charcoal particles were enumerated under a dissecting microscope.

2.5 Diatoms

The LKN 2 core sequence was subsampled every 10 cm for diatom analysis. Approximately 1 g wet weight sediment was digested in 30% hydrogen peroxide in a beaker, on a hotplate, for up to 4 h (Battarbee, 1986). Following digestion, a small amount of HCl was added to remove any carbonates. The suspensions were washed in distilled water and left to settle overnight, before decanting the supernatant (repeated four times). An aliquot of the final suspension was placed onto a coverslip and left to dry. The coverslips were permanently mounted onto slides using Naphrax. Diatoms were identified (where possible) to species level, using a Nikon DIC Microscope. Identifications were undertaken using a range of general (Krammer and Lange-Bertalot, 1991a, b; Krammer and Lange-Bertalot, 1988, 1986) and regional (Foged, 1978; Sonneman et al., 2000) floras. A minimum of 200 valves per sample were counted, and the counts converted to percentage data in C2 (Juggins, 2003). Where possible an ecological preference (i.e. saline, fresh, thalassic) was assigned to each species to create a habitat summary diagram. To further explore patterns in the diatom data, a Detrended Correspondence Analysis (DCA) was carried out using Canoco 4.5.
2.6 Carbon and nitrogen analysis

Sediment from the LKN1 and LKN2 core sediment sample was analysed via mass spectroscopy for % nitrogen, % carbon, C\textsubscript{org}: N, $\delta^{15}$N and $\delta^{13}$C\textsubscript{org}. Samples were dried at 60°C for 30–50 h and placed in 1.7 mL Eppendorf tubes along with Qiagen Tungsten Carbide Beads (3 mm). Samples were shaken for 6–10 min at 25 Hz using a Retsch Mixer Mill MM 200 until a fine, homogeneous powder was produced. Samples for carbon ($\delta^{13}$C\textsubscript{org}) were weighed in silver capsules and placed on a hotplate (60–80°C) to undergo acidification. Aliquots (20 µL) of 10 % HCl were sequentially added to capsules until no effervescence was recorded. Samples for nitrogen ($\delta^{15}$N) analysis were weighed in tin capsules. Once each capsule was prepared it was pinched-closed and pressed into a disk using a pelletiser. Each sample was analysed on an ANCA GSL2 elemental analyser interfaced to a Hydra 20–22 continuous-flow isotope ratio mass-spectrometer (Sercon Ltd., UK). Stable isotope data were expressed in the delta notation ($\delta^{13}$C\textsubscript{org} and $\delta^{15}$N) relative to the stable isotopic ratio of Vienna Pee Dee Belemnite standard ($R_{VPDB} = 0.0111797$) and the air standard ($R_{Air} = 0.0036765$), for carbon and nitrogen respectively. Analytical precision was ±0.1 ‰ for both $\delta^{13}$C\textsubscript{org} and $\delta^{15}$N (SD for $n = 5$).

2.7 Pigments

Pigments were analysed at 5 cm intervals from 0–41 cm and every 10 cm through to 200 cm from the LKN2 core sequence. Freeze-dried sediments were extracted in pure acetone overnight and stored in the dark at −22°C. They were then filtered, dried and re-dissolved, and then injected into a Shimadzu high performance liquid chromatography (HPLC) system. The separation conditions were modified from Mantoura and Llewellyn (1983) and Chen et al. (2001) using a 4.6 mm × 150 mm, 3 µm C8 (Luna, Phenomenex) column. Pigment peaks were identified by retention times and spectra, and then quantified by peak areas at maximum absorbance wavelength using calibrated curves from phytoplankton pigment standards DHI (Denmark). Canthaxanthin
was measured at 475 nm, and the carotenoids lutein, zeaxanthin, diadinoxanthin, dia-
toxanthin, and echinenone were measured at 450 nm. The pigments zeaxanthin, echi-
enone and canxanthin were used as markers for cyanobacteria (Jeffrey and Vesk, 1997), Chlorophyll *a* was measured at 665 nm. Concentrations are reported in micro-
moles (µmol) of pigment relative to the organic material in the sediment as measured
described above.

### 3 Results

#### 3.1 Age model

The unsupported $^{210}$Pb activities for the King Lake core exhibit an overall decay profile with increasing depth (Fig. 2a) indicating the core is suitable for $^{210}$Pb dating. The top 5 cm unsupported $^{210}$Pb activities deviate from a decay profile, which may be due to sediment mixing in the upper section of the core. Below 5 cm the unsupported $^{210}$Pb activities exhibit two distinct zones, where each zone follows a monotonic profile with depth, between 5–20 and 20–90 cm. Using the CIC $^{210}$Pb dating model, the mass accumulation rate between 5–20 cm depth was calculated to be 1 g cm$^{-2}$ year$^{-1}$ (about 0.55 cm year$^{-1}$), and between 20–90 cm depth at 2.4 g cm$^{-2}$ year$^{-1}$ (about 1 cm year$^{-1}$). These mass accumulation rates were used to determine sediment ages between 0 and 90 cm core depth, which were then converted to calendar years (Fig. 2b). The $^{210}$Pb ages were validated by $^{137}$Cs, with the 1964 peak found in the 25–30 cm sediment layer in the LKN1 sequence, dated to 1964–1969 (Fig. 2c). A further validation is the peak in charcoal concentration (24 mL$^{-1}$) at depth 61 cm (Fig. 3), which corresponds to approximately 1937, which is in close agreement with the date of the most widespread bushfires in the catchment in 1939. The sediment age below 90 cm was estimated by extrapolating the measured sedimentation rate over the depth interval 20–90 to 200 cm. Given that sedimentation rates have increased since European settlement, this provides a minimum age for sediments below $\sim$ 140 cm.
3.2 Sediment chronology

Three broad zones within the chronology are identified here.

3.2.1 Zone LK1 (c. 1810–1870, depth 130–200 cm)

This zone covers the period of European settlement in the region. Sediment organic carbon and nitrogen content were relatively stable at 5 and 0.5 % respectively and were the highest observed in the entire record (Fig. 3). Sediment δ\(^{13}\)C and δ\(^{15}\)N were similarly stable at –23 and 4 % respectively. Sediment C\(_{\text{org}}\) : N ratios were ~ 10 over this period. Sediment pigments including diatoxanthin and cyanobacterial pigments were also sporadically high over this period, with total cyanobacterial pigments peaking at 2600 nmol g \(^{-1}\) at 185 cm coinciding with the year ~ 1820 which is the highest concentration in this record. Over this period, thalassic diatom taxa dominated, although to a much lesser extent than in the period LKN1. The diatom *Cyclotella choctawatcheeana* was the dominant species over this period. Charcoal concentrations were much higher over this period than the subsequent periods, and peaked at 170 cm coinciding with the year ~ 1825 (Fig. 4).

3.2.2 Zone LK2 (c. 1870–1920, depth 130–80 cm)

This zone represents an abrupt change in most proxies and encompasses the period of the opening of the artificial entrance in 1889. The sediment δ\(^{13}\)C, %C\(_{\text{org}}\) and %N decreased by 1.5‰, 2 and 0.2 % respectively at 130 cm before increasing again by a similar amount at 80 cm. There was also a distinct jump in the C\(_{\text{org}}\) : N ratio in the sediment from ~ 11 to > 13 over this period. Sediment δ\(^{15}\)N increased abruptly by ~ 1 ‰ at the start of the period, but unlike the other proxies it did not show and marked change at the end of this period as the other proxies did. There was a distinct peak in the dominance of freshwater diatom species over this period. All pigment markers were
relatively low and stable over this period. There was a spike in charcoal at ∼ 100 cm depth coinciding with the year ∼ 1900.

3.2.3 Zone LK3 (c. 1920–2012, depth 0–80 cm)

Over this period, %N and %C\textsubscript{org} increased from approximately 0.2 to 0.5 and 2.6 to 4.3, respectively. The C\textsubscript{org} : N ratio showed an initial decrease and then stabilised at ∼ 11 (Fig. 3). The δ\textsuperscript{13}C and δ\textsuperscript{15}N were relatively constant at −23 and 5 ‰ respectively over the period 1925–1980s; and the sedimentation rate was also constant at 1 cm\textperyear\textsuperscript{−1}. During the late 1980s, the sedimentation rate slowed from 1 to 0.55 cm\textperyear\textsuperscript{−1}, at the same time there was a marked decrease in δ\textsuperscript{15}N, δ\textsuperscript{13}C\textsubscript{org}, %N and %C\textsubscript{org}, and a spike in C\textsubscript{org} : N. This change coincided with the first and largest \textit{Nodularia spumigena} bloom on record in Lake King in 1987, followed by recurring blooms through to the present (Cook and Holland, 2012). In 2006–2007 major bushfires occurred in the East Gippsland catchment, followed by a major flood in 2007.

Concentrations of chlorophyll \textit{a}, diatoxanthin and cyanobacterial pigments increased gradually from 1925 to the late 1980s, before rapidly increasing after this period. Pheophytin \textit{a} showed sporadic peaks between 1950 and 1975, before decreasing in the 1980s, followed by another increase after this. Diatoms were dominated by thalassic taxa throughout the period 1925 to the present. Charcoal concentrations within the sediment were consistently low throughout this period with one spike occurring at ∼ 60 cm which dates to the period of ∼ 1939 coincident with some of the most widespread fires on record in the region.

4 Discussion

Prior to European settlement in the early 1840s, land use by the Aboriginal tribes was of a nomadic hunter gathering nature, and documentary evidence suggests that fire was the principal agent of land management across Australia at this time (Gammage,
This account is however contradicted by most charcoal records from south-east Australia which show an increased incidence of fire after European settlement (Mooney et al., 2011; Mills et al., 2013). The high charcoal counts below 170 cm are consistent with high rates of indigenous burning of fringing reedbeds before tubers were harvested which has been a recognized common practice (Head, 1987). Early European land use was primarily low intensity sheep grazing. Gold mining commenced in the 1850s, followed by increased navigation of the lakes in the 1860s for the purposes of trade, fishing and tourism. By the 1870s there was regular steamer traffic on the Mitchell River and there are regular references to dredging the mouths of the Mitchell, Nicholson and Tambo Rivers from the early 1880s though to the turn of the century and into the 1920s (Synan, 1989). The opening of the permanent entrance in 1889 was one of the pivotal moments in the recent ecological history of the lakes because it led to an increase in the salinity of the Gippsland Lakes (Saunders et al., 2008). Over the period of the opening of the artificial entrance (corresponding to depths of ~ 110 cm), we expected to see a change in the diatom taxa to a greater abundance of thalassic species and a concomitant reduction in freshwater species. Unexpectedly, a spike in freshwater species was instead observed over the period 1870–1925 represented by the LK2 layer. This corresponded with other geochemical proxies which suggested an increase in terrestrial organic matter including a decrease in δ^{13}C, and an increase in the sediment C : N ratio which was also observed (although not discussed) by Saunders et al. (2008). This suggests that the study site had an increased influence from riverine sediments over this period. The most likely explanation for this is remobilization of terrestrial sediments through dredging activities within the delta of the Mitchell River over this period (Fig. 1), which could have led to an increased deposition of terrestrial material at the study site over this period. It appears that this increased input of terrestrial material did not correspond with a changed sedimentation rate as the {superscript 210}Pb decay profiles displayed a similar trend between the LK3 and LK2 layers (Fig. 2a). Irrespective of the exact cause of the LK2 sediment layer, we are confident the LK3 and LK1 layers are representative of post and pre artificial entrance opening periods respectively.
The biogeochemical proxies analysed here provide clear evidence for two periods of eutrophication and cyanobacterial blooms in the Gippsland Lakes: (1) The recent period after the Second World War (LK3) and (2) prior to the opening of the artificial entrance in 1889 (LK1). The latter part of the most recent period has been well monitored and provides an excellent means to calibrate the biogeochemical proxies. The first piece of evidence for the recent period of eutrophication comes from the steady increase in sediment organic carbon content after the 1940s (Fig. 3), consistent with a previous paleolimnological study (Saunders et al., 2008). The $\delta^{13}C_{\text{org}}$ of this organic matter is typically $\sim -23\%$, consistent with organic matter inputs derived from phytoplankton (Fig. 3). This period also coincided with a marked jump in the sum of the cyanobacteria pigments zeaxanthin, echinenone and canthaxanthin from $\sim 500 \text{ nmol g}^{-1} \text{C}_{\text{org}}$ up to $>2000 \text{ nmol g}^{-1} \text{C}_{\text{org}}$ at the top of the core (Fig. 3) consistent with a *N. spumigena* bloom at the site in November 2011–February 2012 (Woodland et al., 2013). The first documented bloom of *N. spumigena* in the lakes occurred in 1965 and the period from 1987 through the 1990s is known to have had severe and regular blooms (Cook and Holland, 2012). Over this period there were also two dips in the $\delta^{15}N$ of $\sim 2\%$ in the 1940s and late 1980s–2000 consistent with the occurrence of nitrogen fixing cyanobacteria. The broad agreement between these cyanobacteria markers and recent recorded blooms gives us confidence that they are appropriate markers of cyanobacteria blooms within the Gippsland Lakes and this is consistent with previous studies (Bianchi et al., 2000).

The biogeochemical proxies for the period prior to the opening of the artificial entrance in 1889 in layer LK1 likewise suggest a period of eutrophication and intense cyanobacteria blooms. The sediment organic content was high ($\sim 5\%$), the $\delta^{15}N$ was low ($\sim 4–5\%$), the $\delta^{13}C_{\text{org}}$ was in the range typical of phytoplankton ($-22$ to $-23\%$) and the cyanobacteria pigments were sporadically high (Fig. 3). Reports from newspaper articles in the late 1870s also suggest the presence of surface scums of cyanobacteria with reference to “noxious and ill smelling weed” on the surface of Lake King and there were anecdotal reports of the greatly improved water quality with the increased
salinity after the opening of the artificial entrance in 1889 (Synan, 1989). We now discuss 3 key factors controlling incidence of cyanobacteria blooms prior to the opening of the artificial entrance and European settlement.

1. Salinity. With no permanent entrance, the inflow of seawater was greatly reduced, and at this time the lakes were considerably fresher (Harris et al., 1988; Saunders et al., 2008). The diatom chronology also supports this reduced marine influence with an increased abundance of *Cyclotella choctawatcheeana* a planktonic diatom characteristic of deep mesosaline (salinities > 10) lakes and brackish marine systems (Fritz et al., 1993), and the reduced dominance of thalassic diatoms (Fig. 4). *N. spumigena* typically blooms at salinities between 9 to 20 in the Gippsland Lakes, and these salinities are currently only reached during late spring-summer in high flow years (Cook and Holland, 2012). Prior to the opening of the artificial entrance it is likely this salinity range was more typical, hence increasing the frequency of blooms. In high flow years when salinities were even lower, it is likely that other nitrogen fixing cyanobacteria, such as *Anabaena* would instead dominate. This species occasionally blooms in Lake Wellington, and it is therefore possible that this cyanobacterium was present in Lake King prior to 1889.

2. Stratification and residence time. At present, the highest stratification is observed during periods of low surface water salinity in the Gippsland Lakes associated with above average river flows (Cook and Holland, 2012). Reduced tidal flushing, combined with low surface salinity would lead to enhanced stratification and residence time of the water column which are both known to favour buoyant slow growing cyanobacteria such as *N. spumigena* (Sellner, 1997; Paerl, 2014). Increased stratification will also lead to increased hypoxia, and the marked increase in sediment organic carbon content to ~ 5% prior to ~ 1870 (below 130 cm, Fig. 4) is consistent with increased hypoxia in this period (Zillen and Conley, 2010). A key effect of this would be to enhance the release of phosphorus from the sediment which
is a key driver of *N. spumigena* blooms in the Gippsland Lakes (Cook et al., 2010; Scicluna et al., 2015).

3. Nutrients. Prior to European settlement it has been estimated that nitrogen and phosphorus loads were a lower by a factor of 2 and 3 respectively (Grayson et al., 2001). At face value, it is surprising that the Gippsland Lakes experienced more blooms of cyanobacteria then, however, this can be reconciled with contemporary studies. Firstly, cyanobacteria such as *N. spumigena* can derive all of their nitrogen requirements from nitrogen fixation, and these blooms can add significant loads to the Gippsland Lakes (Woodland and Cook, 2014). The generally lower sediment $\delta^{15}$N values prior to 1870 (below 130 cm) are consistent with this. Secondly, cyanobacterial blooms are driven by a focused release of phosphorus from the sediments during bottom water hypoxia/anoxia, and it was estimated that a large recent bloom of *N. spumigena* was caused by a release of $\sim$25 t of phosphorus from the sediment in Lake King (Scicluna et al., 2015). Given that phosphorus is trapped and effectively recycled in periodically anoxic and high residence time systems such as the Gippsland Lakes, it is plausible that a pre-European phosphorus catchment load of 50 t$\text{year}^{-1}$ could maintain blooms of at least the same magnitude as currently observed (Grayson et al., 2001).

Following the opening of the artificial entrance there was a $\sim$60 year period ($\sim$80–130 cm) with relatively low cyanobacteria biomass and oxic bottom water, as indicated by the reduction in cyanobacteria pigments and sediment organic carbon content respectively. It is likely that this relatively low productivity period was caused by relatively good ventilation of the bottom water combined with low catchment nutrient inputs. After the 1940s, the modern eutrophication of the Gippsland Lakes commenced as indicated by a steady increase in sediment organic carbon and cyanobacteria pigments. Changes to hydrological, morphological and salinity regimes are unlikely to be a key driver because, apart from a small reduction in river inputs from river diversions ($<20\%$ of total flow), there have been no significant hydrological and morphological
changes to the Gippsland Lakes over this period. Given that fire can lead to increased nutrient loads as previously discussed, it is highly likely that the 1939 wildfires led to a significant pulse of nutrients into the Gippsland Lakes. The subsequent increase in agriculture (Lake Glenmaggie, used for irrigation in the Macalister Irrigation District, a tributary of the Latrobe River, was completed in 1926 and expanded immediately post World War II), industry and urbanization within the catchment have been estimated to have increased nutrient loads by a factor of 1.8 and 3 for total nitrogen and phosphorus respectively (Grayson et al., 2001).

Previous work has shown that the Gippsland Lakes are generally nitrogen limited outside *N. spumigena* blooms (Holland et al., 2012) and increased inputs of this element have most likely resulted in increased productivity in the Gippsland Lakes, particularly during the winter and spring diatom blooms when most of the nutrient load is delivered. The increased settling of phyto-detritus after the collapse of these blooms would have driven increased water column anoxia over the late spring period triggering the release of phosphorus stored in the sediment leading to more favorable conditions for *N. spumigena* blooms (Cook et al., 2010; Scicluna et al., 2015; Holland et al., 2012). We therefore speculate that the recent re-emergence of cyanobacterial blooms is caused (or at least enhanced) by increased nitrogen loads, which drive increased internal release of phosphorus through increased bottom water hypoxia and anoxia in late spring through to summer. These observations are consistent with global studies of coastal waters which consistently show an increased incidence of hypoxia over the past 50 years driven by eutrophication (Diaz and Rosenberg, 2008) and that this may then lead on to blooms of cyanobacteria (Funkey et al., 2014). This finding supports the argument that mitigating coastal eutrophication requires controls on both nitrogen and phosphorus (Paerl, 2014; Conley et al., 2009), even in systems that experience diazotrophic cyanobacterial blooms.
5 Conclusions

In conclusion, blooms of cyanobacteria were a natural feature of the Gippsland Lakes prior to European settlement, most likely driven by strong stratification and phosphorus release from the sediment. We suggest that pre-European phosphorus loads were sufficient to maintain a sediment phosphorus pool capable of driving significant periodic blooms based on contemporary observations. The opening of the artificial entrance in 1889 likely led to increased salinity, flushing and reduced stratification, leading to an increase in bottom water oxygenation, a decrease in sediment phosphorus release and associated cyanobacterial blooms. The re-emergence of *N. spumigena* blooms post world war II may have occurred as a consequence of increased nitrogen inputs which led to increased anoxia occurring as a consequence of increased primary production, triggering sediment phosphorus release during the summer low flow period when blooms occur. This finding provides a mechanism by which decreasing nitrogen loads may also reduce phosphorus limited diazotrophic cyanobacterial blooms, by reducing phosphorus release from the sediment highlighting the need to control both nitrogen and phosphorus loads to estuaries even when they experience blooms of diazotrophic cyanobacteria.

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Figure 1. The Gippsland Lakes, south-eastern Australia. The cores were collected at in Northern Lake King marked with the solid circle.
Figure 2. Unsupported $^{210}$Pb activities (a), $^{137}$Cs activities (b) and the age depth model based on unsupported $^{210}$Pb values using the CIC model (c). The star in (c) refers to the depth of the $^{137}$Cs peak activity.
Figure 3. Depth profiles of diatom salinity indicator species (see Fig. 4 for classification of species) and geochemical proxies at the site Lake King North. Total cyanobacteria is the sum of the pigments, zeaxanthin, canthaxanthin and echineone.
Figure 4. Profiles of diatom species abundance grouped based on water salinity (fresh, saline and thalassic). Fresh indicates diatoms found at < 5 PSU, saline indicates species expected to grow at high salinity within estuaries, and thalassic species are expected to be found exclusively in the coastal ocean. Other refers to species typical of intermediate salinity within estuaries and lagoons.