Dynamics of nutrient and phytoplankton in Epe Lagoon, Nigeria: possible causes and consequences of reoccurring cyanobacterial blooms

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Abstract
The impact of environmental variables and processes of nutrient enrichment on phytoplankton community at Epe lagoon was studied for 18 months (November 2012–April 2014). Two cyanobacterial bloom incidences were observed in the two dry seasons during this period. Rainfall pattern regulated nitrogen limitation which triggered cyanobacterial bloom development and influenced the bloom duration. Nutrient-laden influx from agricultural practices accounted for the increase in nitrate, phosphate and the relatively low silicate. Hypoxic conditions during the two bloom episodes reflected the vulnerability of this lagoon ecosystem. The cyanobacterial blooms were dominated by potentially toxic species; Anabaena circinalis, A. flos-aquae, A. limnetica and A. spiroides. A comparison of ambient nutrient ratios with the Redfield ratio (N/P/Si = 16:1:16) showed clear temporal variations that coincided with phytoplankton dynamics and the bloom regime. Low N/P ratios were recorded during the first 8 months, and there was a shift to higher ratios during the subsequent 8 months, followed by low N/P ratios in the last 2 months. Higher silicate values coincided with decreased cyanobacterial biomass, cyanobacteria bloom collapse and proliferation of diatoms during the first annual cycle. Cyanobacterial abundance exceeded the alert level 1 and almost reached alert level 2 which call for continuous environmental monitoring and management of coastal waters.

Keywords Anabaena • Bloom • Cyanobacteria • Epe lagoon • Nutrient enrichment • Phytoplankton

Introduction
Phytoplankton form the base of aquatic food webs (Sommer 1989; Arrigo 2005) and are important biological component of marine ecosystems and biogeochemical cycles (Malone 1977; Litchman 2007; Litchman et al. 2015). Their excessive growth can cause significant threat to biodiversity and ecosystem functioning as in the case of toxic algal blooms (HABs) (Paerl and Huisman 2008). HABs occur naturally because of increased water temperature and inorganic nutrients (phosphate, ammonium and nitrate) (Carmichael 1997; Hitzfeld et al. 2000; Wetzel 2001). Global reports on algal blooms have increased because of human activities resulting from elevated nutrient loadings (Paerl and Whitall 1999; Van Dolah 2000; Anderson et al. 2002) and water flow modifications (Ferreira et al. 2005; Edwards et al. 2006). HABs result in food web alterations (Jackson 2001) and dominance of introduced species (Hallegraeff and Bolch 1992; Vanderploeg et al. 2001).

Toxic cyanobacterial blooms can potentially serve as signs or ecological indicators of changes in aquatic ecosystem integrity (Jeppesen et al. 1997; Sanchis et al. 2002; Williams et al. 2005; Carmichael 2008; Peretyatko et al. 2010). It is important to identify and understand what triggers bloom events to enable prediction, surveillance and management strategies for bloom occurrences. The relationships among hydrologic discharge (flushing and residence time), vertical/horizontal salinity and thermal gradients, rainfall and drought, wind and tidal mixing determine the frequency, severity, spatial and temporal extent of bloom events in coastal ecosystems (Paerl 2006).

Oxygen depletion in aquatic ecosystem can result from algal blooms. Harmful algal blooms reduce the aesthetic value of coastal and aquatic environment which causes
economic loss from ecotourism, fisheries resources and human health threat (Kevin et al. 2003). Human health threat occur from the consumption of shellfish or fish contaminated with cyanobacterial toxins. Shellfish such as clams, mussels and oysters rapidly accumulate algal toxins in their tissues because they filter large volume of water. Cases of HAB-related neurotoxic shellfish poisoning (NSP), paralytic shellfish poisoning (PSP), amnesic shellfish poisoning (ASP), diarrhetic shellfish poisoning (DSP) and saxitoxin puffer fish poisoning (a finfish poisoning) have been reported (Van Dolah 2000).

Biomasses can produce toxins (e.g., microcystin) which are released by cyanobacteria, diatoms and dinoflagellates (Paerl and Whitall 1999; Hitzfeld et al. 2000; Wetzel 2001; Anderson et al. 2002; Rabalais and Nixon 2002). Extensive study of population dynamics of cyanobacteria in relation to environmental factors (light, temperature, nutrient, pH, mixing or zooplankton grazing) have been documented (Downing et al. 2001; Paerl and Otten 2013; Fernández et al. 2015). Cyanobacteria can outcompete other algal groups because of their resistance to zooplankton grazing, buoyancy and nitrogen fixing ability by heterocystous forms (Graham and Wilcox 2000; Reynolds 2006). In addition, cyanobacteria can thrive in CO₂-deficient and low-pH conditions in aquatic ecosystem (Shapiro 1973; 1997; Sandrini et al. 2016; da Silva Brito et al. 2018). The frequency of eutrophic events has increased over the last several decades in many coastal ecosystems where blooms of phytoplankton are particularly affected by riverine discharge (Paerl et al. 2018). For instance, bloom incidences have been reported in Chesapeake Bay (Officer et al. 1984), the northern Adriatic Sea (Justic et al. 1987), the Baltic Sea (Andersson and Rydberg 1988), Neuse River Estuary (Paerl 2006), Pearl River Estuary (Yin and Harrison 2008), Kopāčki Rit floodplain (Mihaljevic and Stevic 2011), South African inland waters (Matthews and Bernard 2015), Wadden Sea (Vogt et al. 2017), Eurasian Arctic and Hypoarctic large River Estuaries (Davydov and Patova 2017) and Lake Taihu, China (Li and Qin 2019).

In recent years, the south-west coastal Nigeria is one of the fastest developing regions in West Africa. Epe lagoon boarders the eastern section of the largest metropolitan city (Lagos) in West Africa. Lagos consists of over 21 million inhabitants (Ojolowo and Onifade 2016) with associated rapid economic development and coastal degradation by human activities. Epe lagoon is connected to the Lagos lagoon and, hence, receives large loads of anthropogenic nutrients from increased agriculture, fish farming, poorly treated sewage effluent (Onyema et al. 2004; Akagha 2017) and domestic/industrial waste discharge (Chukwu and Nwankwo 2004). In Nigeria, the incidences of cyanobacterial blooms have been documented over the past three decades. Blooms of Anabaena flos-aquae, A. spiroides, Microcystis aeruginosa, M. flos-aquae and M. wesenbergii were reported in the Lagos lagoon (Nwankwo 1993; 1996), Microcystis aeruginosa in Iju, Ogun river (Nwankwo 1993), Microcystis sp. in Kuramo lagoon (Nwankwo 1993), Anabaena flos-aquae in Owo river (Nwankwo et al. 1999), Microcystis aeruginosa in Awba reservoir, Ibadan (Akin-Oriola 2003), Microcystis aeruginosa and M. wesenbergii bloom in Oyan dam, Ogun state (Adesalu et al. 2016) and the proliferation of Oscillatoria sp. in the Lagos lagoon (Onyema and Nwankwo 2009). These bloom series were reported to cause colouration, anoxia, odour and bad taste of the water in these aquatic ecosystems.

Akagha et al. 2019 documented the presence of a new cyanobacterial species Lagosinema tenuis in the Lagos lagoon which further highlights the ecological relevance of cyanobacterial dominance in tropical African lagoon systems. In this paper, our objectives are to elucidate: (1) environmental variables and phytoplankton dynamics, (2) nutrient dynamics and eutrophication and (3) annual cyanobacterial blooms and associated environmental factors in Epe lagoon through an 18-month investigation of three ecological sites.

Materials and methods

Study area

Epe lagoon (Fig. 1) is a tropical water body with a surface area of 243 km² located at the eastern part of the Lagos lagoon complex (Kusenjui 1988). Epe lagoon is fresh, lotic, non-tidal and sandwiched between Lagos and Lekki lagoons (Nwankwo 1998; Uwadibia 2010). It is connected to the Atlantic Ocean via the Lagos lagoon year round. Epe lagoon is influenced by freshwater input from creek and river inflow (Nwankwo 1998). Riparian vegetation such as Paspalum orbiculare, Raphia hookeri, Elaeis guineensis, Acrostichum aureum, Cocos nucifera, Rhizophora and mangroves Rhizophora racemosa, Avicennia nitida are found along the fringing wetland. Notable fauna found in this area are amphipods, oligochaetes, polychaetes, isopods, barnacles, oysters, nematodes, fiddler crabs and migratory birds that feed on exposed biota. The bordering wetland has experienced rapid population growth, agricultural and urban development over the past decade. As a result, anthropogenic activities including domestic waste deposition, fishing, indiscriminate sand mining and inland water transportation are increasing in Epe lagoon. Human influenced sites along the Epe lagoon include the hydrothermal plant at Egbin and agricultural sites on the bordering wetland at Ikosi. In this paper, three sites (Imope, Ikosi and Egbin) were investigated based on their ecological uniqueness and bloom incidences.
Collection of water and phytoplankton samples

Samples of water and phytoplankton were collected monthly for 18 months (November 2012–April 2014) in the morning. Water samples were collected in a well-labelled 1-L plastic bottle with screw cap and transported in a cooler with ice to the laboratory for chemical analysis. Three 250-ml amber bottles were used to collect water samples for analysis of dissolved oxygen, biological oxygen demand and chlorophyll a. Samples for dissolved oxygen were fixed with the Winkler reagent. Phytoplankton samples were collected with a 35 μm plankton net tied unto a motorized boat and towed at low speed (4 knots) for 5 min. The plankton net was hauled in and the samples were emptied into a 500-ml well-labelled plastic container with screw cap and fixed with 4% unbuffered formalin.

Physical and chemical parameters

Water temperature was measured in situ with a mercury thermometer, and water depths were measured with a calibrated pole. A Philips pH meter (Model PW950) and a hand-held Refractometer (model RHS-10 ATC) were used to determine pH and salinity, respectively. Total dissolved solids (TDS) were measured with HANNA instrument (HI98311). Total suspended solids (TSS) were measured following standard methods (APHA 2005). Dissolved oxygen was determined by the standard Winkler method, and biological oxygen demand (BOD) was measured after 5 days of incubation. Chemical oxygen demand (COD) was estimated using closed Reflux method (APHA 2005). Chlorophyll a determination was made using the acetone extraction spectrophotometric method (APHA 2005) with detection limits of approximately 1 µg/L, which was based on three times the standard deviation of the lowest concentrations of the samples. Rainfall data were provided by the Department of Meteorological Services, Oshodi, Nigeria.

Inorganic macronutrients and nutrient ratio

Phosphate and nitrate were analysed using ascorbic acid spectrophotometric method and the phenol disulphonic acid method, respectively (APHA 2005). Sulphate and silicate were determined using the standard methods (Strickland and Fig. 1 Map of study area indicating the three sampling sites: Egbin, Ikosi and Imope
To determine the seasonal trend of nutrient (molar) ratios, data were pooled for each season and the average were taken as follows: dry season I (November 2012–February 2013), early wet I (March 2013–June 2013), late wet I (July 2013–October 2013), dry season II (November 2013–February 2014) and early wet II (March 2014–April 2014).

**Stoichiometric assessment of nutrient limitation**

Stoichiometric assessment of nutrient limitation was made according to the published criteria (Dortch and Whitledge 1992; Justic et al. 1995a, b; Xu et al. 2008). The ratio of two ambient nutrients was calculated, and the Redfield ratio (Redfield et al. 1963) was applied to predict that: (1) N limitation occurs when N/P < 16:1 and Si/N > 1:1 and an estimated threshold value of 1 mM; (2) P limitation was assumed when N/P > 16:1 and Si/P > 16:1 with an estimated threshold value of 0.1 μM; and (3) Si limitation was assumed when N/Si < 1:1 and Si/P < 16:1, with an estimated threshold value of 2 μM. The approach was similar to others, for example, in the Mississippi River estuary (Justic et al. 1995a and in Hong Kong waters where the variations of nutrient stoichiometry in regulating phytoplankton biomass were used (Xu et al. 2008).

**Phytoplankton**

The preserved phytoplankton samples were concentrated by allowing cells to settle for at least 24 h and thereafter decanted. The micro-transect drop count method as described by Lackey (1938) was employed to investigate the phytoplankton species composition and abundance. One drop of each concentrated sample was thoroughly investigated by observing all fields within the cover slip border using an Olympus BX51 binocular microscope with calibrated eye piece at different magnifications (10 × and 40 ×). This was done five times to account for all phytoplankton species in the water samples. The total number of phytoplankton identified was recorded as number of cells per ml. Taxonomy publications including Patrick and Reimer (1966, 1975), Prescott (1961, 1973, 1975, 1979, 1984), Compère (1976, 1977), Komárek and Fott (1983, Round 1981), Vanlandingham (1982), Krammer and Lange-Bertalot (1986, 1988, 1991, 2000), Krammer et al. (1991a), Komárek and Anagnostidis (1998, 2005) and Wołowski and Hindak (2005) were consulted to identify phytoplankton taxa.

**Statistical analyses**

Analysis of variance (ANOVA) was used to test the significant differences of each parameter between the sampling sites. The correlation coefficient, r was calculated using Microsoft® Excel 2015. The statistical test for significance is based on critical values of correlation coefficient, r (Zar 1999).

**Results**

**Physical and chemical variables**

Rainfalls ranged between 0 and 413.6 mm during the study period. In January 2013 (bloom episode I), there was no rainfall, but increased to 80.1–132.7 mm in January and February 2014 (bloom episode II). Water temperature oscillated between 23 and 34 °C with the minimum recorded in August 2013 (Fig. 2). Total suspended solids and total dissolved solids were between (4–3166 mg/L) and (30–5290 mg/L), respectively. pH was 6.1–7.7, indicating acidic conditions, while salinity was in the range of 0–5‰ (Fig. 2). Dissolved oxygen, BOD and COD were in the range of 2–15 mg/L, 0.5–13 mg/L and 15–160 mg/L, respectively (Fig. 3). There were no significant differences (P > 0.05) in the environmental variables across all the sampling sites except for total suspended solids which was significantly different (P > 0.05) across the sampling sites.

**Chlorophyll a**

Chlorophyll a ranged from 1 µg/L (below the limit of detection) at Imope in February 2013 to 201 µg/L at Ikosi in February 2014. Chlorophyll a values were higher (≥ 55 µg/L) in the dry months (November 2012–January 2013) at the three sites during the first annual cycle, and higher values (≥ 153 µg/L) were recorded at Ikosi during the dry month (January 2014–February 2014) of the second annual cycle (Fig. 3).

**Nutrient concentrations and molar ratios**

Nitrate ranged from 1.45 to 20.16 µM, whereas lower values (≥ 1.45 µM) were recorded in January 2013 at all sites. PO4, SiO4 and SO4 concentrations were (0.18–18.76 µM), (0.03–4.24 µM) and (0.10–18.63 µM), respectively. Nitrate, phosphate and sulphate concentrations showed seasonal trends. Higher values of nitrate were recorded in the wet month (≥ 2.42 µM) at the three stations, whereas lower sulphate values (≤ 4.37 µM) were observed in the wet season except in March 2013 when values were higher (≥ 12.08 µM) at Egbin. SiO4 values were highest at Imope (≥ 2.31 µM) in February 2013 and at Ikosi (≥ 4.24 µM) in March 2013 (Fig. 3). N/P ratio at the three sampling sites varied seasonally unlike Si/N ratio. Low N/P ratios (≤ 4.5:1) were recorded during the first 8 months and shifted to higher ratios (≥ 16.7:1) during the subsequent
Fig. 2 Monthly variations in temperature (23–34 °C), pH (6.1–7.7), salinity (0–5‰), TSS (4–3166 mg/L) and TDS (30–5290 mg/L) at the three stations.
8 months (late wet I and dry season II), followed by low ratios (≤ 13.21) in the last 2 months at the three sampling sites (Fig. 4). Overall, N/P ratio values were between 0.22–58.23 at Imope, 0.22–48.1 at Ikosi and 0.21–59.4 at Egbin, while Si/N ratios were between 0.01–0.90 at Imope, 0.01–0.33 at Ikosi and 0.01–0.26 at Egbin (Fig. 5). Based on Si/N = 1:1, N/P = 16:1 and Si/P = 16:1, there were six different areas characterized by the most limiting nutrients. NO₃, PO₄ and SO₄ were not significantly correlated with chlorophyll a at Ikosi, except for SiO₃ which showed a significant and negative relationship with chlorophyll a (Fig. 6).

**Phytoplankton dynamics**

A total of 116 species belonging to 54 genera were recorded during the study, and there were six phytoplankton groups including Bacillariophyceae (40.7%), Chlorophyceae (30.1%), Chrysophyceae (0.89%), Cyanophyceae (20.4%), Dinophyceae (0.89%) and Euglenophyceae (7.08%). The centric diatom, *Aulacoseira granulata*, was the dominant species except in January 2013, 2014 and February 2014.
(dry season I and II) when there was an incidence of cyanobacterial bloom. At Imope, the lowest diatom abundance (48%) was recorded in April 2013, while cyanobacterial abundance was highest (22–47.0%) in May, June and July 2013. At Ikosi, the lowest diatom abundance (37.0%) was recorded in May 2013, while cyanobacterial abundance peaked (≥ 98.0%) in January 2013, January 2014 and February 2014. Diatoms were lowest (5.0%) in April 2013, while cyanobacteria were highest (≥ 36.0%) in April and May 2013 at Egbin (Fig. 7). The prevalent bloom species were *Anabaena circinalis*, *A. flos-aquae*, *A. limnetica* and *A. spiroides*.

Twenty-three cyanobacterial taxa were recorded. Species richness was observed in the Order Nostocales with 10 taxa, Chroococcales with 5 taxa, Oscillatoriales with 4 taxa, Spirulinales with 2 taxa and Synechococcales with 2 taxa, respectively (Table 1). Two cyanobacterial bloom episodes were observed at Ikosi, Epe lagoon, during the two annual cycles of the study when rainfall was low (Fig. 8). The first bloom began in January and ended in February 2013, while the second bloom started in January 2014 and ended in March 2014. There was notable increase in phytoplankton density (> 40,550 cells/ml) and the dominance of cyanobacteria (> 98%). During the first bloom episode, *Anabaena circinalis* (30,000 cell/ml, 73.0%) and *Anabaena limnetica* (10,000 cell/ml, 24%) were dominant, but the bloom collapsed in February 2013. Conversely, during the second bloom episode, *Anabaena flos-aquae* (40,000 cell/ml, 99.0%) was dominant. *Anabaena flos-aquae* (15,050 cells/ml, 27%), *A. circinalis* (30,000 cells/ml, 53%) and *Anabaena spiroides* (10,000 cell/ml, 18%) were recorded in February 2014 at Ikosi. The bloom collapsed in March 2014. At Ikosi, cyanobacterial cells were ≥ 98.0% of the total phytoplankton density which was higher (2000 cell/ml–40,000 cell/ml) than the alert level 1 (≥ 2000 cells/ml) for raw waters (Bartram, 2012).
et al. 1999) during the two bloom episodes. The cyanobacteria bloom did not reach alert level 2 (≥ 100,000 cells/ml) at the sampling sites during the study period (Fig. 9).

Discussion

Environmental dynamics

Seasonality and distribution of terrestrial and aquatic organisms in the tropics are determined by rainfall pattern (Webb 1960). Rainfall is a driving force in aquatic ecosystems of south-west, Nigeria (Olaniran 1969; Oyenekan 1988; Nwankwo 1996) because it influences flow rate, mixing, dilution and nutrient recycling (Valiela et al. 2013). The observed lowest temperature (≤ 27 °C) in the wet season (late wet I) might be attributed to the influx of cooler flood waters from wetlands, adjoining creeks and rivers. These assertions confirm temperature regimes in coastal lagoons of south-west, Nigeria (Onuoha and Vyverman 2010; Nwankwo et al. 2013). The low total suspended solids and total dissolved solids in the wet months agree with observations of Nwankwo (1996, 1997), Onyema and Nwankwo (2006), Adesalu and Nwankwo (2010) and Onuoha and Vyverman (2010) in coastal waters of south-west Nigeria. Flood water intrusion causes dilution, high flushing rate and low retention time during the rainy season, which is different from the situations in the dry season and the onset of the rainfall. pH indicated an acidic condition in certain occasion (July 2013) which could be attributed to the seepage of humic and fulvic acid exudates from surrounding wetlands (Nwankwo et al. 2013; Canini et al. 2013). The slightly acidic to neutral nature of Epe lagoon probably accounted for the dominance of Aulacoseira granulata at Epe during the off-bloom season. Talling and Talling 1965 associated the proliferation of Aulacoseira granulata to low pH in studies of central African lakes. The observed pH remained within the acceptable limits of 6.3–8.5 for inland waters (Antoine and Saadi 1982). At Ikosi, a pH range between 6.7 and 7.4 was found to correspond to the period of high prevalence of Anabaena flos-aquae, A. circinalis, Anabaena limnetica, Anabaena sphaerica and Anabaena spiroides.

Dissolved oxygen is dependent on water temperature and decreases as water temperature increases. Photosynthesis, respiration and other hydrological dynamics such as river influx, flushing rate and mixing influence dissolved oxygen levels in aquatic ecosystems (Watt 2000). In this study, dissolved oxygen and BOD were relatively low especially during the bloom episodes in the dry season. The relatively higher DO and BOD values that were observed during the rainy season might be due to mixing of surface water with atmospheric oxygen by current, wave action and river inflow. Chindah and Nduaguibe 2003 observed high dissolved oxygen levels attributed to current and wave action in a study of the Bonny river in Niger Delta.

In lotic fresh waters, Hynes (1960) suggested that BOD between 6.0 mg/L and 8.0 mg/L indicates moderate pollution while values greater than 8.0 mg/L indicate severe pollution. BOD values at Epe lagoon suggest that Epe lagoon is moderately to highly polluted during the wet season, owing
to influx of land-originated pollutants. The acceptable limit of BOD set by the WHO (1978) for international water quality standards is 15.9–37.5 mg/L with warning limit of 18.9–34.9 mg/L. An increase in the amount of organic materials in aquatic ecosystems results in high levels of COD (WHO 1978). COD values were high at Ikosi in January 2013 and February 2014, Imope in March 2014 and Egbin in April 2014. The observed higher values of COD at these sites during this period may be due to the collapse of the bloom and horizontal mixing of the surface water. There was no significant difference ($P > 0.05$) in water temperature, pH, salinity and TDS across all the sampling sites, whereas significant difference ($P > 0.05$) was observed for total suspended solids which was high at the onset of the rainy season.

**Nutrients and molar ratios**

Nutrient concentrations and variations in stoichiometric nutrient ratio ($\text{Si/N}$, $N/P$ and $\text{Si/P}$) are influenced by rainfall, human activities (Elser et al. 2009; Yadav and Pandey 2018), internal nutrient cycling and regeneration (e.g. denitrification, nitrogen fixation) (Hudson et al. 1999). The rate of nitrogen and phosphorus cycling through sediment influences nitrogen or phosphorus limitation in aquatic ecosystems (Vitousek and Howarth 1991). In freshwater and coastal marine ecosystems, nitrogen is removed in sediments through denitrification (Knowles 1982; Seitzinger et al. 1984; Seitzinger 1988). Phosphorus is readily released from sediments through mineralization (Rowe et al. 1975; Boynton et al. 1982). In tropical waters of west Africa, nitrogen concentrations are mostly introduced through anthropogenic sources. The reduction of nitrogen and/or phosphorus inputs into aquatic systems can improve the quality of the water. Nutrient reduction effort can be challenging especially when it is introduced into the aquatic ecosystem through non-point sources such as agricultural run-off. The improvement of water quality may be slowed by internal loading of nutrients from sediments after external loading of nutrients is reduced (Søndergaard et al. 2003). Nutrient stoichiometry elaborates the role of resource availability in aquatic ecosystems (del Amo et al. 1997) as well as resource competition in phytoplankton (Tilman 1982). N/Si/P ratio of marine diatoms is about 16:16:1 (Redfield et al. 1963; Brzezinski 1985) and deviations from this ratio may result in nutrient limiting for phytoplankton (Hecky and Kilham 1988; Dortch and Whitledge 1992). Silicate was limiting ($\text{Si/N} < 1$) at the study sites, possibly resulting from the absence of tidal sea water influence in Epe lagoon, introduction of organic nitrogen and phosphates by flood water and agricultural run-off. Nutrient-laden influx from agricultural practices could result in increased levels of nitrate and phosphate, as well as relatively low or declining silicate concentrations (Justic et al. 1995b). Nwankwo (1996) attributed higher silicate values in the dry season to the cessation of flood water discharge and the influx of tidal sea water into the brackish Lagos lagoon. van Bennekom and Wetsteijn (1990) reported that Si concentration at the Rhine River remained constant during a long-term study. Comparably, Si concentrations were reduced by 50% during 1955–1987 in the Mississippi River (Turner and Rabalais 1991). In this study, silicate values were higher than the threshold value 2 µM (Justic et al. 1995a) at Imope ($\geq 2.31$ µM) in February 2013 and at Ikosi ($\geq 4.24$ µM) in March 2013. These values coincided with the lower cyanobacterial biomass at Imope in February 2013, the collapse of cyanobacterial bloom and the proliferation of diatoms (84.38%) at Ikosi in March 2013 after the bloom regime (Fig. 7).
NO₃ values (≥ 1.45 µM) at the three sampling sites were higher than the threshold value 1 µM (Justic et al. 1995a). The increase in nitrate levels during the wet season could be attributed to the introduction of floodwaters that contained organic materials from adjoining wetland and nutrient rich agricultural run-off due to poor farm practices. Of all the five seasonal partitions, low N/P was recorded during the dry season I (November 2012–February 2013), early wet I (March 2013–June 2013) and early wet II (March 2014–April 2014) during this study. The observation during the dry season I might be a consequence of faster utilization of nitrate by phytoplankton in aquatic environments, while low N/P during early wet season I and early wet season II might be a sign of the effect of high flushing rate and low retention time during this period. N/P was high (≥ 16.72) at the three sites during the late wet season I (July 2013–October 2013) and dry season II (November 2013–February 2014). PO₄ values (≥ 0.18 µM) at the three sampling sites were higher than the threshold value 0.1 µM (Justic et al. 1995a).

Rainfall values were high during the late wet season I which caused intense nutrient-laden flood water intrusion into the lagoon. However, the absence of rainfall in August 2013 possibly slowed flushing and increased residence time of the lagoon during this period. Flood water inflow influenced eutrophication in coastal waters because nitrate and phosphate from agricultural run-offs (fertilizers) are released (Qian et al. 2016). The resulting effect is the change in N/P, Si/P and Si/N leading to potential silicate limitation because of the increased nitrate and phosphate concentrations. The dry season II period was associated with reduced volume of freshwater inflow from adjoining river, less perturbation stress related to mixing, reduced flushing and higher residence time in the lagoon. This situation probably explained the high N/P ratio values and cyanobacterial bloom during this period. Relationships between riverine nutrient input, land based flood water and eutrophication in coastal waters have previously been reported in the Chesapeake Bay (Officer et al. 1984), the northern Adriatic Sea (Justic et al. 1987), some areas of the Baltic Sea (Andersson and Rydberg 1988), Mississippi river (Turner et al. 1998), Neuse River Estuary (Paerl 2006), Pearl River Estuary (Yin and Harrison 2008) and Kopački Rit floodplain (Mihaljevic and Stevic 2011).

Sources of sulphate in aquatic environments can be either natural or anthropogenic. Industrialization, burning of fossil fuel and agricultural practices contribute to sulphate availability in coastal ecosystems (Rhode et al. 1995; Zhao et al.

| Cyanobacteria species | Imope F% | Ikosi F% | Egbin F% | Order       |
|-----------------------|----------|----------|----------|-------------|
| Anabaena affinis Lemm. | –        | –        | 5.6      | R           | Nostocales  |
| Anabaena azollae Strasburger | –  | –        | 5.6      | R           | Nostocales  |
| Anabaena catenula Strasburger | – | –        | 11.1     | C           | Nostocales  |
| Anabaena circinalis Rabenhorst ex Bornet and Flahault | – | 22.2      | C        | –           | Nostocales  |
| Anabaena flos-aquae (Lynge) | – | –        | 22.2     | C           | Nostocales  |
| Anabaena limnetica Smith | – | –        | 11.1     | C           | Nostocales  |
| Anabaena spheerica Bornet and Flahault | – | –        | 5.6      | R           | Nostocales  |
| Anabaena spiroides Lemm. | 11.1     | C        | 27.8     | C           | Nostocales  |
| Aphanocapsa delicatissima West and West | 5.6 | R        | 16.7     | C           | Synechococcales |
| Chroococcus dispersus (Keiss) Lemm. | 5.6 | R        | 16.7     | C           | Synechococcales |
| Chroococcus limneticus Lemm. | 5.6 | R        | 16.7     | C           | Synechococcales |
| Merismopedia elegans Braun | – | –        | 5.6      | R           | Synechococcales |
| Merismopedia glauca (Ehr) Nageli | 11.1 | C | –        | –           | Synechococcales |
| Microcystis aeruginosa (Kützing) Kützing | 38.9 | C | 33.3     | C           | Chroococcales |
| Microcystis flos-aquae (Wittrock) Kirchner | 11.1 | C | 44.4     | C           | Chroococcales |
| Nostoc sp. | 5.6     | R        | 11.1     | C           | Nostocales  |
| Oscillatoria curviceps Agardh ex Gomont | 5.6 | C | –        | –           | Oscillatoriales |
| Oscillatoria limosa (Roth) Ag. | 22.2 | C | 16.7     | C           | Oscillatoriales |
| Oscillatoria princeps Vaucher | 5.6 | R | 5.6      | R           | Oscillatoriales |
| Oscillatoria tenius Agardh | 11.1 | C | 11.1     | C           | Oscillatoriales |
| Spirulina laxa Smith | 16.7    | C | –        | –           | Spirulinales |
| Spirulina major Kützingii | – | 11.1      | C        | –           | Spirulinales |
Sulphate enters aquatic ecosystems through leachates from soil, precipitation, petroleum spill and ammonium sulphate fertilizers. Phytoplankton utilize sulphate for physiological and metabolic processes (Giordano et al. 2005) although little is known about its role in phytoplankton bloom. Sulphate concentrations regulate the flux of phosphorus from sediment to an extent. At low sulphate levels, phosphorus is usually adsorbed to sediment while high sulphate levels support the release of phosphorus to the water column (Caraco et al. 1989, 1990). Sulphate concentrations in this study were generally higher in the dry than the wet season because of the reduced flood water influx and longer retention time during the dry season. The presence of petroleum products in this lagoon could be the source of observed SO$_4$.

**Phytoplankton dynamics and controlling factors**

Chlorophyll $a$ followed a seasonal trend and was closely related to phytoplankton abundance. Similarly, Suzuki et al. (2002) reported that low chlorophyll $a$ value implied limited phytoplankton growth in a turbid Mexican lagoon. The rise in chlorophyll $a$ during the dry season may be related to increased insolation, photosynthetic depth, retention time as well as less perturbation stress from flood waters. Flushing of planktonic microalgae by flood could lead to the low chlorophyll $a$ values in the rainy season. A single major rainfall peak in the dry season (January–February) and a minor peak in the late rainy season (August–November) were documented at Ikpoba reservoir, Edo state, Nigeria (Kadiri 1993).

The dynamics and seasonality of chlorophyll $a$ pointed to the interplay between phytoplankton growth and loss rate which are caused by multiple mechanisms in aquatic ecosystem (Cloern 1996). Phytoplankton population dynamics are dependent on changes in the proportion of dissolved Si, N and P. Officer and Ryther (1980) hypothesized that decreasing Si/N ratio may increase eutrophication by reducing the potential for diatom growth in favour of harmful phytoplankton species. Long-term decline in Si/P ratios was responsible for significant blooms of non-siliceous algae in coastal waters worldwide (Smayda 1990).

Phytoplankton successional pattern at Ikosi showed a clear transition from diatom to cyanobacterial dominance in the dry season during the two annual cycles. The coincidence of the cyanobacterial bloom in the dry season could be attributed to higher insolation, water temperature, photosynthetic depth and increased stability of water (Fig. 10). *Anabaena catenula, A. circinalis, A. limnetica* and *A. spiroides* which are nitrogen-fixing cyanobacteria dominated the phytoplankton community during the bloom episode in the dry season I and II at Ikosi. The bloom episode during the dry season I coincided with low N/P ratio, possibly due to improved water stability and phosphorus.
being more available as a result of biogeological process and mineralization (Rowe et al. 1975). The prevalence of N fixing cyanobacteria is a response to nitrogen limitation in aquatic ecosystems (Paerl et al. 2014). In this study, higher nitrogen concentration (N/P = 16.7:1) during the bloom episode in the dry season II may suggests an input from the surrounding wetland. In the tropics, rainfall controls physical, chemical and biological dynamics in aquatic ecosystem. Consequently, this might be the reason for the low nitrogen during the bloom episode I (January 2013) when there was no rainfall. However, rainfall values (≥ 80.1 mm) accounted for the nitrate availability during the bloom episode II (January and February 2014) which lasted longer because of the associated nutrient-laden river influx.

Despite the increased stability of the water column during the bloom episode I when there was no rainfall, the bloom duration was shorter than the bloom episode II. This suggests that in addition to optimal environmental conditions (light, temperature, stability etc.), riverine and land-based nutrient influx might be an important factor that influenced the bloom duration in the tropical aquatic environments.

**Eutrophication, cyanobacteria bloom and consequences**

A visible greenish colouration was seen on the surface water at Ikosi on three sampling occasions. The two cyanobacterial bloom regimes occurred in the dry season and were related to environmental conditions. The duration of the bloom was determined by the rainfall pattern which controlled nutrient influx, flushing rate and residence time. Higher insolation and improved stability of the lagoon during the dry months probably favoured the proliferation of N fixing cyanobacteria. Domestic waste discharge, agricultural run-off and poor sewage systems are sources of organic nutrient enrichment in coastal waters of Nigeria. Coastal lagoons are residual sink for large nutrient loading associated with anthropogenic and agricultural activities. These introductions cause an imbalance in the aquatic ecosystem which alter food chain that lead to loss of biodiversity and productivity (Nwankwo and Akinsoji 1992; Kevin et al. 2003). Nutrient loading has led to intensified eutrophication causing reoccurring algal bloom episodes including harmful species, accrual of organic matter and oxygen depletion (hypoxia and anoxia). In addition to eutrophication, climatic and hydrogeological factors such as rainfall pattern, flood, temperature rise, flow rate, river and storm water discharge influence anoxia and hypoxia conditions in aquatic ecosystems. The complex interactions of these environmental factors determine the magnitude of the temporal and spatial dynamics of algal blooms and oxygen depletion (Cloern 2001, Thomas and Litchman 2016). The complexity of these systems affects the physical, chemical and biological processes controlling the production and accumulation of organic matter, oxygen dynamics and nutrient cycling (Paerl 2006). DO depletion (anoxia and hypoxia) is one of the major consequences of eutrophication in coastal waters (Harrison et al. 2008). The variation between relatively high rates of oxygen consumption and low rates of oxygen introduction results in a decrease in DO levels. Very low or hypooxic conditions (DO ≤ 2.6 mg/L) were observed at the three sites in January 2013 and at Ikosi (DO ≤ 5 mg/L) in January 2014 due to the bloom episode and subsequent oxygen utilization by microbial degradation of organic matter.

**Anabaena catenula, A. circinalis, A. limnetica and A. spiroides** were the bloom species observed in the lagoon. Some of these species have been reported to be potentially toxic. For instance, potentially toxic cyanobacteria *Snowella fennica, Microcystis aeruginosa, Planktothrix agardhii* and *Anabaena circinalis* were reported in Paso de las Piedras reservoir, Buenos Aires, Argentina (Fernández et al. 2015). Furthermore, blooms of *Aphanizomenon*, *Dolichospermum*, *Nodularia* and *Phormidium* have been documented in ecological studies of the Eurasian Arctic and Hypoarctic large river estuaries (Davydov and Patova 2017). In reference to the alert framework for cyanobacteria in coastal waters (Bartram et al. 1999), alert level 1 (≥ 2000 cells/ml) was exceeded at Epe lagoon (Ikosi). Cyanobacterial biomass was high (40,000 cell/ml), almost reaching alert level 2 (≥ 100,000 cells/ml). These observations call for an alarming concern and demand continuous environmental monitoring and management of this coastal lagoon.

**Conclusion**

The factors regulating the severity of algal blooms in aquatic ecosystems are complex. In this study, rainfall, anthropogenic activities along bordering wetlands and hydrodynamics are forcing factors which influence phytoplankton
communities, nutrient dynamics and bloom development. The hypoxic conditions observed during the two bloom episodes reflect the vulnerability of the biological component of this lagoon. Rainfall controlled nutrient dynamics which triggered cyanobacterial bloom development and influenced the bloom duration. Nitrate, phosphate, sulphate and other environmental factors were related to the seasonality whereas silicate was relatively low. Blooms were dominated by potentially toxic species: *Anabaena circinalis*, *A. flos-aquae*, *A. limnetica* and *A. spiroides*, and were related to nitrate dynamics. This study provides an important scientific information to the development of tools to assess, manage and mitigate risk of noxious bloom occurrences in tropical lagoons.

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**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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