Oligotrophication after a nutrient reduction in a shallow sand-pit lake (Créteil Lake, Paris suburbs, France) : a case of rapid restoration

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Keywords : shallow lake, sand-pit, nutrient reduction, oligotrophication.

A limnological study was performed out, over 8 years from 1979 to 1986, in a shallow recently created sand-pit lake, Lake Créteil (Paris suburb, France). The lake is mainly supplied by phreatic waters. Typical of shallow waters, thermal stratification which occurred from May to October was intermittently broken by wind, leading to high seasonal fluctuations and allowing a reoxygenation of the total water column. The interannual development of the lake was characterized by a regular increase in transparency (from 1 m in 1979 to 2.7 m in 1985) which was partly due to not only stabilization of the bottom sediment and the banks, but also to a reduction of the biological component. A decrease in chlorophyll a concentrations resulted from nutrient diversion of domestic inputs collected through the rainwater pipe coming into the lake. The rapid restoration of water quality shows that this small artificial lake is very sensitive to human interventions.

Oligotrophisation d’une sablière en eau (Lac de Créteil, région parisienne, France) après une réduction des apports en éléments nutritifs : un cas de rapide restauration

Mots clés : lac peu profond, sablière en eau, réduction en éléments nutritifs, oligotrophisation

1. Introduction

Limnological studies have mainly focused on natural lakes ; some of them are particularly well documented in literature (Constance, Geneva, Victoria, Balaton or Windermere lakes…) whereas the knowledge of others have even been recently integrated into books (Mendota Lake : Billings et al. (1985), Mirror Lake : Likens (1985), Stechlin Lake : Caspers (1985)). More recently, a book was proposed analysing reservoir functioning (Thornton et al. 1990).

Although sand-pit lakes have become more and more numerous in the last few years, due to the increasing need in building materials, less attention has been paid to these kind of ecosystems. Some ecological characteristics of several sand-pit lakes have however been given in Garnier et al. (1987).
The sand-pit lake of Créteil (Paris suburb, France) has been intensively studied for 8 years. The study started in 1979 soon after the quarrying of sand has finished in 1976. The study period can therefore be considered as a field experiment around the following question: how long would be necessary for this young system to achieve a relative stability? Such a recently created system was a priori appropriate to interpret the possible process of naturalization of an artificial aquatic system. However, situated in a town, the role of Lake Créteil as a recreational and ornamental area has required several management measures which interfered with natural processes. The paper analyses the seasonal and interannual changes of that young artificial aquatic system.

2. Site study

Créteil lake is situated in an urbanized area near the confluence of the Marne and Seine rivers 15 km SE of Paris, France (Fig. 1). Parisian Basin has a temperate climate and has been rather stable for the period studied, although some year-to-year variability has been observed. A summary of annual mean and extreme values is given for global incident radiation, temperature, rainfall and wind (Table 1).

The Créteil sand-pit lake originated from the excavation, between 1947 to 1976, of the alluvial sediments. The alluvial plain is delimited by the Seine (West) and Marne rivers (North) and by Tertiary outcrops (East, South). Créteil Lake is not connected with the rivers. The lake is mainly supplied by anoxic phreatic waters of the alluvial aquifer, water circulating mainly through diverse filling materials which have replaced the natural alluvium dug out in the past. Besides direct rainfall, the lake also receives rainwater runoff from the 0.5 km² of the Mont Mesly urbanized zone. The water residence time in the lake was estimated at 6-12 months (Chestérikoff 1981). The high total salinity (1.4 g l⁻¹) of Lake Créteil water originates from the composition of the supplying underground water (Table 2). The dominance of calcium and sulfate is mainly caused by the gypsum strata of Mont Mesly outcrops whereas the high concentration of sodium and chloride is due to the filling materials. Total salinity in the water coming from the rainwater pipe is lower (0.87 g l⁻¹). Data collected from 1978 to 1980 and again in 1986, show that the ionic composition of the lake water has been rather stable for the period studied, although it slightly decreased in 1986 (Table 2).

The surface area of the lake is 0.42 km²; maximal length and width are 1.5 and 0.4 km respectively following a NS axis. The banks are steep so that macrophytes occupy only 15% of their length. The lake is surrounded by buildings at East and North-East, an artificial hill at West; it is exposed to wind from the South and South-East. The depth of the lake is homogeneous within most of its surface area but maximal depth has varied between 4 and 5 m.

During the study period, different management measures were successively taken, which all perturbed the trend of the lake functioning.

1981: The rainwater runoff collected on the urbanized watershed and brought to the lake by a pipe was accompanied by polluted domestic water at the beginning of the study. In May 1981, the polluted inputs were lowered.

1983: The water level of the lake regularly increased (1.27 cm/month) from 1979 to 1982 due to the consolidation of ground materials following the construction of an artificial hill (Chestérikoff & Chevreuil 1987), although this increase might also be caused by the increase in rainfall observed during that period (Table 1). At the beginning of 1983, a drainage pipe of 500 l/sec in capacity was set to regulate the level of the lake, decreasing the residence time.

1985: From December 1985 to August 1986, intensive pumpings of phreatic waters led to a regular decrease of the water level (3 cm/week, i.e. 1 m in total); initial values were then found when pumping ceased.

3. Material and methods

The study was carried out in the central area of the lake from 1979 to 1986 at intervals ranging from a week in summer to a month in winter.

Measurements of temperature and oxygen were performed in depth profiles with intervals of 50 cm, with a YSI (model 57) probe. Wind speeds were recorded at a meteorological station in the nearby
NUTRIENT REDUCTION IN A SAND-PIT LAKE

Fig. 1. Situation of Créteil Lake. Cross-section of the plain of Créteil (according to Chestérikoff & Chevreuil 1987).

Fig. 1. Situation du lac de Créteil. Coupe géologique de la plaine de Créteil (D’après Chestérikoff & Chevreuil 1987).
Table 1. Interannual variations, from 1979 to 1986, of annual means (x) and extreme values (min, max) for global incident radiation (Io : cal cm\(^{-2}\) day\(^{-1}\)), air temperature (T : °C), wind speed (wind : m sec\(^{-1}\)) and rainfall (mm year\(^{-1}\)).

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Table 2. Comparative conductivity (µS cm\(^{-1}\)) and major elements (mg l\(^{-1}\)) of the waters. Phreatic waters : range of values for the 7 to 9 different piezometers and exceptional high or low values in parentheses (Chestérikoff et al., 1981). Pipe waters : extremes values (ext) and mean (x) (Chestérikoff et al. 1981). Lake waters : mean values in 1978-1979 (Chestérikoff et al. 1981), in 1980 (Chestérikoff 1981) and in 1986.

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<td>830-1260</td>
<td>1155</td>
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<td>HCO(_3^-) mg l(^{-1})</td>
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<td>Cl(^-)</td>
<td>50 - 200 (1600)</td>
<td>64 - 220</td>
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<td>90 - 200</td>
<td>141</td>
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<td>Ca(^{2+})</td>
<td>144 -1000</td>
<td>100 - 154</td>
<td>127</td>
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<tr>
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<td>Na(^+)</td>
<td>23 - 100 (506)</td>
<td>52 - 148</td>
<td>98</td>
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<td>K(^+)</td>
<td>20 - 50 (422)</td>
<td>10 - 21</td>
<td>16.8</td>
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environment. Global radiation was also continuously recorded with a pyranograph (Eppley) equipped with an integrator (Mecilec) at the meteorological station. Light attenuation profiles were established every 20 cm, with a quartz meter coupled with an integrator (LI 193 S, LI 188). Vertical attenuation coefficient (K) was calculated from a linear regression given by the equation: \( \ln Q_z = Kz + \ln Q_0 \) where \( Q_0 \) and \( Q_z \) are the irradiance under the surface and at the depth \( z \). Transparency was estimated with a Secchi disk (white, 30 cm diameter).

Nutrient analyses were carried out on the water taken at 1.5 m from 1979 to 1981 and on a mixed water sample of six depths (0, 0.5, 1, 1.5, 2.5 and 4 m) from 1982 to 1986. Water samples for P-PO\(_4\), N-NO\(_3\) and N-NH\(_4\) were filtered through GF/F membrane filters and frozen until analyses. P-PO\(_4\) was measured by spectrophotometer at 630 nm after a reaction with ammonium molybdate and a reduction to molybdenum blue by ascorbic acid. Total Phosphorus (P-tot) was determined as above on unfiltered lake water after sodium persulfate digestion and mineralization at 110°C in acidic phase. N-NO\(_3\) was spectrophotometrically measured at 415 nm after the formation of Na-paranitrosalicylate whereas N-NH\(_4\) was determined at 635 nm by the indophenol method. As relatively small values of N-NO\(_2\) were measured at the beginning of the study, total inorganic nitrogen (N-tot) was considered to be the sum of N-NO\(_3\) and N-NH\(_4\), although N-NO\(_2\) might have developed transiently as a result of degradation of organic matter.

Chlorophyll \( a \) was determined spectrophotometrically after filtration through a Whatman GF/C filter, extraction in 90% acetone and centrifugation (Lorenzen 1967). The volume of filtered water varied from 0.5 to 3 L, and samples were duplicated. Values represent an average of the six depths.

4. Results

4.1. Physical and chemical parameters of the lake

Water temperatures were lowest in January (2°C within the whole water column). Then, together with the increase of global radiation and day length, water temperature regularly increased from mid-February onwards and reached 15°C in April (Fig. 2). At that time (end of April to beginning of May), a thermal gradient, (4-5°C as a maximum) regularly appeared; it disappeared entirely in mid-October when the temperature had decreased to 15°C. Maximal surface temperatures varied from 22°C (1982) to 25°C (1983 and 1986). Intermittent mixing occurring during that period under wind effect was of major significance for phytoplankton seasonal periodicity (Gamier 1989, 1992). Periods of stratification were generally short and never lasted for more than 6 weeks. In 1985 and especially in 1986 when the level of the water was lowered, the amplitude of the thermal gradient was reduced.

A continuous increase in water transparency, \( Z_s \) (a decrease in light attenuation coefficient, \( K \)) occurred throughout the whole period of the study (Fig. 3). Increases of \( Z_s \) between 1979 and 1985 were significant according to Kruskal-Wallis test and multiple comparisons. A decrease of \( Z_s \) was observed in 1986 when the water level was lowered. Also, during the first four years of the study (1979 to 1982), a seasonal trend in water transparency was observed, with values lower in summer (May to

![Fig. 2. Interannual variations of temperature (T: °C) and percentage saturation of oxygen (O\(_2\): %) at the surface (—) and 4 m (—) from 1979 to 1986.](image-url)
Fig. 3. Interannual variations of water transparency ($Z_s : \text{m}$), of the light attenuation coefficient ($K : \text{m}^{-1}$) and of chlorophyll $a$ concentrations ($\text{chl } a : \mu \text{g l}^{-1}$) from 1979 to 1986.

Fig. 3. Variations interannuelles, de 1979 à 1986, de la transparence de l'eau ($Z_s : \text{m}$), du coefficient d'extinction ($K : \text{m}^{-1}$) et de la concentration en chlorophylle $a$ ($\text{chl } a : \mu \text{g l}^{-1}$).

October) than in winter (November to April) (Wilcoxon-Mann-Whitney bilateral test). From 1983 onwards, the increase in water transparency was more pronounced, so that no seasonal tendency was observed in 1984 and 1986, mean summer values being even higher than winter values.

At a seasonal scale, three periods can be distinguished with respect to water oxygenation (Fig. 2). From January to April, the whole water column was saturated (values between 90 to 100 %) ; from May to October, depending on the stratification-destratification episodes, water was at times super-saturated in the upper column whereas oxygen depletion sometimes occurred deeper ; at the end of October, oxygen depth profiles were homogeneous but a slight depletion (saturation at 70 %) was observed, saturation being progressively restored towards December.

From May to October, oxygen depth profiles depended on the thermal structure. Oxygen depletion at the bottom was more pronounced when the thermal gradient has been important for a long period ; oxygen depletion at the bottom was accentuated by the supply of anoxic phreatic waters, circulating through strongly reducing filling material. In addition, when transparency was low at the beginning of the study, as photosynthetic processes were restricted within the upper water layer, differences between surface and bottom were accentuated. Therefore, wind induced intermittent mixing events had an important role in the re-oxygenation of the water column in summer. The undersaturation during autumnal mixing was caused by a considerable demand for dissolved oxygen for the decomposition of organic material produced during the period of maximum biological activity.

Although, as a general rule, oxygen followed the same year to year seasonal pattern, changes have nevertheless been observed throughout the studied period. Differences between surface and bottom have become of lesser importance at the end of the study when the increase in water transparency deepened.
photosynthetic activity (Fig. 2). As an average for the summer period, saturation percentage regularly increased from 48.8 % in 1979 to 88.7 % in 1985. In 1985 and 1986, minimum values at 4 m equalled 50 % whereas they decreased to 5 % from 1979 to 1982 (Fig. 2).

4.2. Changes in chlorophyll a level

Whereas transparency increase, concentrations in chlorophyll a decrease significantly during the period of the study; values, in annual means, were the highest (about 15 $\mu$g l$^{-1}$) from 1979 to 1981, decreased significantly in 1982 and 1983 (7 and 9.5 $\mu$g l$^{-1}$ respectively) and decreased still further in 1984 and 1985 (4.1 and 3.6 $\mu$g l$^{-1}$) whereas a slight increase occurred in 1986 (5.4 $\mu$g l$^{-1}$), (Fig. 3). Wide variations in chlorophyll a concentrations were observed at a seasonal scale. High value in winter were often associated to early increases in abundance (not in biomass) of small algal species (Garnier 1989); the highest algal biomass found in summer and determined from cell volume (Garnier 1989) were not accompanied by the the highest values in chlorophyll so that a seasonal trend did not appear clearly.

4.3. Nutrients in the lake and sources of nutrients

A nutrient budget is difficult to establish for the Lake Créteil because the hydrodynamic and chemistry of the supplying phreatic water are very complex. The rainwater pipe furthermore complicates the system because, until May 1981, it also brought polluted domestic waters besides rainwaters.

The year-to-year trends in total phosphorus were clearly related to human intervention on the rainwater pipe which decreased the polluted effluents. Total phosphorus concentrations steadily increased in lake water from January 1979 to May 1981, values averaging 32.8 $\mu$g l$^{-1}$ in 1979, 48.5 $\mu$g l$^{-1}$ in 1980 and 71 $\mu$g l$^{-1}$ from January to May 1981 (Fig. 4); the nutrient diversion caused a decrease to an average of 37.6 $\mu$g l$^{-1}$ in the second part of 1981. From 1982 to 1986, annual average of total phosphorus (between 42 and 50 $\mu$g l$^{-1}$) stabilized in the range of the values found before the increase, although high values were occasionally still observed. The concentrations of orthophosphates decreased at the beginning of the study period and remained at a low level since then (Fig. 4). There was no seasonal trends for both total phosphorus and orthophosphates even when averaging summer (from May to October) and winter data (November to April).

At the beginning of the study, the rainwater pipe constituted a considerable input of phosphorus, but fortunately 80 % of the phosphorus have been shown to precipitate in the sediment (Chestérikoff et al. 1981). The precipitation would explain the relatively low increase in total phosphorus concentration in the lake compared to that in the rainwater pipe: from 1979 to 1980, concentrations in total phosphorus in the rainwater pipe increased indeed from 1500 to 9000 $\mu$g l$^{-1}$, (Table 3, Chestérikoff et al. 1981). In 1985, total phosphorus was analysed in lake water, just below the pipe: high values of phosphorus (P-tot : 119 $\mu$g l$^{-1}$ in annual average ranging from 36 to 250 $\mu$g l$^{-1}$; P-P04 : 60.6 $\mu$g l$^{-1}$ ranging from 4 to 193 $\mu$g l$^{-1}$) showed that urban waters brought by the rainwater pipe at the end of the study period still constituted a source of nutrients, despite the sewage diversion in May 1981. No apparent increase in phosphorus content has been observed in the lake sediment from 1980 (0.33 mg cm$^{-3}$) to 1987 (0.34 mg cm$^{-3}$).

Phreatic waters also appeared to be a source of phosphorus for the lake; measurement performed at the beginning of the study from piezometers were indeed higher than those of the lake (Table 3, Chestérikoff et al. 1981).

Total inorganic nitrogen (N-tot) decreased at the beginning of the study, from 1.08 to 0.75 mg l$^{-1}$ between 1979 and 1980 as an annual average (from 3.9 to 1.15 mg l$^{-1}$ for maximal values) but also drastically decreased in May 1981; values stabilized around 0.3 mg l$^{-1}$ from 1982 to 1986 (Fig. 5). The decrease in total inorganic nitrogen resulted from a simultaneous decrease of both ammonium and nitrate representing respectively 2/3 and 1/3 of total inorganic nitrogen.

Despite some irregularity in the seasonal pattern, nitrate tended to decrease in summer, this tendency being apparent at the beginning of the study when higher values were found (Fig. 5). No seasonal trend was found for ammonium, excepted in 1979.

Similarly to phosphorus, the pipe and phreatic waters constituted a source of nitrogen; concentration in total inorganic nitrogen (N-tot) in the pipe waters (18.7 mg l$^{-1}$) as well as in the phreatic waters (from zero to 150 mg l$^{-1}$ depending on the situation of piezometers, median value of 12 mg l$^{-1}$ (Table 3) were much higher than in the lake water (1 mg l$^{-1}$). In 1985, nitrate concentrations (N-NO$_3$)
just below the pipe (0.37 mg l$^{-1}$, from 0.09 to 0.56 mg l$^{-1}$ in extreme values) higher than those in the lake show that the rainwater pipe was still a source of nitrogen.

There was no coincidence between maximum values of total phosphorus and inorganic nitrogen in the lake.

### 5. Discussion

#### 5.1. Typical seasonal features of the physical and chemical environment

As shown by the vertical distribution of oxygen concentrations, a slight thermal gradient is sufficient to offer a resistance to exchanges between surface and bottom but, mixing events invariably reduced the duration of bottom deoxygenation. The increase in water transparency, favouring a relative increase in photosynthetic activity deeper, has attenuated the strong heterogeneity of oxygen profiles observed at the beginning of the study.

The decrease in transparency during summer for 1979 to 1982 coincided to the maximum of phytoplankton biomass. However, after 1982, the summer phytoplankton biomass was too low to influence the seasonal trend in transparency. Seasonal fluctuations of chlorophyll $a$ concentrations were clearly associated to the main phases of phytoplankton growth and decline, although high chlorophyll values, compared to phytoplankton biomass, were found in winter (Garnier 1989, Lacroix et al. 1989). These wide variations appear to be typical of shallow waters, where mixing events create new conditions (nutrient release) which enhance phytoplankton growth.
The summer decrease in nitrates at the beginning of the study must be related to phytoplankton uptake but also, as anaerobic conditions developed, to denitrification processes important in sandpit lake (Labroue et al. 1988). The other forms of nutrients have not shown any clear seasonal pattern. This could indicate a rapid biological uptake together with rapid exchanges between water and sediment during mixing events.

5.2. Characteristics of the physical and chemical changes over the 8 years studied

From 1979 onwards, the increase in water transparency has been the most significant and directly observable characteristic of the development. Suspended matter (dry weight) have not been continuously analysed but between 1978 (Chestérikoff et al. 1981) and 1985 (Mourelatos 1988), a decrease from 13 to 6 mg l$^{-1}$ as an annual average has occurred. The reduction in suspended matter partly corresponded to a decrease of the organic fraction (a decrease in chlorophyll $a$ concentration from 17 to 3.5 $\mu$g l$^{-1}$ in annual means) throughout the period studied. However, as the decrease in water transparency has been observed as early as 1980 whereas chlorophyll $a$ concentrations began to decrease only in 1982, the biological component alone cannot explain the increase in transparency. It must also be related to a progressive stabilization of bottom sediment and banks, after the quarrying of sand has stopped, thus reducing resuspension by wind and littoral turbulence. In 1986, when the lake water level was lowered, the reversed trend can conversely be explained by erosion of the banks as well as by increased resuspension of the bottom sediments. An indirect but important effect of the increase in water transparency was to improve the oxygenation of the water column.

Whereas direct superficial inputs in nutrients are obviously an important source for eutrophication, the rapid decrease in chlorophyll after nutrient reduction has shown the lake to be very sensitive to restoration.

The diversion of domestic water brought by the rainwater pipe until May 1981 is the main cause for the decrease in phosphorus in the lake. The early decrease in nitrogen, but also in orthophosphates, tends to support the idea of a reduction of nutrients brought by the phreatic waters, after progressive wash out of nutrients from polluted material. Later on in 1983, the regulation of the water level of the lake by a drainage pipe, decreasing the residence time, coincided with an accentuation of oligotrophication, but no effect of residence time can be shown for the nutrient conditions found in the lake in 1983 (Garnier & Billen, in press). Other processes including colonisation of macrophytes could have contributed to a progress in the oligotrophication of the lake.

Internal processes must have also contributed to the elimination of nutrients.

The hard waters of Lake Crétail represented favourable conditions for a transfer of large fraction of the phosphorus input to calcium, explaining the precipitation as calcium phosphates and thus
limiting the increase in phosphorus concentration in the lake (Tiren & Petersson 1985; Herodek & Istvanovics 1986). Whereas a large part of phosphorus was therefore immobilized in the sediments (Bates & Neafus 1980, Bostrom et al. 1982, Foy 1986), another part must be however permanently recycled either under intermittent stratification-destratifications, by mineralization of organic phosphorus under oxic conditions (Kamp-Nielsen 1975, Lee et al. 1977) or by biological uptake and adsorption-desorption processes on suspended matter within the lake water (Capblancq et al. 1986).

Considering nitrogen, denitrification could be a major process for its elimination. In Lake Créteil chiefly at the beginning of the study, the supply with anoxic water and intermittent oxygen depletion together with a high algal biomass in summer represented suitable conditions for denitrification (Barroin 1985, Labroue et al. 1988). Denitrification might partly explain the difference between nitrate concentrations in the lake (1 mg l⁻¹) and in phreatic waters and in waters brought by the pipe (12 and 18 mg l⁻¹ respectively); it is improbable that the difference could have been mobilized by phytoplankton. In 1985, particulate nitrogen for the fraction less than 35 μm were 0.1 mg l⁻¹; taking into account the reduction in phytoplankton biomass (from 17 to 3.5 μg l⁻¹ in chlorophyll a) particulate nitrogen should be less than 1 mg l⁻¹ at the beginning of the study.

References