

Paleoclimate changes inferred from stable isotopes and magnetic properties of organic-rich lake sediments in Arctic Norway

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Abstract Stable isotope measures in organic matter are frequently used as indicators of past climate change. Although such analyses can provide valuable information, there is considerable uncertainty associated with studies of organic-rich sediments, especially those from Arctic lakes and bogs. We studied stable isotopes of carbon and nitrogen, and magnetic properties in a sediment core from a small alkaline lake with a high sedimentation rate, Lake Nattmålsvatn, Norway. There is good correspondence among the different sediment variables during the late

glacial, and they seemingly reflect major climate variations such as the Allerød Interstade and the Younger Dryas, as well as the transition into the current interglacial. During the early Holocene, however, these relationships are more complex and $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values do not stabilize until $\sim 7,500$ cal year BP. A significant excursion in all variables occurs between 6,850 and 6,500 cal year BP and is interpreted to represent climate deterioration. Holocene $\delta^{13}\text{C}$ values vary little and indicate that isotopically-depleted dissolved inorganic carbon (DIC) in the lake, possibly influenced by methanotrophy and high pCO_2 , dominated the lake's carbon cycle. Holocene $\delta^{15}\text{N}$ is similarly muted, likely due to the availability of abundant dissolved nitrogen. Bulk organic matter is probably dominated by phytoplankton remains produced beneath the ice cover in late spring and during ice breakup when isotopically-depleted DIC, pCO_2 and ammonium availability were maximal. Thus, use of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ as indicators of Holocene paleoclimate and paleoproductivity variation can be challenging in a lake such as Nattmålsvatn, where ice cover isolates the basin for large parts of the year, allowing dissolved respiratory gases to accumulate in the water column. In contrast, magnetic variables appear to better track climate variations. In particular, runoff-driven influx of minerogenic sediments shows high variability that can be attributed to regional changes in Holocene winter precipitation. The most striking shifts occur between 4,000 and 2,300 cal year BP.

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Introduction

Small lakes respond quickly to environmental changes, adjusting immediately to seasonal temperature shifts, catchment runoff, and spring snow melt. Responses may include algal blooms, precipitation of authigenic calcite, and changes in the composition of the chironomid community. These combined seasonal influences, in addition to external factors, determine the variations in sediment organic matter over time.

In organic-rich lake sediments, organic matter (OM) content can be assessed by weight loss on ignition (LOI) or by measure of total organic carbon (TOC), the latter a measure of the carbon in OM (Dean 1974). Because bulk organic matter has multiple potential sources within and outside the lake, other analyses are required to interpret shifts in OM content. The residual (inorganic) component of the sediment, whether detrital, biogenic silica (diatoms), authigenic carbonate, or a combination thereof, should also be determined.

When organisms die, their remains are subject to microbial decomposition. Net loss of organic matter can be great, and there may be alteration of the primary material. Despite these transformations, the carbon/nitrogen ratio (C/N) and $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values of bulk organic matter can be used to track the origin of the OM, providing information about the relative contributions of autochthonous versus allochthonous OM to the deposits (Meyers and Lallier-Vergés 1999; Talbot and Johannessen 1992; Filippi and Talbot 2004). Furthermore, the magnetic properties of minerogenic matter that reaches the lake can be used to constrain the external influx of OM identified from isotope measures. Inferring the minerogenic input by means of rock magnetism can also be used to track flood history and other important past environmental events in the catchment (Thompson and Oldfield 1986; Evans and Heller 2003).

The ability of lakes to respond rapidly to environmental changes and to record those changes in their sediment makes lacustrine deposits attractive archives for study of high-frequency paleoclimate oscillations (Bradley 1999). Coastal lakes in Scandinavia are especially suitable for such purposes, as they generally

display high sedimentation rates. Many of these lakes were beyond the ice sheet margin during the Younger Dryas (12,900–11,700 cal year BP), suggesting that their sediment records may extend beyond the last glacial-interglacial transition. Several studies have shown that these waterbodies are sensitive to climate fluctuations and that their sediments record regional climate changes (Vorren et al. 1988; Birks et al. 2005; Bjune et al. 2005; Paasche et al. 2007a, b). To our knowledge, no previous study in northern Norway has used carbon and nitrogen isotopes in combination with rock magnetic properties to infer past climate changes on a sub-centennial time scale.

This study addresses the prospects and problems associated with using carbon and nitrogen isotopes as indicators of past climate change in high-latitude lakes with organic-rich sediments. We conducted analyses on a well-dated lacustrine record that spans the entire Holocene and late glacial, at multi-decadal resolution. In addition to recording the withdrawal of the inland ice sheet and/or a local ice cap, the sediment record documents several distinct climatic phases during the Holocene, including a prominent climatic excursion $\sim 6,700$ cal year BP. The challenges associated with using $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ in organic-rich lake deposits are highlighted and we emphasize how methanogenesis and seasonal ice cover may dampen the climate sensitivity of the lake.

Study area

Lake Nattmålsvatn is located in the southeastern part of the large island of Senja, northern Norway (Fig. 1), 170 m above sea level and 1 km from the coast (69°10'45.58"N 17°43'23.40"E). The lake lies outside the marginal moraines of the Younger Dryas (Andersen 1968) as well as those of the Older Dryas, $\sim 14,450$ cal year BP (Sveian pers. commun.). The catchment is composed of Ordovician and Silurian bedrock consisting of garnet mica schist and calcite marble, the latter being dominant. The large amount of carbonate rock, which is mostly calcite, but also contains small amounts of dolomite, contributes to the high pH and “hardness” of surface water runoff entering Lake Nattmålsvatn. The lake has a summer (July) pH of 8.0 (A. Langangen, pers. commun.). Maximum water depth is ~ 13 m and the lake is 200 m wide and 250 m long.

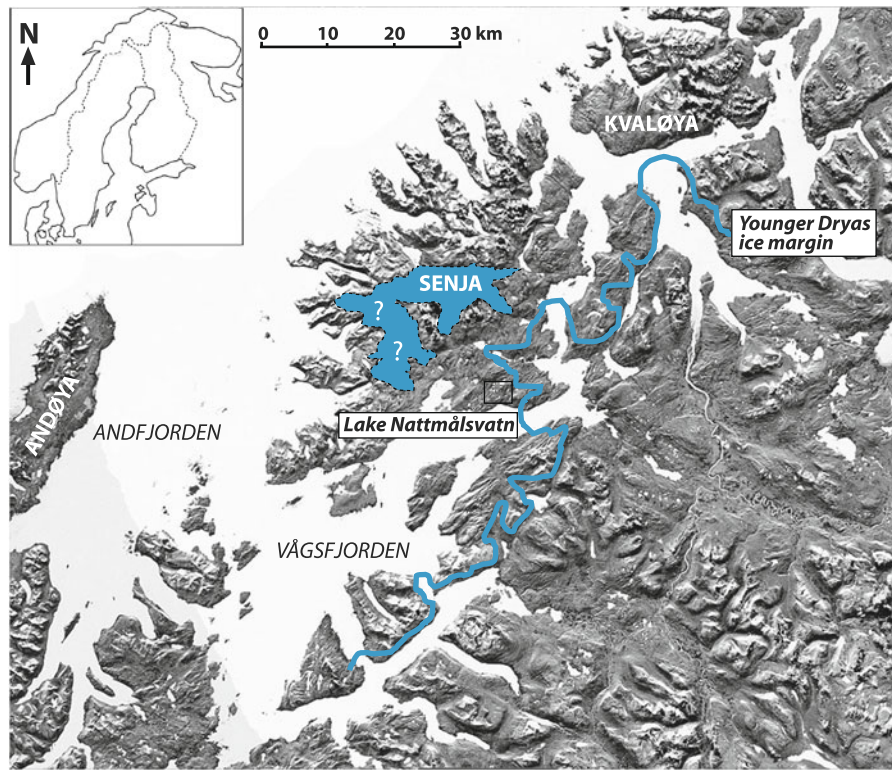


Fig. 1 Outline of Scandinavia (*upper left*) and satellite image of the study site and its catchment. Lake Natnmålsvatn is situated on the large island Senja (1,586 km²) in northern Norway. The blue lines show the reconstructed ice-sheet margin of the Fennoscandian ice sheet during the Younger

Dryas (~12,500 cal year BP), whereas the dotted blue line indicates a presumed advance of local glaciers on Senja according to Vorren and Plassen (2002), possibly older than the Younger Dryas

The nearest meteorological station lies 11 km to the northeast, at Finnsnes. For the period 1961–1990, it recorded a mean annual temperature of 3.2°C, a July mean of 12.5°C and a January mean of –4.3°C. Lakes in the region are commonly ice covered from October/November until late May. Mean annual precipitation is approximately 1,000 mm, of which ~35% falls from May to September, while ~65% falls from 1 October through April (Klimaavdelingen 1993a, b). Chemical and meteorological data are presented in Table S1.

Lake Natnmålsvatn is currently considered an alkaline *Chara* lake, with charophytes and marl precipitation during summer (Langangen 1995, 2004). Marl is common in the shallow waters and along the shore. During earlier centuries, local farmers collected the marl for their fields, leaving

holes along the banks. Marl lake sediments are excellent archives of environmental change (Kelts and Hsü 1978; Hollander and McKenzie 1991; Bradbury and Dean 1993; Noe-Nygaard 1995; Hammarlund et al. 2003; Teranes et al. 1999; Leng and Marshall 2004; Diefendorf et al. 2008) and one of our original objectives in this study was to compare the isotopic records from the carbonate and OM fractions in the Lake Natnmålsvatn sediment. Two cores retrieved from the deepest part of the lake, however, yielded almost 6 m of blackish, organic-rich sediments with no apparent carbonate. The cores also smelled of H₂S. Despite the fact that Lake Natnmålsvatn is today an alkaline *Chara* lake according to pH measurements, there is no evidence in the sediment record for its present state. Our isotope analyses were thus carried out only on the sediment OM fraction.

Materials and methods

Core collection and LOI and carbon analyses

Two cores, designated Core I and Core II, were taken ~100 m apart in March 2002. The cores were collected from the deepest part of the lake using an 11-cm-diameter piston corer (Nesje 1992) operated from surface ice. Both cores were sampled for LOI₅₅₀ at 0.5-cm intervals by extracting 1 cm³ of wet material. Samples were weighed, dried overnight at 105°C, and re-weighed to assess percent dry weight. Next, samples were subjected to rising temperatures for half an hour and then combusted at 550°C for 1 h in a Nabertherm controller C6 muffle furnace. After ignition, samples were cooled in a desiccator and re-weighed to estimate OM content (Dean 1974). Twenty samples were analysed for TOC using a LECO furnace.

Smear slides

A full set of smear slides was prepared from Core I to examine the organic material in the sediment. Smear slides were prepared from samples taken every 2–10 cm, except across identifiable sedimentological boundaries, where closer-interval samples were taken. Small samples of moist sediment were prepared for visual examination under a petrographic microscope. Smears were mounted beneath a cover slip using standard epoxy adhesive. Plane- and cross-polarised light and incident UV radiation were used to examine the slides. Analysis showed that sediments are composed of a variety of minerogenic and organic components, including authigenic minerals, notably pyrite and feldspar at some depths in the core.

Isotope and elemental analysis

Core I was subsampled every 4 cm for organic carbon ($\delta^{13}\text{C}$) and nitrogen ($\delta^{15}\text{N}$) isotope analysis of bulk organic matter (OM), total organic carbon (TOC) and total nitrogen (TN) content. Sample aliquots were reacted with 1 M HCl at 50°C to remove possible traces of carbonate. After 24 h, samples were rinsed in distilled water five times and dried in an oven at 40°C. Dried samples were then crushed in an agate mortar to homogenize them and reduce grain size. There has been concern about

whether components of the nitrogen isotopic composition are lost when samples are reacted in HCl and oven-dried (Meyers and Teranes 2001). We thus ran 20 freeze-dried and homogenised bulk samples from the same levels that were prepared using HCl and by oven-drying, to test if there were differences between results derived from the two preparation methods (Table S2). Because we found no large discrepancy between the methods, the HCl treatment was applied to the nitrogen isotope samples throughout the core. All isotope samples were analysed according to standard methods at Iso-Analytical Inc., England, with an elemental analyser connected to a continuous flow isotope-ratio mass spectrometer. Analytical precision was $\pm 0.3\%$ for $\delta^{15}\text{N}$ and $\pm 0.1\%$ for $\delta^{13}\text{C}$. The elemental analyser provides measurements of total organic carbon (TOC) and nitrogen content (TN). There was no evidence for presence of inorganic nitrogen (Talbot 2001), and TN thus reflects total organic nitrogen. In this study, C/N ratios are expressed as weight percent. Carbon isotope values are reported in standard δ notation (‰) relative to VPDB (PeeDee Belemnite) (Coplen 1982), and nitrogen values are reported with respect to atmospheric nitrogen (AIR).

Magnetic variables

Magnetic susceptibility (χ_{bulk}), anhysteretic remnant magnetization (ARM), and saturation isothermal remnant magnetization (SIRM), were measured to characterize the minerogenic material deposited in the lake. Such measurements permit discrimination between, for instance, glacial and non-glacial sediments, as reflected in changes in the strength of the signal or shifts in the S-ratio ($\text{IRM}_{100\text{mT}}/\text{SIRM}_{2\text{T}}$). Such data can be used to identify the end of deglaciation and to characterize the input of detrital material throughout the Holocene.

Core I was sampled for magnetic analyses with plastic boxes every 1 cm throughout the core ($n = 489$). Initial magnetic susceptibility (χ_{bulk}) was determined on both wet and dry samples using a KLY-2 induction bridge with a sensitivity of 4×10^{-8} SI. ARM was imposed with a 0.1 mT DC field and a 100 mT AC-field in a 2G af-demagnetizer on contiguous samples collected from the basal 89 cm of the core (489–400 cm depth; $n = 89$). SIRM was obtained for the same interval (489–400 cm depth;

$n = 89$) using an ASC IM10-30 Impulse Magnetizer for which saturation levels (SIRM) were assumed by applying a 2T field. Because there was little difference between measurements performed with 1T and 2T fields ($n = 20$), we assume that “saturation” was obtained with a 2T field. Backfield measurements were performed after exposing the samples to a 100mT field (IRM_{100mT}). Subsequently, the S-ratio was used to indicate the dominant magnetic mineralogy (Stober and Thompson 1979). Low values represent the presence of canted-ferromagnetic minerals such as α -Fe₂O₃ (haematite) and high values represent the presence of ferromagnetic minerals such as Fe₃O₄ (magnetite). Magnetic susceptibility was measured on splits from Cores I and II using a Bartington instrument equipped with a MS2E surface sensor. Measurements were performed in the Paleomagnetic Laboratory at the University of Bergen, Norway or at the Magnetic Laboratory at ETH, in Zürich, Switzerland.

Core chronology

Terrestrial plant macrofossils were carefully hand-picked from the core for AMS radiocarbon dating. Samples were washed with distilled water, dried and sealed in sterile glass containers before dating at the Poznań Radiocarbon Laboratory, Poland. Dates were calibrated using CALIB 5.1 (Stuiver et al. 2005) (Table S3). All reported dates are calibrated intercept dates that used the one-sigma range (weighted mean). Because the upper 60 cm in core I are missing (Fig. 2), the three dates from this core were shifted accordingly.

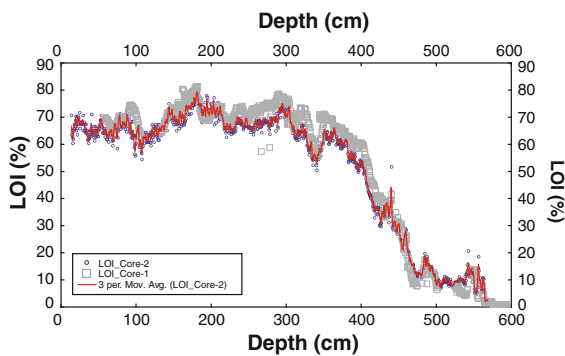


Fig. 2 Loss-on-ignition (LOI) stratigraphies from Cores I and II. The stratigraphies display very good co-variation in both LOI trends and amplitude. This comparison illustrates that Core I is missing 60 cm at the top

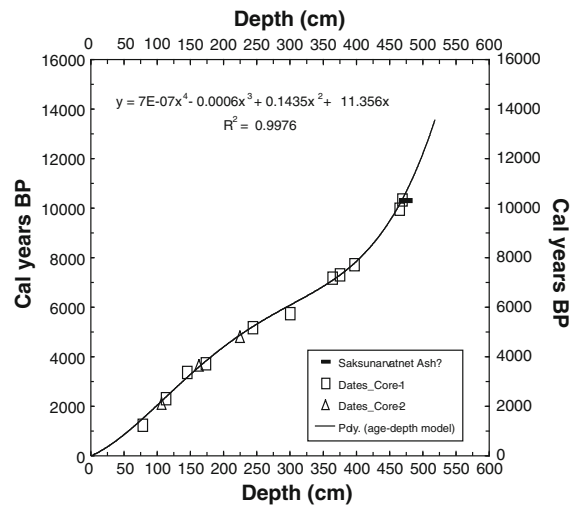


Fig. 3 Terrestrial macrofossil dates from Core I (squares) and Core II (triangles) were combined to form a composite age-depth model for Lake Natnålsvatn. A polynomial fit was used to establish the model. Based on downcore variations in LOI and the missing material at the top of Core I (Fig. 2), dates from Core I were adjusted by 60 cm with respect to their initial depth

A polynomial was used to construct an age-depth model for both cores. None of the 14 dates on macrofossils show an age-depth reversal (Fig. 3). The sedimentation rate has remained high and stable for most of the Holocene. The model age for a peak in magnetic susceptibility, which might represent the Saksunarvatnet Ash (Mangerud et al. 1986), is in good agreement with the known age of the ash, 10,347 years before AD 2000 (Rasmussen et al. 2007).

Results

The late glacial record

Visual inspection of the lower part of Core I revealed firm grey clay and silt with isolated clasts, with a sharp transition to overlying, finely laminated, greenish-grey silt. The next 50 cm consist of alternating silty laminae and silty gyttja laminae, characterised by abundant mm-size flakes of muscovite. From ~405 cm, there is a transition from greenish silt to dark-brown gyttja that continues from 370 cm to the top of the core (Fig. 4).

No macrofossils were located in the lowest part of the core. Ages before 10,370 cal year BP were

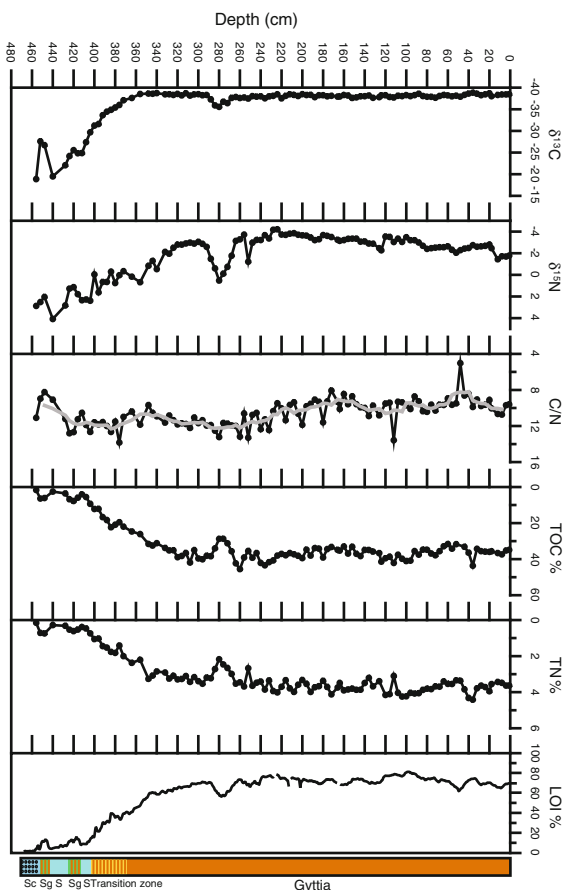


Fig. 4 Stable isotope values ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$), carbon/nitrogen (C/N) ratio, total organic carbon (TOC%), and total organic nitrogen (TN%) plotted against depth (cm). Sc Silty clay (this unit also contains clasts), Sg Silty gyttja, and S Silt

obtained by extrapolation and therefore have large associated uncertainties. Mean late glacial sampling resolution is ~ 30 years/cm and somewhat better for the Holocene. Relatively large clasts embedded in silty clays in the lowermost part of the core are thought to reflect deglaciation.

Throughout the late glacial part of the core, χ_{bulk} , ARM and SIRM values are generally higher than in the Holocene section (Figs. 5, 6). A stepwise decline culminates in the interval from 456 to 446 cm, which may correspond to the Allerød Interstade. The magnetic mineralogy, inferred from the S-ratio, suggests a gradual shift from “harder” to “softer” minerals, with a few exceptions indicated by lower values (Fig. 7). A prominent peak in ARM, SIRM, and particularly χ_{bulk} , obscures what would otherwise be a clear transition from the late glacial to the

Holocene (Fig. 6). This peak is anomalous compared to the remainder of the record, with χ_{bulk} values reaching $10^{-6} \times 1.97 \text{ m}^3 \text{ kg}^{-1}$.

Smear slides revealed a variety of grains derived from the catchment, notably quartz, feldspar and mica. In addition, the section between about 445 and 400 cm is characterised by abundant authigenic pyrite framboids (Fig. 8).

Organic matter is absent below 455 cm, so isotopic analysis of OM was only possible on samples from that depth and above (Fig. 4). The earliest part of the record shows a negative $\delta^{13}\text{C}$ excursion during the Allerød Interstade. Thereafter, following a return to higher $\delta^{13}\text{C}$ values, $\delta^{13}\text{C}$ decreases steadily upward through this section of the core, a trend that is also apparent in the early-late Holocene $\delta^{13}\text{C}_{\text{OM}}$ records from other marl lakes (Diefendorf et al. 2008). $\delta^{15}\text{N}$ also decreases upward, but in a more irregular fashion than the carbon isotopes (Fig. 5).

LOI₅₅₀ values tend to be correlated inversely with magnetic susceptibility during the late glacial (Fig. 6). A rapid increase in LOI₅₅₀, from 1–2 to 10–11%, occurs at the onset of the Allerød, $\sim 13,500$ cal year BP. There is an equally swift return to lower values (5%) at the onset of the Younger Dryas, $\sim 12,700$ cal year BP. At 11,500 cal year BP, values rise again, but a temporary return to lower values is associated with the previously described anomaly in magnetic properties. Based on the results presented above, three sedimentary phases (P4–P6), representing late glacial changes, were recognized in Core I (Fig. 7).

The Holocene record

From 10,300 to 6,800 cal year BP, LOI values gradually increase, from less than 10 to more than 70% (Figs. 5, 6). During the first 1,000 years of this period, the LOI increased stepwise, but during the latter 2,500 years of this period, the increase was smoother. At 6,800 cal year BP, LOI declined by $\sim 15\%$ (Fig. 6) for about 500 years. During the rest of the Holocene LOI values are extremely high, reaching a maximum of $\sim 81\%$ at 3,500 cal year BP, after which they declined for 1,300 years, to 60%.

Holocene variations in χ_{bulk} indicate that the influx of minerogenic matter after 9,500 cal year BP was low, although values averaged $10^{-7} \times 1.48 \text{ m}^3 \text{ kg}^{-1}$. There were marked strengthenings of magnetic susceptibility twice during the late Holocene (Fig. 6).

Fig. 5 Stable isotope values ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$) and magnetic susceptibility (χ_{bulk}) measurements through the Holocene. Note the similar decreasing trends between 10,000 and 7,500 cal year BP, and also the similar response in all three variables between 6,850 and 6,300 cal year BP. There are two-fold increases in χ_{bulk} at 4,000 and 2,300 cal year BP

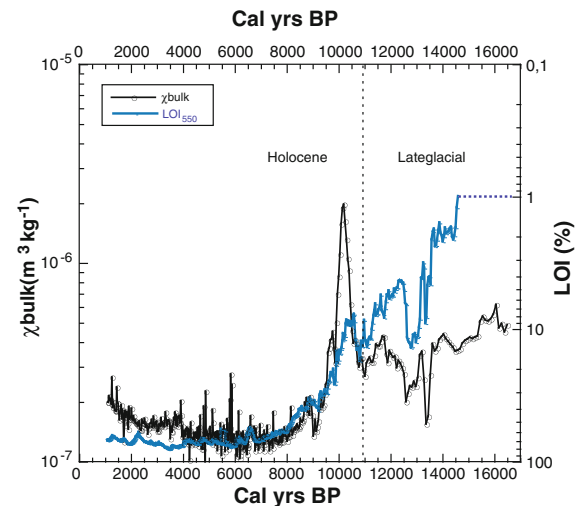
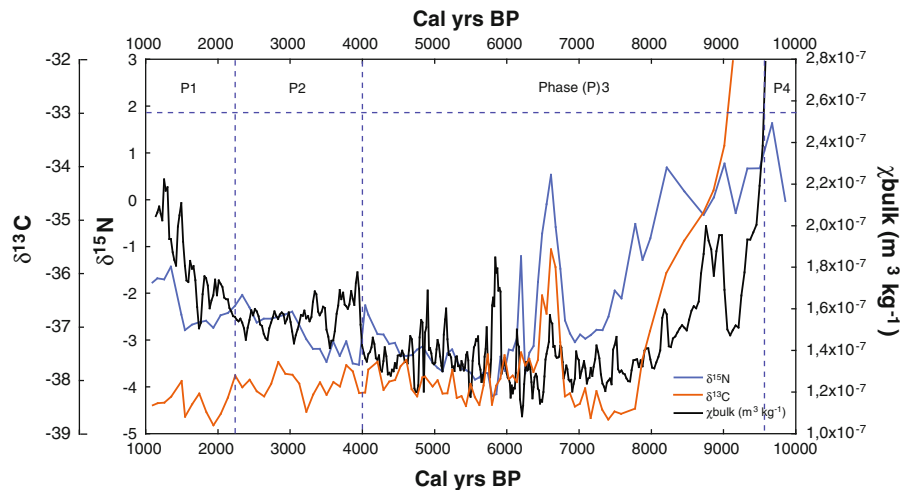


Fig. 6 Magnetic susceptibility (χ_{bulk} , left y-axis and log-scale) and LOI (% , right y-axis, values are reversed and on log-scale) variations through the late glacial and Holocene. Note the strong inverse relationship between LOI and χ_{bulk} during the entire late glacial period

The first increase began around 4,000 cal year BP, at which time higher values were attained rapidly. After 2,300 cal year BP, values began to increase further, a trend that continues towards the top of the core with χ_{bulk} values of $10^{-7} \times 2.15 \text{ m}^3 \text{ kg}^{-1}$.

Smear slides illustrate the organic-rich nature of the Holocene sediments and contain little siliclastic material (mainly quartz), suggesting minimal input of mineralogenic grains from the surrounding landscape. The organic fraction contains unstructured (amorphous) material of probable algal origin, as well as

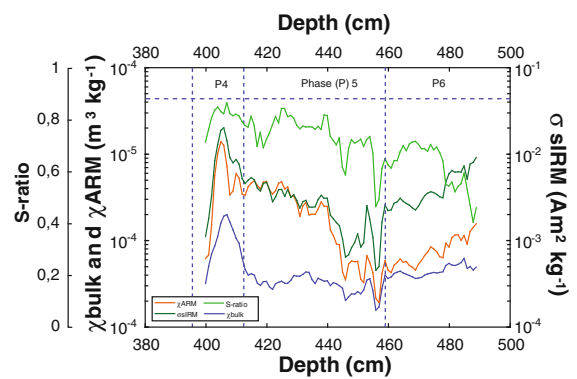


Fig. 7 Variations in χ_{bulk} and χ_{ARM} (left y-axis, log-scale) and SIRM (right y-axis, log scale) throughout the lower part of Core I. The peak ~ 405 cm depth might represent the Saksunarvatn Ash (10,347 before AD 2000). A change in magnetic mineralogy is indicated by the S-ratio (left, y-axis), where ‘softer’ minerals such as magnetite (Fe_3O_4) are associated with values close to 1

from higher plants, in particular well preserved remains of the moss *Drepanocladus* sp., and from a variety of aquatic invertebrates (Fig. 8). Diatoms are few, small and poorly preserved from the beginning of the Holocene until ca. 5,500 cal year BP, but thereafter become abundant, with a pronounced increase in large species. Early Holocene sediments contain euhedral, in some cases elongate, crystals of feldspar, showing typical albite twinning (Fig. 8). Their perfect crystal form strongly suggests they are authigenic feldspar. The crystals become rare and eventually disappear around 5,000 cal year BP, at the same time diatoms become abundant.

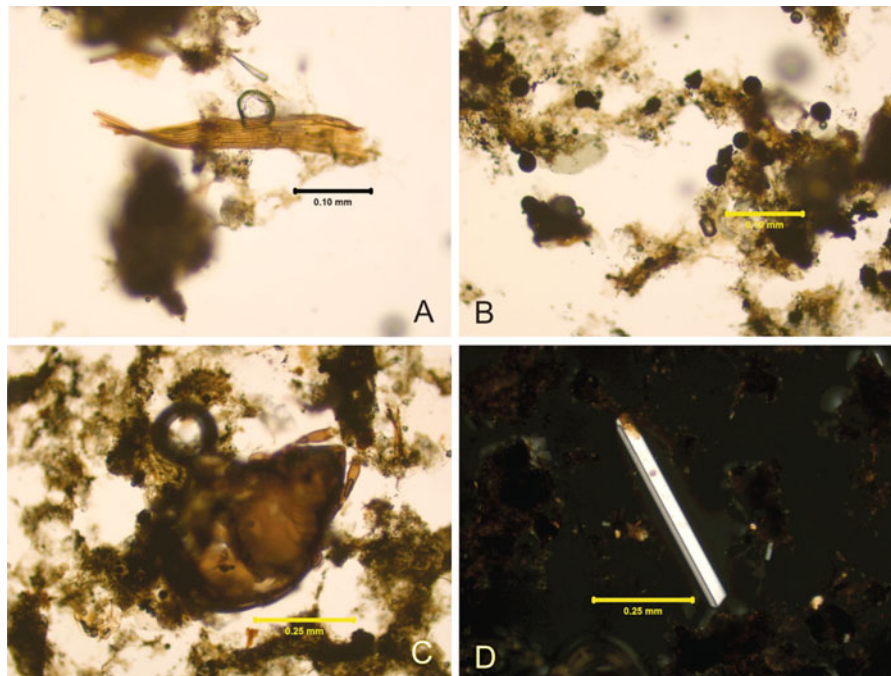


Fig. 8 Photomicrographs of smear slides showing some notable features of the Lake Nattmålsvatn sediments. **a** Fragment of the moss *Drepanocladus* sp., a characteristic component of the preserved organic matter. **b** Pyrite framboids (*black*

spherules) produced by microbial sulphate reduction. **c** Unidentified aquatic invertebrate. **d** Elongate, euhedral authigenic feldspar showing typical albite twinning. Cross-polarised light

The $\delta^{13}\text{C}$ record from the earliest Holocene continues the trend seen in the late glacial. From the Younger Dryas, there is a transition to lower $\delta^{13}\text{C}$ values in the Holocene. The $\delta^{13}\text{C}_{\text{org}}$ curve stabilizes $\sim 8,000$ cal year BP and values remain low ($< -38.3\text{‰}$) for the rest of the Holocene, except for a short period of isotope enrichment at 6,700–6,400 years BP, the Atlantic period, corresponding to the lower LOI values described above (Fig. 5).

Nitrogen isotope values ($\delta^{15}\text{N}_{\text{org}}$) also display a trend to progressively lower values into the Holocene, but not as smooth a trend as for $\delta^{13}\text{C}_{\text{org}}$ (Fig. 5). The Younger Dryas and perhaps the Preboreal Oscillation (PBO) have the highest values, around +2.2 to +4‰, probably because immature soils provided the dominant source of nitrogen during this period (Wolfe et al. 1999; Talbot 2001). $\delta^{15}\text{N}$ values stabilise at relatively low values, -4 to -2‰ , at around 7,500 cal year BP, and remain low for the rest of the Holocene, except for a period of isotope enrichment in the Atlantic period, as seen for carbon isotope ratios (Fig. 5). There are, however, slight

temporal differences between the trends in carbon and nitrogen isotope signals during this period (Fig. 5). After about 3,200 cal year BP the nitrogen isotope ratios increase slightly, but display relatively low variability.

Discussion

Sources of organic matter in sediments

In Lake Nattmålsvatn sediments, C/N ratios range from 8 to 13 (Fig. 4), which indicates that organic matter comes predominantly from aquatic sources and that in-lake production is high. Organic matter delivered to the lake from the catchment is moderate by comparison. The shift from a C/N ratio of ~ 12 at $\sim 6,000$ to ~ 9 at $\sim 4,000$ cal year BP indicates a decline in delivery of organic matter from the terrestrial environment, increased lake bioproduction, or both (Figs. 4, 5). Prior to 8,000 cal year BP, TOC and TN are highly correlated ($r = 0.99$, $P < 0.001$). A linear fit of the data indicates a mean C/N of ~ 11

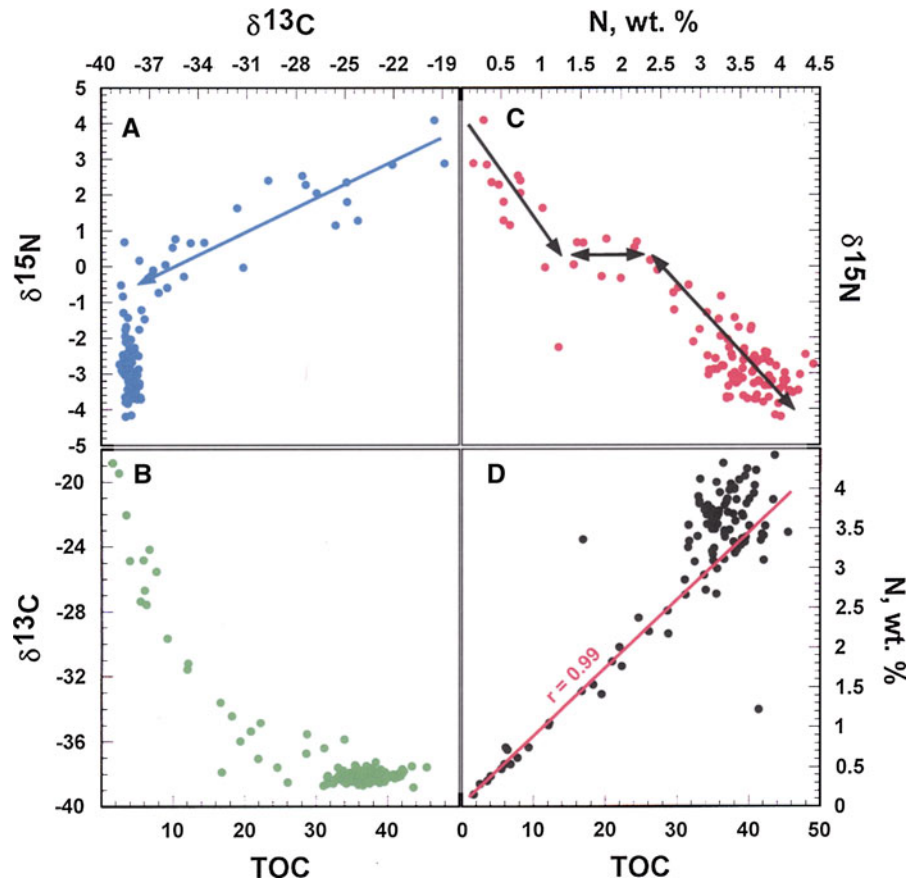


Fig. 9 Cross-plots of the most significant isotope and elemental variables. **a** $\delta^{13}\text{C}$ versus $\delta^{15}\text{N}$ showing a positive correlation for the late glacial to earliest Holocene section. This correlation breaks down once $\delta^{13}\text{C}$ values stabilise at -38.1% . The arrow shows the trajectory followed by the isotopic data during the late glacial–Holocene transition. **b** $\delta^{13}\text{C}$ versus TOC. An inverse correlation exists in the early part of the record, but the relationship breaks down, as in plot A, when $\delta^{13}\text{C}$ values drop below -38% . **c** $\delta^{15}\text{N}$ versus TN%. Similar to the $\delta^{13}\text{C}$ versus TOC plot (**b**), there is an overall inverse correlation, however in the case of nitrogen, the relationship is

(Fig. 9). After *ca.* 8,000 cal year BP, and in recent times, this relationship breaks down and the data are poorly correlated ($r = 0.22$) and show slight N enrichment (mean C/N = 10.2) in relation to the older deposits (Fig. 9). This change may reflect conversion of Lake Nattmålsvatn from an oligotrophic to a eutrophic waterbody, a change that was probably driven by the onset of warmer climate.

Organic matter is frequently a minor constituent of lake sediments. In Lake Nattmålsvatn, organic matter dominates the sediment mass, with maximum LOI values near 82% (Fig. 2). We assume that the residual

stepped, with a transition to higher TN at *ca.* 8,000–7,000 cal year BP. This was due to a change from oligotrophy to eutrophy. The single arrow shows the direction of change during the late glacial, the double arrows show the range of variation during the transition period and the Holocene. **d** TOC versus TN. A strong positive correlation is seen for the late glacial to earliest Holocene part of the record and mean C/N was 11. Once the lake became eutrophic the TOC/TN relationship broke down and the data formed a poorly correlated cluster with a mean C/N of 10.2, i.e. slightly N-enriched relative to earlier deposits

material from the loss-on-ignition procedure consists largely of diatoms, given the low MS signal and the smear slide observations.

Stable isotopes

Both $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ display large ranges, ~ 20 and $\sim 9\%$, respectively, suggesting major changes in carbon and nitrogen cycling throughout the post-glacial history of Lake Nattmålsvatn. In the late glacial, there were $\delta^{13}\text{C}$ values greater than -25% . Sediments of that phase consist of silt with low TOC

content, alternating with silty gyttja. This reflects a pioneer phase with little catchment vegetation. Dissolution of detrital carbonate and exposed marble bedrock would have been the principal source of DIC at that time, contributing isotopically enriched runoff to the lake, which in turn would generate OM with relatively high $\delta^{13}\text{C}$ values. A similar scenario is apparent in Lough Inchiquin (Eire), where the limestone bedrock dominated the isotopic composition of lake OM during the immediate post-glacial period (Diefendorf et al. 2008).

Evidence for weathering of labile glacial detritus is also provided by abundant pyrite framboids in parts of this section, which were formed by microbial sulphate reduction. The dissolved sulphate required for pyrite formation probably came from the oxidation of detrital pyrite in tills derived from mica schist. Pyrite is absent from younger deposits, despite the anoxic conditions that must have prevailed in such organic-rich sediments. Once all detrital pyrite had been removed from the glacial deposits, there presumably was no other source of sulphate, and bacterial sulphate reduction was no longer an important process in Holocene lake sediments.

The isotope and elemental data display some interesting relationships (Fig. 9). During the initial postglacial period, there is a positive correlation between $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ (Fig. 9), but this breaks down around 8,000–7,000 cal year BP when $\delta^{13}\text{C}$ stabilises at $-38.1 \pm 0.3\text{‰}$, while $\delta^{15}\text{N}$ varies over a range $\sim 5\text{‰}$ (Fig. 9). At the same time there is a shift in the $\delta^{15}\text{N}$ vs. TN relationship (Fig. 9), as the OM displays a much higher TN content. This change supports our inference that the transition from oligotrophic conditions that prevailed during the late glacial, to eutrophic conditions, occurred at ca. 8,000 cal year BP (Fig. 5).

The $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ in the early part of the record show an inverse correlation with TN and TOC, respectively. This is opposite to what is expected if variations in isotopic composition are driven by primary productivity (Hollander and McKenzie 1991; Bade et al. 2004). The inverse relationships suggest, instead, that the DIC and DIN pools available for primary production were not limited. Development of ice cover for 5–7 months each year may have contributed to this DIC and DIN availability. Respiration of sediment OM during winter months can lead to accumulation of dissolved and free gases beneath the ice, notably carbon dioxide, methane, ammonia

and nitrous oxide (Wetzel 2001), with concentrations well in excess of those resulting from equilibrium with the atmosphere (Kortelainen et al. 2000; Striegl et al. 2001; Huttunen et al. 2003a, b). Primary production may also continue beneath the ice (Wetzel 2001). Although it is likely that production in Lake Nattmålsvatn was low during the dark winter months, high insolation at these northern latitudes during April and May could have stimulated high rates of primary production beneath the ice at a time when dissolved nutrients would have been at their highest concentrations. Very low Holocene $\delta^{13}\text{C}$ values may reflect assimilation of this DIC pool under the ice and during the ensuing period of ice-out, when primary production reaches a peak (Boström and Pettersson 1977). Carbon isotope values lower than about -32‰ suggest that methane oxidation has contributed CO_2 to the DIC pool (Rau 1978; Whiticar and Faber 1986). Another reason for the low values may be elevated pCO_2 , as fractionation increases with increased CO_2 concentration (Degens et al. 1968). The remarkably constant $\delta^{13}\text{C}$ record for the Holocene suggests that Lake Nattmålsvatn was, and still is, a well-buffered system. We do not, however, have a good explanation for why the $\delta^{13}\text{C}$ value for Holocene OM stabilized at $-38.1 \pm 0.3\text{‰}$.

Abundance of dissolved nitrogen species may also explain the low $\delta^{15}\text{N}$ values ($<0\text{‰}$). Like C isotopes in organic matter, N-isotope fractionation increases with rising N availability (Talbot 2001). Unlike the $\delta^{13}\text{C}$ record, however, $\delta^{15}\text{N}$ did not stabilise at a fixed value, but varied over a range of $\sim 2\text{‰}$, probably due to variations in N availability, likely in the form of ammonia (Billaud 1968), and drawdown of the DIN reservoir as the spring bloom proceeded.

Low $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values in Holocene OM were interrupted only by higher values from 6,850 to 6,500 cal year BP (Fig. 5). Nitrogen isotopes, in particular, suggest a return at that time to near-late-glacial conditions. Concomitant reductions in TOC, TN and LOI, accompanied by the rise in $\delta^{15}\text{N}$ and a small rise in C/N (Figs. 4, 5), are consistent with a reduction in lake productivity and an increased supply of plant and soil OM to the lake. As mentioned, we interpret this excursion to be the result of a previously unrecognised brief period of cooling in Arctic Norway.

Feldspar authigenesis also provides insight into the state of the lake. The paucity of diatoms in the

feldspar-bearing section suggests that the silica necessary for feldspar precipitation may have been provided by diatom dissolution. The solubility of opaline silica increases dramatically at $\text{pH} \geq 8$. Diatom dissolution probably occurred when the lake was at least seasonally alkaline, with a pH greater than the current 8. Such conditions could have been achieved in a climate warmer and drier than today's, when high rates of primary production and greater summer evaporation led to elevated surface water pH. Increased diatom abundance and the accompanying disappearance of feldspar after *ca.* 5,500 cal year BP suggest a change to less alkaline conditions, probably due to a transition to a cooler, wetter climate that characterised the later Holocene in the Senja region.

Correlation of TOC and LOI

Overall, there is a strong correlation between LOI and TOC (Fig. 10), but Fig. 10 also reveals a result not previously described in the literature. Above $\sim 60\%$ LOI, corresponding to $\sim 32\%$ TOC, there is a wide scatter of values, and correlation between the two variables in this group of high values is poor (Figs. 9, 10). Lacustrine sediments with such large amounts of

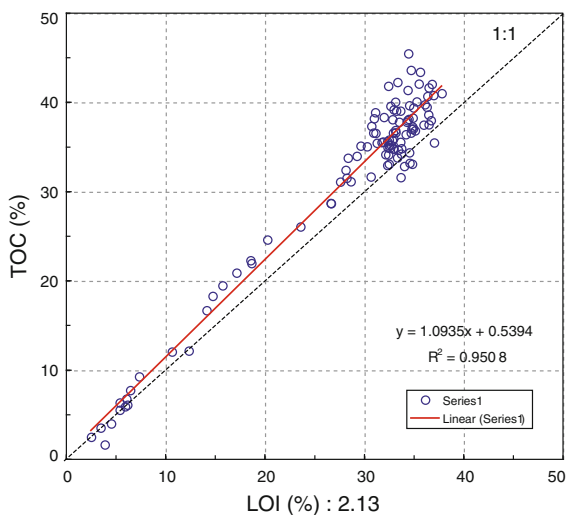


Fig. 10 Total organic carbon (TOC %) plotted against Loss-on-ignition (LOI %). Original LOI values were divided by 2.13, according to Dean (1974) to approximate the amount of TOC in bulk organic matter. TOC values from Lake Nattmålsvatn are consistently higher than the 1:1 relationship, indicating that Dean's (1974) empirical constant (2.13) is too high for Lake Nattmålsvatn. The relationship generally breaks down when TOC values are $>30\%$

OM are rare, and the LOI and TOC values exceed those that Dean (1974) used to develop the LOI-TOC relationship. There are several possible explanations for the breakdown of the correlation at high concentrations: (1) LOI was measured with error at such high concentrations of OM, perhaps due to incomplete ignition at 550°C , (2) high concentrations of TOC overwhelm the elemental carbon analyser, resulting in artificially low carbon counts, (3) The linear relationship between LOI and TOC does in fact break down when OM content exceeds $\sim 60\%$, (4) The concentrations above which the relationship breaks down, 60% LOI and 32% TOC, occur at a depth (age) in the core when there is a shift from low values in the deglacial and early Holocene to higher values in the middle and late Holocene, perhaps suggesting a change in OM processing starting $\sim 7,000$ cal year BP (312 cm).

We speculate that 3 and 4 above, i.e. extremely high LOI values ($>60\%$) in combination with a change in OM composition, are responsible for the poor relationship between LOI and TOC. Below 60% LOI there is a very strong correlation between the two variables ($r = 0.99$, $P < 0.001$) (Fig. 10). As noted above, there is a gradual decrease in C/N ratio after the middle Holocene, indicating an increasing proportion of nitrogen-rich OM in sediments, originating from lake primary production. This is also evident in the %TN curve, which shows the highest nitrogen concentrations in the latest part of the Holocene (Fig. 5). Alternatives (1) and (2) were rejected for several reasons. Sample size was small (1 ml wet material), so incomplete ignition is unlikely to have been a problem. Furthermore, the strong stratigraphic correlation between LOI values in Cores I and II also suggests complete ignition was not a problem. Furthermore, measurement errors with the elemental analyzer are highly unlikely. The instrument is routinely tested using carbon-rich materials such as wheat flour, with TOC values $>40\%$, and should therefore be capable of handling organic-rich lacustrine sediments. In addition, TOC values are not consistently low or high relative to what might be expected from LOI. Instead, there is simply wide scatter.

TOC and LOI values from the Lake Nattmålsvatn cores display corresponding trends over the long time frame (Fig. S1), but the signals diverge in some sections, particularly when OM exceeds 60% . The

TN and C/N records enable us to conclude that poor correlation occurs in lake sediments consisting of more nitrogen-rich OM, in the middle and late Holocene.

Calcite dissolution

Calcite was observed in shallow waters and along the lake shore during fieldwork in July 2002. Greyish-white coatings on submersed rocks and twigs and marl on the lake bottom confirmed that Nattmålsvatn is a *Chara* (marl) lake. Langangen (1995) supports this observation, but nonetheless, there is no evidence of carbonate in the core. Several samples were checked for carbonate by XRD analysis and all yielded negative results.

Calcite dissolution in hard-water lakes has been described from other sites, e.g. the Swiss Alps (Ohlendorf and Sturm 2001). Biogenic calcite precipitation in Lake Hagelsewli occurs in late summer as a result of photosynthetic uptake of CO₂. The following winter, the precipitated calcite is completely dissolved. The long period of ice cover leads to bottom waters strongly undersaturated with respect to calcite. During ice cover, bottom waters become anoxic and pH declines.

In the northern part of Norway examined in this study, ice cover on lakes of small to medium size and depth averages 5–7 months (Økland and Økland 1998), but can be longer. Such conditions are similar to those in the Swiss Alps, and winter processes are probably similar in the lakes of both regions. Neither Hagelsewli nor Nattmålsvatn is a “typical” oligotrophic, alkaline water body. For instance, there is considerable decomposition of OM in both lakes. In this study, such decomposition was evident from the strong H₂S odor, as well as methanogenesis in the sediments. The organic matter is subject to microbial decomposition, with consequent consumption of O₂. Even in an oligotrophic hard-water lake such as Nattmålsvatn, oxygen depletion at the sediment–water interface may occur, but does not necessarily lead to oxygen depletion of the entire hypolimnion. Similar scenarios are described by Økland and Økland (1998).

In Lake Nattmålsvatn, surface sediments are dominated by particles from high autochthonous organic production during spring and summer, along with precipitated calcite. There are only minor

amounts of detrital material. Decomposition of the OM produces slightly acidic conditions at the sediment–water interface with a sufficiently low pH to dissolve the precipitated calcite. A higher influx of detrital material could possibly ‘protect’ the calcite from dissolution.

Minerogenic influx and environmental implications

The lowermost part of the core is characterized by greyish silt containing larger clasts, most likely related to the local retreat of the inland ice sheet and/or a local ice cap at Senja (Fig. 1). The sequence reveals high values for ARM, SIRM and χ_{bulk} that appear to drop relatively rapidly (Fig. 7). Such high values are only surpassed by the brief period centred at 10,250 cal year BP, which might represent the Saksunarvatnet Ash layer (Mangerud et al. 1986, Rasmussen et al. 2007). Although exact dating of the deglaciation may not be possible, its timing can be constrained. The lake was positioned between two ice-sheet margins, the youngest and best defined being the Younger Dryas, referred to locally as the Tromsø-Lyngen stage (Andersen 1968). The older one is commonly referred to as the D-event (Vorren and Plassen 2002) (Fig. 1). According to Vorren and Plassen (2002), this event occurred between 13.8 and 13.2 ¹⁴C years BP. Recalibrating the raw ¹⁴C dates ± 150 year with CALIB 5.0.2, the event dates to between $15,890 \pm 270$ and $15,120 \pm 250$ cal year BP. Consequently, deglaciation of Lake Nattmålsvatn took place after $15,120 \pm 250$ and before 12,900 cal year BP. Our age-depth model suggests that deglaciation occurred earlier, around $15,400 \pm 250$ cal year BP, but this age is tentative. A more likely scenario for deglaciation is the Bølling warming, which started around 14,700 cal year BP, when cirque glaciers in the region started receding (Paasche et al. 2007a).

Holocene minerogenic influx to the lake can be broadly divided into four phases (Fig. 5). The early Holocene transition (P4) included what might be the Saksunarvatnet Ash (10,347 before 2000 AD) (Fig. 7). This interval was followed by a long and stable phase from 9,500 to 4,000 cal year BP, during which there were eight brief (<50 years) episodes with greater minerogenic material. In addition to these short periods, there were two prominent events that lasted

much longer. The first one spanned the period between 6,850 and 6,500 cal year BP, and the second lasted from about 6,000–5,750 cal year BP (Fig. 5). These episodes of higher minerogenic matter influx probably reflect an increase in runoff from a sustained increase in winter precipitation (Bakke et al. 2010).

At 4,000 cal year BP, a shift occurred in <100 years, marking the onset of inferred increased runoff that prevailed for 1,700 years. At 2,300 cal year BP, minerogenic matter increased again and the positive trend continued to the end of the record at ca. 1,100 cal year BP. We also interpret these two shifts to indicate wetter winter conditions which might have been related to large-scale atmospheric circulation anomalies. This interpretation is in good accordance with results from northern Fennoscandia (Vorren et al. 2007; Bakke et al. 2005; Paasche et al. 2004, 2007a, b).

Timing of events in the sediment record

In the late glacial period, variations in isotope values ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$), MS, LOI and C/N in the Lake Nattmålsvatn cores occurred synchronously (Figs. 4, 6). Vegetation was poorly developed at the time and there was a large influx of detrital particles from the lake catchment. There was a strong correlation between $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$, and an inverse correlation between the isotope signatures and their respective elemental concentrations (TOC and TN), from the late glacial up to the levels where methane oxidation had a profound influence on the isotope results (Figs. 4, 9). These results are explained by the profound effect of prolonged ice cover on carbon and nitrogen cycling in Lake Nattmålsvatn. The termination of the Younger Dryas and the beginning of the Holocene are not reflected synchronously in variables LOI, TOC and C/N. This is because the relative increase in OM did not result from increased soil erosion and input of terrestrial plant material, which implies that the gradual depletion of both $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values was not a result of changes in OM source, but rather, changes in DIC and DIN. The decrease in χ_{bulk} in the early Holocene contrasts with the gradual increase in LOI, indicating less transport of detrital material to the lake (Fig. 6). The increasing LOI values signal a smaller relative amount of detrital material in the sediment. The MS values record a diminishing magnetic signal in sediment that already displayed a weak signal.

In the Holocene, C/N ratios were stable. TOC and TN values increased, whereas $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values declined. The $\delta^{13}\text{C}$ reached “steady” values earlier than other variables, indicating that methanogenesis and oxidation of methane began early in the Holocene. Variables other than $\delta^{13}\text{C}$, however, indicate that variations did occur in the Holocene.

The 6,850–6,500 cal year BP event, evident in the isotope and the LOI₅₅₀ stratigraphies, is the most striking excursion in the Holocene (Fig. 5). The C/N signal, however, showed little change. This may indicate sustained lake production in combination with reduced influx of terrestrial OM. Although the event is recorded in both the $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ datasets, it is more pronounced in the latter. This probably does not reflect differences in response time between these variables, but may instead indicate that the DIC changed little in the system and that a threshold must be surpassed before a shift in $\delta^{13}\text{C}$ occurs. If winter ice cover helped maintain low $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values, then ice out probably lead to ventilation of depleted DIN and DIC, perhaps changing the $\delta^{13}\text{C}$ values.

In recent years, winters in northern Norway have been mild, and persistent low temperatures have not set in until January and February, resulting in a very short period of ice cover on lakes. Comparable conditions may have existed during the 6,850–6,500 cal year BP event, with mild, wet winters and poorly developed ice cover. Plant growth and algal blooms in summer may have remained unchanged, despite milder winters. Under such a scenario, the composition of OM would not change. For example, the C/N ratio would not be affected. Altered winter conditions, however, would change DIN and DIC, and also cause a stronger MS signal, given that greater winter precipitation leads to higher runoff from the catchment, bringing more minerogenic matter into the lake.

From 6,300 to 4,000 cal year BP there was a decline in C/N ratio. This decline was probably associated with a relative decrease in the amount of terrestrial OM accumulating in the lake sediments or a gradual increase in aquatic primary production.

Different climate proxies in lake sediments have different thresholds and sensitivities to climate changes that affect a lake and its catchment. Some proxy variables may be highly sensitive, whereas others only respond to major climate and

environmental changes. Additionally, there may be temporal differences in response among the climate proxies in a paleoclimatic study. The core chronologies and inter-site correlation within a lake may further compromise interpretations. In the Lake Nattmålsvatn records, only the deglaciation events produced synchronous changes among most of the proxies. The deglacial was a period with increased detrital influx and moderate to low amounts of OM in the sediments. Holocene deposits are extremely organic-rich, which seems to have lowered the sensitivity of the climate proxies used in this study, yielding relatively constant values. The exception to this generalization was the 6,850–6,500 cal year BP event, which is recorded by all proxies except C/N ratio. As suggested, most of the proxy variables were probably affected by shifts in winter conditions, whereas the C/N ratio stayed rather constant because it reflects organic matter production during summer.

Conclusions

We presented a high-resolution sediment record from Lake Nattmålsvatn, northern Norway, that spans the last <15,000 cal year. The Holocene sequence, in particular, is organic-rich, with TOC values up to 45%. Our main conclusions are:

1. Stable isotopes of organic matter ($\delta^{13}\text{C}$ and $\delta^{15}\text{N}$) in sediments do not serve as strong paleoclimate proxies in lakes such as Lake Nattmålsvatn. The large amount of organic matter production and deposition, in combination with methanogenesis and prolonged seasonal ice cover, makes the lake less sensitive to changing environmental conditions. Magnetic variables are better suited to track climate and environmental variations, predominantly fluctuations in winter precipitation that affect catchment runoff.
2. A period of microbially-mediated pyrite formation during the late glacial probably reflects weathering of newly exposed, pyrite-bearing glacial detritus. Once removal of the detrital pyrite was complete, sulphate concentrations in the lake were insufficient to permit formation of early diagenetic pyrite.
3. Isotopic and elemental data suggest that Lake Nattmålsvatn changed from an oligotrophic to a eutrophic waterbody about 8,000 cal year BP.
4. Authigenic feldspar formation during the early Holocene was the result of alkaline conditions in the lake. This was probably due to a warmer and drier climate, which promoted diatom dissolution, thus providing the silica required for feldspar precipitation.
5. Climatic deterioration from 6,850 to 6,500 cal year BP has not been described previously in Scandinavia. This anomaly most likely represents a period of increased winter precipitation and reduced ice cover. A two-step increase in winter precipitation, at 4,000 and 2,300 cal year BP, is indicated by magnetic susceptibility, and corresponds to the ‘Holocene Turnover’ evident in other paleoclimate studies.
6. Loss-on-ignition (LOI) and elemental analysis revealed a previously undescribed relationship between TOC and LOI₅₅₀ in organic-rich sediments. At LOI₅₅₀ values >60%, there is no longer a linear relationship between LOI₅₅₀ and TOC. This may reflect variations in the C/N ratio of the organic matter. This decoupling of LOI₅₅₀ and TOC makes it difficult to interpret changes in both proxies or link their variations to specific climatic conditions. Overall, however, long-term major trends in LOI₅₅₀ and TOC were in agreement.

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