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Combining Limnology and Palaeolimnology to Investigate Recent Regime Shifts in a Shallow, Eutrophic Lake

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Abstract

In this study, we demonstrate that an integrated approach, combining palaeolimnological records and limnological monitoring data, can increase our understanding of changing ecological patterns and processes in shallow lakes. We focused on recent regime shifts in shallow Lake Krankesjön, southern Sweden, including the collapse of the clear-water state in 1975 and its subsequent recovery in the late 1980s. We used diatom, hydrocarbon, and biogenic silica sediment records, in concert with limnological data sets on nutrient concentrations, water clarity, chlorophyll-a and water depth, to investigate the shifts. The shift from clear to turbid conditions was abrupt and occurred over 1 to 2 years, whereas recovery of the clear-water state was more gradual, taking 4–5 years. In 1978, shortly after the first regime shift in water clarity, the diatom community underwent a significant shift. It

became less diverse, with decreased abundance of epiphytic and planktonic taxa. Despite rising phosphorus concentrations and lower abundance of submerged macrophytes, Lake Krankesjön has remained in the clear-water state over the past 20 years, although this state seems to be increasingly unstable and susceptible to collapse. The complex reactions of the entire lake ecosystem to major changes in lake-water clarity, as shown by the palaeolimnological variables investigated in this study, emphasize the importance of careful lake and catchment management if a stable, clear-water state is desired.

Keywords: regime shift, monitoring, ecosystem, alternative stable state, diatoms, hydrocarbons

Introduction

Abrupt shifts in ecosystem structure and function are often described as "regime shifts," i.e., sudden and large events that lead to rapid ecosystem reconfiguration from one alternative state to another (Lees et al. 2006; Andersen et al. 2009). Regime shifts occur in multiple ecosystem types (Scheffer et al. 2001), including marine environments (Collie et al. 2004; Spencer et al. 2011), Arctic lakes (Smol et al. 2005), brackish lagoons (Jeppesen et al. 2007), terrestrial systems (Dearing 2008), and coastal areas (Conley et al. 2009; Duarte et al. 2009). These shifts are, however, especially well documented in shallow lakes and are often associated with anthropogenic eutrophication (McGowan et al. 2005; Hargeby et al. 2007; Scheffer and Jeppesen 2007; Scheffer and van Nes 2007; Zimmer et al. 2009; Hobbs et al. 2012). The transition of a shallow lake from a clear-water, macrophyte-dominated regime to a turbid, phytoplankton regime has been described theoretically by, for example, Scheffer et al. (1993) and Scheffer and Carpenter (2003). Theory predicts that increasing nutrient concentrations decrease the stability of the clear-water regime until a critical threshold is reached, at which time the lake shifts to the turbid regime. Reestablishment of the clear-water regime is not possible unless nutrient concentrations are reduced substantially and the system displays hysteresis (Scheffer et al. 2001; Andersen et al. 2009). An external perturbation can cause a regime shift in the lake at moderate nutrient levels, but at low and high nutrient concentrations, clear and turbid regimes are the only possible states (Scheffer et al. 1993; Scheffer and Carpenter 2003).

Lake Krankesjön, in southern Sweden (fig. 1), is a shallow, eutrophic lake with a history of structural shifts between two alternative states, one clear and macrophyte-dominated and the other turbid and algal dominated, in spite of supposedly very little change in external nutrient loading (Blindow et al. 2006). These shifts have been documented for almost a century, in reports on the abundance of herbivorous waterfowl and notes on water clarity (Swanberg 1931; Almestrand and Lundh 1951; Thomasson 1956; Forsberg 1964; Hansson et al. 2010), which are summarized in figure 2b. During the 1920s, there were dense beds of charophytes in the shallow lake, which continued to be dominated by charophyte vegetation until the mid- 1940s, after which there was a short period when submerged plants were absent (Almestrand and Lundh 1951; Blindow 1992b). By 1948, the submerged vegetation had recovered in Lake Krankesjön, and in the 1950s and 1960s submerged vegetation was well developed (Karlsson et al. 1976). A shift to the turbid state occurred in the mid-1970s, and the lake had sparse submerged vegetation for almost a decade (Blindow et al.

2006). In the mid-1980s, submerged vegetation rapidly started to recolonize the lake, and since the late 1980s, the lake has been in a clear-water state, with abundant macrophyte beds. Periods of temporary instability of the clear-water state, however, occurred in 1997 and 1999 (Blindow et al. 2002), and most recently, during 2011–2012 (Brönmark, C. and Hansson L-A., unpublished data).

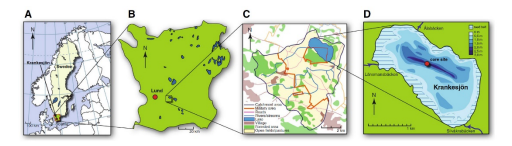


Figure 1. (A) Map of Scandinavia showing the location of Lake Krankesjön in southern Sweden. (B) Map of the province of Scania. (C) Map of the catchment area surrounding Lake Krankesjön. (D) Close-up map of the lake, showing inlets, outlet, bathymetry, and sediment sampling site.

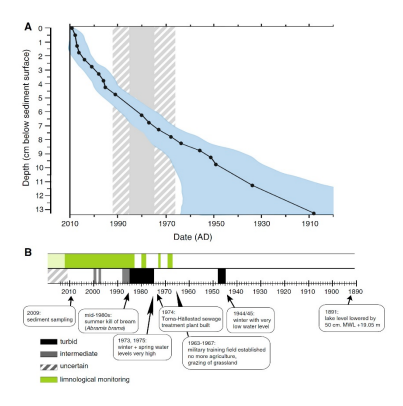


Figure 2. (A) Age-depth model for the Krankesjön sediment sequence based on 210 Pb dating. The *blue shading* indicates dating uncertainty envelopes, with 2σ error. The *grey shading* indicates a period with turbid conditions, with the *dashed shading* indicating the dating

uncertainty. (B) Overview of the development of Krankesjön during the studied time period. Data from Blindow (1992a), Hargeby et al. (2007), The Swedish Fortification Agency (Fortifikationsverket), and recent monitoring.

Lake Krankesjön is considered a waterfowl habitat of international importance and is included in both the international Ramsar Convention of Wetlands and the EU network Natura 2000. The clear-water state of the lake is generally preferred from a human perspective because greater water transparency, higher biodiversity, and fewer problems with toxin-producing phytoplankton produce higher recreational value. The clear-water state is also favored from a conservation standpoint for managing waterfowl in Lake Krankesjön.

Sayer et al. (2010) suggest that the combination of contemporary ecology and palaeolimnology can be a powerful means to assess the dynamics of shallow-lake ecosystems on multiple timescales. Records of multiple variables obtained from sediment successions can be used to pose and test ecological theories across time scales beyond the reach of monitoring data, whereas studies of contemporary aquatic ecology provide insights into habitat preferences, seasonality, and interactions of organisms. The combination of long-term observational data sets and recent sediment records can yield long-term records of lake histories (Battarbee et al. 2005). In this study we combined limnological monitoring data with palaeolimnological diatom, hydrocarbon, and biogenic silica records to make an integrated assessment of the shifts between turbid- and clear-water states in Lake Krankesjön. We focused on the regime shifts during the mid-1970s and mid-1980s, and examined how they were recorded by the monitoring data and whether they were recorded in the sediment. Considering that regime shifts have been documented since the 1920s in Lake Krankesjön, we focused on the recent lake sediments, covering the period from ca. 1900–2009, which enabled us to compare the sediment record to the limnologically documented changes.

Materials and methods

Site description, core collection, and chronology

Lake Krankesjön is situated in the southernmost part of Sweden (55°42′N, 13°28′E), approximately 20 km east of the city of Lund (fig. 1). The lake is shallow (maximum depth 3.2 m, median depth 1.5 m), with a water retention time of 2.3 months, and it is moderately eutrophic and calcium rich. It has a surface area of 3.4 and a 53-km² catchment that is dominated by arable land (59%), pastures and open fields (17%), and forest (16%). Much of the land in forest and open fields is part of a military training area that surrounds the lake. The lake has two inflows, the River Länsmansbäcken, which enters from the west, and the River Silvåkrabäcken, which enters from the southeast. The River Ålabäcken is the outlet, in the northern part of the lake.

A 76-cm sediment sequence was collected from the ice in February 2009, using a crust-freeze sampler (Renberg 1981). The frozen sediment sequence was subsampled at the Department of Ecology and Environmental Science at Umeå University into 151 contiguous samples, each 0.5 cm thick, except the uppermost 1-cm-thick sample, using methods described by Renberg (1981).

The uppermost 14 cm of the sediment sequence was analyzed, providing 27 samples that were freeze-dried and homogenized prior to other analyses. The sequence was dated by measurement of ²¹⁰Pb, ²²⁶Ra, and ¹³⁷Cs, using gamma spectrometry at the Gamma Dating Centre, Institute of Geography, University of Copenhagen. ²¹⁰Pb ages were calculated using the constant rate of supply (CRS) model (Appleby 2001).

Diatom analysis

Approximately 0.01 g of freeze-dried sediment was digested using the water-bath technique with 30% H₂O₂ (Battarbee et al. 2001). Diatom concentrations (valves mg⁻¹) were determined by adding DVB (divinylbenzene) spheres to the cleaned samples (Battarbee and Kneen 1982). Cleaned subsamples were dried onto cover slips and mounted onto microscope slides using Zrax (R.I~1.7+). At least 600 diatom valves (minimum 200 non-Fragilariaceae) were counted in each subsample under a light microscope at ×1,000 using phasecontrast optics. Diatom taxonomy followed Krammer and Lange-Bertalot (1986, 1988, 1991a, b), Williams and Round (1987), Lange-Bertalot and Metzeltin (1996), Round and Bukhtiyarova (1996), Lange-Bertalot (1999), and Håkansson (2002). In total, 72 taxa were identified. All diatom data are expressed as relative percent abundances and were reduced to 13 taxa. Only taxa with abundances exceeding 3% of the total were included in the statistical analyses. Fragilarioid taxa comprised up to 89% of the valves counted in each sample, and therefore the non-fragilarioid taxa were also examined separately, using the 24 taxa with abundances exceeding 3% of total non-fragilarioid taxa. The diatom accumulation rate (DAR) and Hill's N2 diversity index (Hill 1973) were calculated using all identified taxa.

Geochemical analyses

Hydrocarbons were extracted from approximately 1–2 g of freeze-dried sediment with a mixture of dichloromethane and methanol (9:1 v/v). A Dionex Accelerated Solvent Extractor 300 was used (programmed for three extraction cycles at 1,000 psi and 100°C). The lipid extracts were reduced in volume by using a Büchi Rotovapor and injected in pulsed splitless mode into an Agilent 6890 gas chromatograph equipped with a CIS-4 Gerstel inlet and a DB5- MS column (30 m × 0.25 mm i.d. × 0.25 μ m film). Oven temperature was held at 35°C for 6 min, increased to 300°C at 5°C min⁻¹ and held there for 20 min. The chromatograph was interfaced with an Agilent 5973 mass spectrometer operated at 70 eV in full-scan mode (m/z 50–500 amu). External and internal standards (S-4066 from CHIRON, Norway and deuterated perylene from Cambridge Laboratory, USA) were used for quantification.

Biogenic silica was analyzed following the wet chemical digestion technique described by Conley and Schelske (2001). Results are presented as wt% SiO₂ of total dry weight.

Monitoring data

Lake Krankesjön has been the focus of numerous research projects, as well as national and regional monitoring, since the late 1960s (fig. 2b). We obtained and merged data sets from the Swedish University of Agricultural Sciences and the Aquatic Ecology Unit of the Department of Biology, Lund University, and selected variables that aid in describing and

understanding the shifts between clear and turbid states (fig. 3). Sampling was less consistent at times, and the variables analyzed have differed, but since the mid-1980s, the sampling protocol has been fairly stable. Before 1983 there was only one water quality sample collected per year, in July–August. Water-level data (year-round monthly measurements) from 1946 to present (excluding 1990–1994) were obtained from the County Administrative Board of Skåne (Länsstyrelsen) and the Swedish Fortification Agency (Fortifikationsverket).

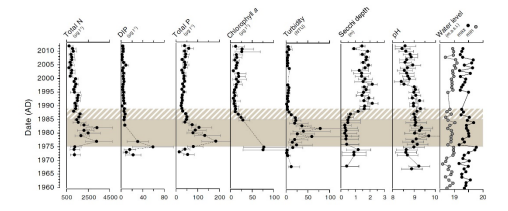


Figure 3. Yearly means (Mar–Sep) of monitoring data from Lake Krankesjön, and water level range (min and max) from The County Administrative Board of Skåne (Länsstyrelsen) and the Swedish Fortification Agency (Fortifikationsverket). The *brown shading* indicates turbid conditions with the *dashed shading* indicating intermediate conditions. *Error bars* mark the 95% confidence interval of the means.

Statistical analyses

Yearly means (March–September) of nutrients, chlorophyll-a, turbidity, pH, and Secchi depth were calculated from the monitoring data (all log-transformed except pH and Secchi depth) as marginal means from a General Linear Model that accounted for differences in months of sampling across years (Carstensen et al. 2006). Means of log-transformed variables were back-transformed using the exponential function to represent geometric means.

Stratigraphic diatom assemblage zones (DAZ) were determined through regression tree analysis (De'ath and Fabricius 2000), using the mvpart package (De'ath 2012) for the R statistical software. The smallest tree within one standard error of the tree with the lowest cross-validation relative error was chosen, producing two zones. A detrended correspondence analysis (DCA), performed using the vegan package (Oksanen et al. 2012) in R, revealed that the gradient length in the diatom data was short (–2 SD units), and therefore principal component analysis (PCA) was performed on Hellinger-transformed diatom data using the same package. One PCA axis was identified as explaining significant proportions of the variance in the stratigraphic data when compared to the broken stick (null) distribution. This axis explained 46% of the variance.

Results

Chronology

Unsupported ²¹⁰Pb activity in the sediment sequence is generally low, with activities consistently <0.19 Bq g⁻¹ dry material, which could indicate temporary erosion at the core site. Low activities, in combination with small sample sizes, led to rather large error bars on the radiometric data (fig. 2a). Application of the CRS dating model (Appleby 2001) to the unsupported ²¹⁰Pb data, constrained by ¹³⁷Cs, suggests a mean sedimentation rate of 0.15 cm year⁻¹ over the last ~100 years, with increasing rates, ranging from 0.4 to 0.7 cm year⁻¹, in the surface sediments (0–2 cm depth). Sampling resolution in the core is, on average, ~3 years sample⁻¹.

Monitoring data

During the 1960s and early 1970s, Lake Krankesjön was in a clear-water state (fig. 2b), but by 1975 submerged vegetation was completely absent (Karlsson et al. 1976). This was accompanied by a shift to turbid conditions. Secchi depth decreased from 0.9-1.2 to 0.3-0.5 m and turbidity increased (fig. 3). All nutrient concentrations increased along with the shift to the turbid state, in particular dissolved inorganic phosphorus (DIP), which reached 61 lgL-1 in August 1975 and total phosphorus (TP), which peaked at 186 μ g L⁻¹ in August 1977. These increases in lake-water phosphorus content suggest large releases of ironbound phosphorus from sediments, as well as release associated with decomposition of organic matter following the decline of submerged macrophytes. DIP concentrations had returned to low levels at the end of the turbid phase (although there are no monitoring data for the period 1977–1983) and total phosphorus and total nitrogen concentrations remained high until the early 1980s. Around 1984, Potamogeton pectinatus and characeans started to recolonize the lake, and from 1985 recolonization was rapid and the lake shifted to a clear-water state (Blindow 1992b). This is evident from turbidity values, which were high during the turbid state and decreased over 3-4 years following macrophyte recolonization. Secchi depth similarly increased to ~1.5 m during those years and has remained between 1.5 and 2.0 m to the present (fig. 3). TP concentrations increased around 1975 and then decreased during the remainder of the turbid state. After that, total phosphorus remained low, but displayed an increasing tendency since 1990. On the other hand, total nitrogen concentrations decreased steadily since the turbid state. Chlorophyll-a levels were not measured during the turbid period but declined significantly during 1985-1990 relative to measurements taken in 1974 and 1975. After 1990, chlorophyll-a increased, similar to total phosphorus. Water level was unusually high prior to the shift from clear-water to turbid conditions in 1975 (fig. 3), especially during spring and summer (not shown).

Palaeo-data

The diatom community of Lake Krankesjön was completely dominated by benthic species (93–99% relative abundance) throughout the investigated period. Thus, benthic primary production seems to have been an important contribution to whole-lake primary production, in both the turbid and the clear-water states. The studied sediment section was divided into two zones at ca. 1978 (fig. 4), with different diatom community structures

(Diatom Assemblage Zones; DAZ). The earlier zone (DAZ1) shows a lower abundance of diatoms but a more diverse flora, whereas the later zone (DAZ2) shows slightly increasing diatom abundance but lower diversity.

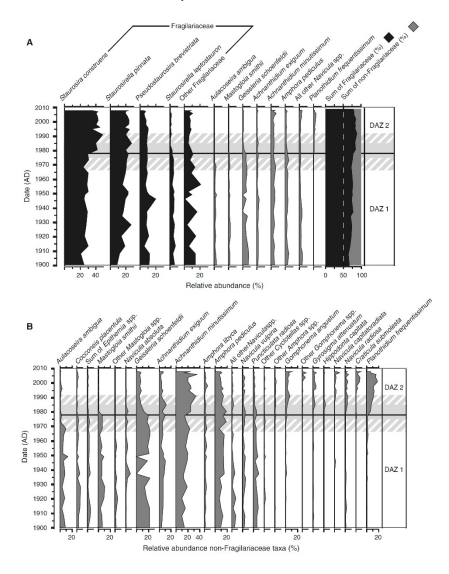


Figure 4. (A) Diatom stratigraphy of the Lake Krankesjön sediment sequence, showing the major (>3%) taxa as relative abundance (%), diatom assemblage zones (DAZ), and proportion of Fragilariaceae and non-Fragilariaceae taxa (%). (B) Major (>3%) non-Fragilariaceae taxa as relative abundance (%) within the non-Fragilariaceae community. *Grey area* turbid conditions, *hatched* dating uncertainty, as in figure 2.

Daz1 (1900-1978)

Fragilariaceae make up most of the diatom assemblage (65–76%) in this zone, with *Staurosira construens* (Ehr.) Williams & Round 1987, *Staurosirella pinnata* (Ehr.) Williams & Round

1988, Pseudostaurosira brevistriata (Grunow in Van Heurk) Williams & Round 1987, and Staurosirella leptostauron (Ehr.) Williams & Round 1987 the most abundant fragilarioid species (fig. 4a). The non-fragilarioid community is dominated by Geissleria schoenfeldii (Hustedt) Lange-Bertalot & Metzeltin 1996, Achnanthidium minutissimum (Kützing) Czarnecki 1994, Amphora pediculus (Kützing) Grunow in Schmidt 1875, Achnanthidium exiguum (Grunow) Czarnecki 1994, Mastogloia smithii Thwaites 1856, and planktonic Aulacoseira ambigua (Grunow) Simonsen 1979, and Puncticulata radiosa (Lemmerman) Håkansson 2002 (fig. 4b). Diatom productivity and abundance are often estimated from biogenic silica (BSi; Conley and Schelske 2001; Struyf and Conley 2009), and these concentrations are low throughout the zone (1.8–2.9 wt%; fig. 5). Similarly, diatom accumulation rates (DAR), another estimation of productivity and/or preservation of diatoms, are relatively low throughout the zone, with a DARtotal of 40–320 × 10⁴ valves cm⁻² year⁻¹ (fig. 5).

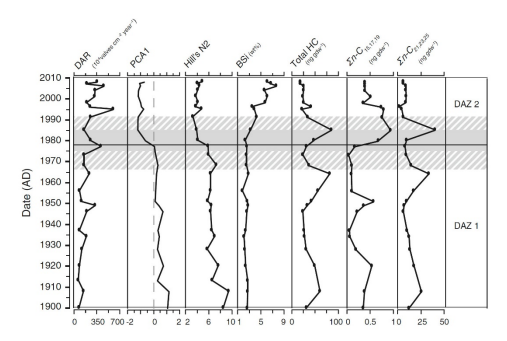


Figure 5. Diatom accumulation rates (DAR), diatom PCA axis 1 scores, Hill's diversity index N2, wt% biogenic silica, total hydrocarbon concentration (ng per g dry weight), concentration of low and medium odd numbered *n*-alkanes (C_{15,17,19} and C_{21,23,25}). Diatom assemblage zones (DAZ) and *grey shading* follow figure 4.

The hydrocarbon compositions of many algae and cyanobacteria are dominated by low molecular weight n-alkanes (n-C_{15,17,19}). In contrast, submerged and floating-leaved macrophytes have n-alkane distribution maxima at C₂₁, C₂₃, or C₂₅ (Ficken et al. 2000; Meyers 2003). Σn -C_{15,17,19} values are around 0.4–0.5 ng gdw⁻¹ (gram dry weight) during the 1910s–1920s (fig. 5), decline in the late 1920s and remain low (0.04–0.16 ng gdw⁻¹) throughout the zone, with the exception of a rise in the mid-1940s, a peak at 0.57 ng gdw⁻¹ around 1950,

and a rapid decline thereafter. The concentration of Σn -C_{21,23,25} follows largely total hydrocarbon (HC) concentration variations, with fairly stable values between 7 and 18 ng gdw⁻¹ throughout the zone, except for a minor peak of 25 ng gdw⁻¹ around 1908 and a more prominent peak of 33 ng gdw⁻¹ in the mid-1960s (fig. 5).

Daz2 (1978-2009)

The proportion of fragilarioid taxa increases slightly after ca. 1978, to between 78 and 89%, but the dominant species remain the same. In the non-fragilarioid community, *Achnanthidium minutissimum*, *A. exiguum*, and *Amphora pediculus* remain dominant, whereas *G. schoenfeldii*, *M. smithii*, and the two dominant planktonic species, *A. ambigua* and *P. radiosa*, decrease markedly and are replaced by, above all, *Planothidium frequentissimum* (Lange-Bertalot) Round & Bukhtiyarova 1996, *Gomphonema angustum* Agardh 1831, *Navicula radiosa* Kützing 1844, and *Craticula submolesta* (Hustedt) Lange-Bertalot 1996 (fig. 4). The BSi content increases throughout the zone (from 2.4 wt% to a maximum of 7.7 wt% in 2006), indicating increased diatom productivity/abundance (fig. 5). Diversity, as measured by Hill's N2 index, declines at the transition from DAZ1 to DAZ2 and remains low throughout this zone, with a slight peak ca. 1995 (fig. 5). The change in sediment diatom composition is also visible on PCA axis 1, where a shift from positive to negative values occurs at the beginning of DAZ2 (fig. 5).

The Σn -C_{15,17,19} alkane concentrations show a second rapid increase starting at the beginning of DAZ2, with a peak value of about 0.94 ng gdw⁻¹ in 1985. The values remain high for nearly 10 years before declining around 1998 to values around 0.40 ng gdw⁻¹ (fig. 5). Σn -C_{21,23,25} values are low (2–10 ng gdw⁻¹), with the exception of one extreme value of 40 ng gdw⁻¹ around 1985.

Discussion

The historically described and previously investigated regime shifts in Lake Krankesjön from clear to turbid conditions in 1975, and turbid to clear conditions in the mid-1980s, are apparent in the monitoring data (fig. 3) but rather unclear in the palaeolimnological data (figs. 4, 5). The shift to turbid conditions in 1975 was registered as a rapid decrease in Secchi depth, an increase in turbidity, and increases in total nutrient concentrations (fig. 3). At roughly the same time, Σn -C_{15,17,19} values increased. Because low molecular weight n-alkanes dominate the hydrocarbon composition of most aquatic algae (i.e., both diatoms and green algae) and photosynthetic bacteria (Choudhary et al. 2010), the increase in these compounds, combined with low diatom productivity, as indicated by low BSi values, could reflect dominance of cyanobacteria and green algae during most of the turbid state. The temporary increase in DAR, suggesting increased dominance of diatoms around 1977 (fig. 5), may be connected to the high levels of dissolved inorganic phosphorus (DIP), which occurred at the beginning of, or possibly just before, the turbid period (fig. 3). DIP concentrations are usually near detection limits in Lake Krankesjön, but in 1975 a considerable phosphate release occurred, which could be linked to the massive loss of submerged macrophytes and their subsequent decomposition. In addition, the enhanced accumulation of organic matter may have reduced redox conditions in the sediments, thereby enhancing

releases of iron-bound phosphate. As indicated by the HC, DAR, and BSi data, cyanobacteria and green algae, which can maintain relatively high growth rates at low light intensities (Mur et al. 1999), seem to have outcompeted the diatoms and remained dominant throughout the turbid period, a pattern commonly seen under such conditions (Brönmark and Hansson 2005).

Within the diatom community, a significant shift occurred ca. 1978. Both PCA and regression tree analysis data show that the diatom assemblage of DAZ2 differs significantly from the previous assemblage (fig. 4). At that point, periphytic G. shoenfeldii, M. smithii, and Cocconeis placentula decreased in abundance along with planktonic species A. ambigua and P. radiosa. The periphytic forms P. frequentissimum, G. angustum, C. submolesta, and S. construens increased at the same time. Whereas the decline in plant-attached species such as C. placentula can be interpreted as a direct response to the loss of submerged macrophytes, and thus the loss of habitats for epiphytic taxa (Sayer et al. 2006), increased turbidity and hence reduced light conditions may have been important in inhibiting planktonic production. The decline of P. radiosa and A. ambigua, species associated with mesotrophic rather than strongly eutrophic waters (Bennion and Appleby 2001), may also suggest eutrophication, and the diatom assemblage changes may in fact suggest slow eutrophication, with the macrophyte loss in the mid-1970s acting as a perturbation, which caused the apparent regime shift detected in the sediment data. As the diatom community changed, diversity, recorded by Hill's N2 diversity index, declined, indicating a less diverse diatom community post-1978. This, too, could be an indication of the slow eutrophication of the lake (Bennion et al. 1996, 2000).

The shift back to a clear-water state in the late 1980s was more gradual, with Secchi depth increasing and turbidity decreasing over a period of ca. 4–5 years, from 1985, when macrophytes were reestablished in the lake (fig. 3). The shift to the clear-water state is, however, not easily identified stratigraphically, and it appears that the lake did not revert to the same clear-water conditions that prevailed prior to the turbid period. Increasing values of BSi and decreasing concentrations of low molecular weight *n*-alkanes imply lower influence of cyanobacteria and green algae and greater influence of diatoms in the basic food web during the past 20 years. The diatom record shows that as water clarity improved and diatom production increased, the community that was established around 1978 persisted into the clear-water state. Within the cyanobacterial community, however, large changes occurred in 1988, when small, noncolonial cyanobacteria were replaced by colonial, toxic *Microcystis botrys* and *M. aeruginosa*, as well as by the golden alga *Dinobryon sociale* (Blindow et al. 2006).

By combining monitoring and sediment data across the shifts from clear-water to turbid conditions, and back again from the mid-1970s to the mid-1980s, we found that regime shifts that were very apparent in the monitoring data were not clearly reflected in the palaeolimnological record. It appears that these regime shifts between clear and turbid states, which in some sense have become the model of regime shifts in shallow lakes (Scheffer et al. 1993, 2001), mainly affected the macrophyte community and higher food web of Lake Krankesjön, but not the entire, complex lake ecosystem. The diatom community was not clearly affected by the changes in water clarity and the record instead shows an underlying, slower change towards more eutrophic conditions, possibly amplified by the die-off

of macrophytes in the mid-1970s. This eutrophication could have acted as an underlying mechanism behind the major regime shift, reducing the resilience of the system and making the clear-water state more vulnerable, hence matching the "classical" pattern of regime shift in shallow lakes (Scheffer et al. 1993, 2001; Scheffer and van Nes 2007). There are, however, suggestions that shifts between turbid- and clear-water states in shallow lakes in general are gradual and not necessarily sudden processes (Scheffer and van Nes 2007; Sayer et al. 2010). van Nes et al. (2007) proposed a theory for cyclic shifts between alternative states in some shallow lakes, whereby submerged plants create a positive feedback for their own growth conditions by clearing the water but in the long run undermine their position as the dominant primary producers by increasing nutrient retention and creating slow "internal eutrophication." This results in periods of low resilience, when a weak disturbance can trigger a shift to the turbid state. Hargeby et al. (2007) argued that this behavior may be a factor behind the shifts in Lake Krankesjön and could explain the difficulties in identifying the specific causes behind the shifts in the lake.

Using additional sediment proxies for past limnological conditions, such as plant macrofossils and zooplankton remains, the natural lake state and variability, as reflected in the sediment record, might be better understood. It might make it possible to determine if and why the entire lake ecosystem did not respond to the major changes in water clarity. Extension of well-dated records beyond the last century could also shed light on whether the recent shifts between clear-water and turbid conditions are anomalies or part of a natural, cyclic lake behavior. This, however, might prove challenging because the lake is calcareous and terrestrial macrofossils in the sediment are scarce.

In recent years, Lake Krankesjön seems to be changing, with increasing TP levels and lower abundance of characeans (fig. 3). This suggests increasing instability and lower resilience of the current clear-water state, meaning that the probability of a shift to turbid conditions has increased. We posit that the combination of contemporary and palaeolim-nological data yields a fuller understanding of changing ecological patterns and processes in shallow lake ecosystems. In the present case, palaeolimnological data provide insight into components of the ecosystem that were not measured in contemporary surveys and that were generally stable, despite shifting water clarity and macrophyte cover.

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