

River flooding as  
a driver of polygon  
dynamics

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# River flooding as a driver of polygon dynamics: modern vegetation data and a millennial peat record from the Anabar River lowlands (Arctic Siberia)

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## Abstract

The spatial and temporal variability of a low-centred polygon on the eastern floodplain area of the lower Anabar River (72.070° N, 113.921° E, northern Yakutia, Siberia) has been investigated using a multi-method approach. The present-day vegetation in each square metre was analysed revealing a community of *Larix* shrubby *Betula* and *Salix* on the polygon rim, a dominance of *Carex* and *Andromeda polifolia* in the rim-to-pond transition zone, and a predominantly monospecific *Scorpidium scorpioides* coverage within the pond. The TOC content, TOC/TN ratio, grain-size, vascular plant macrofossils, moss remains, diatoms, and pollen were analysed for two vertical sections and a sediment core from a transect across the polygon. Radiocarbon dating indicates that the formation of the polygon started at least 1500 yr ago; the general positions of the pond and rim have not changed since that time. Two types of pond vegetation were identified, indicating two contrasting development stages of the polygon. The first was a well-established moss association dominated by submerged or floating *Scorpidium scorpioides* and/or *Drepanocladus* spp. and overgrown by epiphytic diatoms such as *Tabellaria flocculosa* and *Eunotia* taxa. This stage coincides temporally with a period in which the polygon was only drained by lateral subsurface water flow, as indicated by mixed grain sizes. A different moss association occurred during times of repeated river flooding (indicated by homogeneous medium-grained sand that probably accumulated during the annual spring snow melt), characterized by an abundance of *Meesia triquetra* and a dominance of benthic diatoms (e.g. *Navicula vulpina*), indicative of a relatively high pH and a high tolerance of disturbance. A comparison of the local polygon vegetation (inferred from moss and macrofossil spectra) with the regional vegetation (inferred from pollen spectra) indicated that the moss association with *Scorpidium scorpioides* became established during relatively favourable climatic conditions while the association dominated by *Meesia triquetra* occurred during periods of harsh climatic conditions. Our study revealed a strong riverine influence (in addition to climatic influences) on polygon development and the type of peat accumulated.

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# 1 Introduction

Polygon mires are the most common type of arctic mire. Often these mires occur in an advanced stage of succession, known as low-centred polygons (Minke et al., 2007). Low-centred polygons evolve when frost-heave processes form ridges (the polygon rims) above ice wedges and depressions between the ridges that are often water-filled (the polygon pond). **Since these polygons have an important role in the arctic carbon cycle (Hobbie et al., 2002); although the mires of the arctic tundra only cover about 3% of the arctic land area, they contain more than 15% of the world's soil carbon (Post et al., 1982).** Improving our understanding of polygon dynamics is therefore a key scientific objective. Due to the harsh living conditions plants with short life cycles are common as well as those capable of asexual reproduction. Woody shrubs and herbs are consequently replaced at higher latitudes by cryptogams, as a result of their lower requirements for light, temperature and nutrients (Frahm, 2001). Most of the biomass in the arctic tundra is therefore fixed in peat from decomposed bryophytes (Longton, 1997).

Climate and the nature of the soil substrate are the main factors influencing the formation of polygons while their dimension, form and position within landscapes are controlled by complex positive feedbacks between vegetation, ice, water, and peat (Minke et al., 2007). Investigations have previously been conducted into the ecology of polygonal landscapes and the way in which they function. For example, the hydrology of the active layer has been investigated in relation to the form of the polygon (Riordan et al., 2006; Yoshikawa and Hinzman, 2003) and the displacement of soil by frostheave processes (Mackay, 2000). The dependence of the energy and water balances in the polygonal tundra on precipitation and weather conditions have already been investigated (Boike et al., 2008). **Especially in the Lena River Delta (Samoylov Island) which is also located in an area of river influence form and sediment characteristics have been investigated (Boike et al., 2012; Fiedler et al., 2004).** In the last decade studies with a special focus on the carbon content (Zubrzycki et al., 2013) and the net

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ecosystem CO<sub>2</sub> exchange have been **done**, e.g. in low-centred polygons on Samoylov Island (Runkle et al., 2013). Other investigations have used polygon sediments to trace short-term changes in vegetation (De Klerk et al., 2011) and long-term changes in landscape dynamics (Minke et al., 2009; Zoltai and Vitt, 1990, 1995). While previous studies have identified the general processes that lead to the formation of polygon mires, the polygon dynamics and associated driving factors remain only poorly understood.

In the North Siberian Lowland, polygon mires are particularly common along rivers or in the vicinity of lakes. We therefore selected an individual low-centred polygon (designated 07-SA-LY) on the flood plain of the lower Anabar River as study site using a multi-method approach that included analysis of vegetation, fossil moss tissue, pollen, diatoms, grain-size, and organic matter, together with radiocarbon dating. The objectives of this research were (1) to investigate the spatial pattern of vegetation in relation to abiotic factors (2) to reconstruct past changes in biotic communities and abiotic environments, and (3) to identify the drivers behind the polygon dynamics.

## 2 Geographic setting

The study area in north-western Yakutia (Siberia) is characterized by an extreme continental climate. Winters are dominated by a stable high pressure system over the Siberian landmass resulting in low temperatures (mean January temperature: -36.1 °C) and only a thin snow cover. Summers are dominated by Icelandic and Aleutian lows; they are short and relatively cool, with mean July temperatures of 12 °C (Saskylakh Climate Station, 71.967° N, 114.083° E: Rivas-Martinez and Rivas-Saenz, 2007).

The study area is underlain by 500 to 600 m of permafrost (Yershov et al., 1991). Temperature increases at the beginning of the Holocene initiated thermokarst processes that led to permafrost degradation and widespread subsidence (Czudek and Demek, 1970). The study area is characterized by meander terraces, thermokarst lakes

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5 alas depressions, and polygon mires. The uppermost parts of the soils are wet and organic-rich and are classified as Histic Fluvisols (Jones et al., 2010).

10 Polygon fields in the Anabar River lowlands were observed from helicopter flights to be restricted to areas along the flanks of rivers. The studied polygon is located within a polygon field on the Anabar River floodplain about 800 m to the east of the river bank and within the curve of an abandoned meander. It lies about 15 km north of Saskylakh on the North Siberian Lowland in the north-eastern part of Yakutia (Fig. 1b). The Anabar River is fed by the Bol'shaya Kuonamka and the Malaya Kuonamka rivers, which originate on the Anabar Shield. It drains into the western part of the Laptev Sea ~ 400 km west of the Lena River Delta and has a discharge that is highly variable, both seasonally and annually. In contrast to rivers in southern Yakutia, the Anabar River freezes to the bottom during winter followed by a rapid increase in discharge in spring that reaches a maximum of  $5000 \text{ m}^3 \text{ s}^{-1}$  (data from Saskylakh station; Huh and Edmond, 1999).

15 The lower part the Anabar River in particular is a meandering and partly braided river system (Fig. 1b). The polygon field in which our site is located is drained by several small streams flowing into the Anabar River in summer but can be flooded in spring. Thus the rapid melting of snow in spring as result of the sharp increase in temperature combined with the vicinity of the site to the large Anabar River results in a water supply to the polygon field that is both seasonally and inter-annually variable.

20 Our field vegetation survey has indicated that the study site lies within an area with *Larix* trees up to 4 m high. The area belongs to the geobotanical sub-province "Anabar-Lena subarctic" (Isaev et al., 2010) with a "widely spaced dwarf shrub tundra (*Betula exilis*, *Salix pulchra*), dwarf shrub tundra (*Cetraria cucullata* together with *Alectoria ochroleuca*, *Coelocaulon divergens*) and tundra bog" vegetation type according to the agriculture map from Yakutia (Matveev, 1989).

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### 3 Materials and methods

#### 3.1 Fieldwork

Fieldwork was completed in August 2007. A typical low-centred polygon (07-SA-LY; 72.070° N and 113.921° E; Fig. 1c, d) was selected for detailed study. The polygon was divided into 156 square meter plots, using a grid labelled A to L and 1 to 13 m (Fig. 2). We measured the elevation of the land surface (including the floor of the pond) relative to the water level in the pond as well as the depth of the active layer, at the centre of each plot (method according to Donner, 2007). The coverage of the main vascular plants was recorded for each plot in modified per cent classes according to the Braun-Blanquet floristic approach. Several taxa could only be recorded to genus or family level because the flowering period of most taxa was already over.

We also collected pH and conductivity measurements in the field for both soil and surface water along Transect G (Fig. 2a) using a WTW Multi 350i measuring instrument. We obtained a short core (Core C) of the upper 54 cm of sediment from plot G8 (pond) using a simple plastic tube (Fig. 2a). The core became compressed to 27 cm in the plastic tube during the coring process due to the loose organic material. Sub-samples were taken from Core C at 1 cm intervals. Since we cannot assume a homogenous compression over the whole length of the core, we present the data for the 27 cm core length and not for the original 54 cm length. We also sampled vertical soil sections of 42 cm length (Section A) and 50 cm length (Section B) in plots G1 and G3, respectively, at 2.3 cm resolution. The materials were transported to the Alfred Wegener Institute in Potsdam, Germany and stored at 4 °C.

#### 3.2 Radiocarbon dating

Handpicked, alkali-residue plant remains from twelve sediment samples were dated by radiocarbon accelerator mass spectrometry (AMS) at the Leibniz Laboratory for Radiometric Dating and Stable Isotope Research in Kiel (Table 1), using a Tandemtron

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4130 AMS system (High Voltage Engineering) with a single caesium sputter ion source. "CALIB rev 5.01" software modified with the IntCal04 dataset (Reimer et al., 2004) was used to calculate the calibrated ages.

### 3.3 Geochemical, stable carbon isotope, and granulometric analyses

5 Total carbon (TC), total organic carbon (TOC) and total nitrogen (TN) contents were measured using an elemental analyzer (Elementar Vario EL III). The TOC content was measured on carbonate-free samples (treated with 4 % HCl). The TC, TOC and TN results are given as weight per cent (wt %) and the C/N ratio is expressed as TOC/TN.

10 Stable carbon isotope ratios ( $\delta^{13}\text{C}$ ) in TOC were measured with a Finnigan DELTA S mass spectrometer coupled to a FLASH element analyser and a CONFLO III gas mix system. The analyses were accurate to  $\pm 0.2\%$ . The  $\delta^{13}\text{C}$  values are expressed in delta per mil notation ( $\delta$ , ‰) relative to the Vienna Pee Dee Belemnite (VPDB) standard.

15 Grain-size distribution analyses (for grain sizes between 0.375 and 1000  $\mu\text{m}$ ) were conducted on freeze-dried carbonate-free (treated with 10 %  $\text{CH}_3\text{COOH}$  for 12 h) and organic-free (treated with 3 %  $\text{H}_2\text{O}_2$  for a maximum of 4 weeks) subsamples from Section A, Section B (excluding the sample from 9.2 to 6.9 cm) and Core C using a laser diffraction particle sizer (Beckmann-Coulter LS 200). Grain-size fractions are given in volume per cent (vol %).

20 An end-member modelling algorithm was used to calculate robust end-members (EMs) from the total set of grain-size distributions with MATLAB software (R2012a; 7.14.0.739), using conceptual modifications proposed by Weltje (1997). The minimum number of potential EMs is thereby represented by the cumulative explained variance. We used the maximum value of the mean coefficient of determination ( $r^2$ ) to determine  
25 the maximum feasible number of EMs. The final EMs (robust EMs and the residual member) were determined with a test for robustness (Dietze et al., 2012).

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### 3.4 Analyses of moss remains and vascular plant macrofossils

Moss remains were analysed for every second sample from Core C (available volume of sieving residue after the KOH washing step for pollen preparation: 5–12 mL), the upper and the lower samples from Section A (50 mL), and 15 irregularly distributed samples from Section B (50 mL). The material was washed through 250 and 850  $\mu\text{m}$  mesh sizes. Moss remains larger than 850  $\mu\text{m}$  were analysed using ZEISS light and stereo microscopes (AxioLab 40, Stemi 2000-C) with 25 $\times$ , 100 $\times$ , and 400 $\times$  magnification; vascular plants were separated from mosses in order to evaluate their relative proportions. The identification of mosses was based on the relevant literature (Frahm and Frey, 2004; Lobin et al., 1995) and descriptions of Hedenäs (2008) from the Bryophyte Flora of North America website (<http://www.mobot.org>). Amblystegiaceae taxa were not differentiated in detail in Section B.

Vascular plant macrofossils were extracted from all depths in sections A and B by washing 25 mL subsamples through sieves with the same mesh sizes as used for mosses. Material in the +850 and 850–250  $\mu\text{m}$  fractions was again analysed using ZEISS light and stereo microscopes (AxioLab 40, Stemi 2000-C) and identified with the help of vegetation atlases and seed identification manuals (Anderberg, 1994; Beijerinck, 1947; Berggren, 1969, 1981; Katz et al., 1965) together with a regional reference collection at AWI Potsdam.

### 3.5 Pollen analysis

Pollen analyses were performed on all 27 samples from Core C. Processing of pollen samples in the laboratory included treatment with HCl (10%), KOH (10%) and HF (50%; 2 h boiling), followed by acetolysis, sieving (7  $\mu\text{m}$ ) in an ultrasonic bath, and mounting in glycerine. At least 350 terrestrial pollen grains were counted in all except three samples (from 20 to 17 cm) which had extremely low pollen concentrations. Pollen identifications were based on the relevant literature (Beug, 2004; Moore et al., 1991) and on a regional type slide collection of pollen types. The total pollen from

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arboreal and terrestrial non-arboreal taxa (excluding Cyperaceae) identified in each pollen spectrum is taken as 100% for the calculation of the pollen percentages. In addition, common non-pollen palynomorphs (NPPs, e.g. *Pediastrum*, *Botryococcus*, *Delitschia*, *Gelasinospora*, *Geumannomyces*, *Glomus*, *Macrobotus hufelandi*, and *Sordariales*) were counted from the pollen slides and their abundances expressed relative to the total terrestrial pollen.

### 3.6 Diatom analysis

About 0.5 g of each sample from Core C was used for diatom analysis. The processing of diatoms was conducted in accordance to Pestryakova et al. (2012): the calcareous and organic components were removed by heating with HCl (10%) and H<sub>2</sub>O<sub>2</sub> (30%). Cleaned diatom samples were mounted on microscope slides with Naphrax<sup>®</sup>. About 500 diatom valves were counted per slide along random transects. Samples were analysed using a ZEISS light microscope (AxioScope.A1) equipped with a differential interference contrast at a magnification of 1000×. The following literature was used for the determination of diatom taxa: Krammer and Lange-Bertalot (1986), Lange-Bertalot (1993), Lange-Bertalot and Metzeltin (1996), Lange-Bertalot et al. (1996), and Lange-Bertalot and Genkal (1999). The following regional works were also consulted: Komarenko and Vasilyeva (1975) “The diatoms of the USSR” (1974, 1988, 1992), “The diatoms of Russia” (2003), and Loseva (2000). The classification of diatom species used in this paper is based on the latest revisions of the *Achnanthes*, *Fragilaria*, *Navicula* and *Cymbella* genera (Krammer, 2000, 2002, 2003; Lange-Bertalot, 2001).

### 3.7 Statistical analysis

Statistical analyses were performed to deduce patterns in surface vegetation and in the diatom assemblages of Core C. We applied a non-metric multidimensional scaling (NMDS) analysis of species percentages using the Bray-Curtis dissimilarity index implemented in R version 2.15.0 (R Project for Statistical Computing, using the “vegan”

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package, Oksanen et al., 2007). Various supplementary parameters were added to the NMDS plots

## 4 Results

### 4.1 High-resolution spatial characteristics of the investigated polygon and its vegetation pattern

The 07-SA-LY low-centred polygon has dimensions of about 12 × 13 m (Fig. 2b). The outer margin of the investigated polygon approximately traces the ice-cracks that separated from its neighbouring polygons. The distance from the crack to the rim margin is 3–4 m. The polygon centre is filled by an intrapolygonal pond of water with approximate dimensions of 5 × 7 m. The ground surface measurements indicated that the polygon is not hydrologically closed. Transects A to C are up to 44 cm above the water level in the pond in contrast to transects H to K (plots 5 to 10), which cover a transition zone to a sunken crack in Transect L that has a depth of 62 cm below the water level in the pond. The maximum height of the other rims was 35 cm, and only minor depressions within these rims were detected e.g. from D11 to G11 and from K2 to K11 (Fig. 2b). The active layer depth ranges from 21 cm beneath the rims to 73 cm below the floor of the polygon's pond (Fig. 2c). This depth varies considerably beneath the rims but is relatively consistent beneath the waterbody (Fig. 2c). In general, the elevation correlated well with the active layer depth ( $r^2 = 0.81$ ). Along Transect G, the pH was found to be about 6.14 in the rim, 5.70 in the rim-to-pond transition zone, and 6.25 in the pond. The conductivity was highest on the rims (up to 84  $\mu\text{S cm}^{-1}$  in G1) and lowest in the pond water (e.g. 16  $\mu\text{S cm}^{-1}$  in G8).

A vegetation survey over the grid revealed differences in plant taxa between the rim, the rim-to-pond transition zone, and the pond (Fig. 3), as is clear from the NMDS analysis which yielded a stress value of 16 % (Fig. A1). Axis 1 correlates well with elevation and active layer depth and arrange the polygon vegetation along a rim-to-pond

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transect. The composition of the rim vegetation (appearing on the left side of the biplot) is highly variable and diverse, but is always dominated by vascular plants such as the ubiquitous shrub *Betula nana* (which is abundant over 2/3 of the grid), the two different shrubby *Salix* species (Type A with maximum abundances of 20 to 40 %, and Type B with a mean coverage of approximately 20 % in non-aqueous habitats), *Dryas punctata*, *Ledum palustre* and *Vaccinium* sp. Eight *Larix gmelinii* trees (with a maximum height of 4 m) were recorded from the upper rim sites. The rim-to-pond transition zone (the central part of the NMDS biplot) is characterized by the appearance of *Andromeda* and/or *Carex*. Relatively dry zones situated only few centimetres above the water level are covered by Type B *Salix* and *Vaccinium* sp., with abundances between 5 and 80 %. The pond vegetation (on the right side of the NMDS biplot) consists almost exclusively of the moss *Scorpidium scorpioides*; only in the shallowest parts is it intermixed with *Carex* taxa. In addition, individual plants of *Pedicularis* sp. appear in a small depression between the pond and the crack on Transect L.

## 4.2 Geochronology and age-depth relationships

The oldest ages were obtained from the bottom samples from Section A (420–539 cal AD) and Section B (417–568 cal AD) (Table 1). The calibrated ages obtained for Core C range between 1213 and 1635 cal AD, but do not show a clear age-depth relationship; at least the near surface date (from 3.5 cm depth) is the youngest of the geochronological record.

## 4.3 General properties of the sedimentary fill

The two sections (A and B) and the sediment core (Core C), from plots G1, G3, and G8, respectively, were analysed for their TOC content, TN content, and grain-size composition (Fig. A2 for sections A and B, Fig. 4 for Core C). The TOC values are relatively low in the lower parts of the profiles (0.3 to 1.3 wt. % in sections A and B; 1 to 4 wt. % in Core C) and higher in the uppermost (most recent) samples (32 to 33 wt. % in sections

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A and B; 26 wt. % in Core C). Due to the low TN contents (below the detection limit, i.e. < 0.1 %) reliable TOC/TN values are not available for the lower parts of sections A and B. The TOC/TN ratios in Core C vary between 7 and 26. The  $\delta^{13}\text{C}$  values range between  $-29.0$  and  $-26.3\text{‰}$  in Section A between  $-29.8$  and  $-25.3\text{‰}$  in Section B and between  $-30.6$  and  $-25.9\text{‰}$  in Core C. The ratio ranges differ markedly between the profiles, and also vary within each profile (Fig. 4, Fig. A2).

The grain-size composition of the polygon sediments is dominated by the fine to medium sand fraction (Fig. 4, Fig. A2). Only in the uppermost parts is silt about twice as abundant as sand. The inversion algorithm for end-member modelling of compositional data suggests the minimum number of EMs considering all weight transformations to be 3 or 4, and the maximum number to be 6. We therefore used different weight transformations in the models (with percentile ranges of  $P_4$ – $P_{96}$ ,  $P_5$ – $P_{95}$ ,  $P_6$ – $P_{94}$ ,  $P_7$ – $P_{93}$ ) and EMs ranging from 3 to 6 in order to extract final robust EMs and the residual member. One robust EM (EM1) and a residual member (RM; describing the remaining noise) were finally identified. They are presented in the loading plot (Fig. 5) together with their range of confidence as calculated from 16 end-member models. EM1 represents 75.3% of the total variance within the original data and corresponds to a medium to coarse sand fraction (180–720  $\mu\text{m}$ , peaking at  $\sim 270 \mu\text{m}$ ). Considering the goodness of variable and sample fit, we found a reasonably good modelling for Section A ( $r^2 = 0.92 \pm 0.14$ ), Section B ( $r^2 = 0.85 \pm 0.19$ ), and Core C ( $r^2 = 0.89 \pm 0.21$ ), for the robust end-member model. The median of EM1 scores for the whole of Core C (0.88) and Section A (0.75) are higher than the median score for Section B (0.58). All profiles show obvious variations in the EM1 to residual ratio (Fig. 4 for Core C, and Fig. A2 for sections A and B). Based on this information, Section B (rim-to-pond transition zone) and Core C (pond) can be correlated, indicating that both records cover an approximately similar period of time. In contrast, Section A shows a different sedimentation pattern and cannot be correlated with Section B, or with Core C.

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## 4.4 Bioindicators

We identified 12 moss taxa in Core C within the identifiable moss leaf fraction (Fig. 6). Most abundant were *Meesia triquetra* and *Scorpidium scorpioides*, while subdominant were *Drepanocladus* spp., *Hamatocaulis vernicosus* and *Calliergon* sp. *Bryum pseudotriquetrum* and *Rhizomnium pseudopunctatum* occurred only in trace quantities. Only a single leaf of *Aulacomnium turgidum* was recorded (not included in Fig. 6). The vertical distribution pattern of the main moss taxa in Section B (Fig. A3) was similar to that in Core C, which supports the correlation inferred from the grain-size record. *Meesia triquetra* occurred in most samples from Section B and Core C, but its abundance is highly variable. The upper parts of Section B (except for the uppermost 3.5 cm) and Core C are dominated by, or entirely composed of, *Scorpidium scorpioides* (from 10.5 to 0 cm in Core C and from 25 to 0 cm depth in Section B). The moss spectrum from the rim, i.e. from Section A (Fig. A4), which was only determined in the uppermost (1.2 cm) and lowermost (40.3 cm) samples, is completely different from the spectra in Section B and Core C. The moss community in the lowermost sample from Section A was composed of *Distichium capillaceum* (~ 15%), *Hylocomium splendens* (~ 14%), *Tomentypnum nitens* (9.5%), *Rhizomnium pseudopunctatum* (5%), *Bryum pseudotriquetrum* (~ 8%), *Hamatocaulis vernicosus* (7%), and *Campylium stellatum* (~ 2%). The genus *Aulacomnium* was only present in small quantities (< 0.2%). In contrast, *Aulacomnium* (with *Aulacomnium palustre* and *A. turgidum*; each ~ 13%) dominates the moss spectrum in the uppermost sample from Section A, which consists also of *Campylium stellatum* agg. (20%), *Hamatocaulis vernicosus* (< 10%) and *Dicranum acutifolium* (< 10%).

Needles of *Larix gmelinii* dominate the identifiable vascular plant macrofossil spectra in the upper parts of Section A (0.1–6.2 needles cm<sup>-3</sup>; Fig. A5) and Section B (5.7–19.0 needles cm<sup>-3</sup>; Fig. A6) while the lower part of Section B is dominated by *Carex* seeds (0.1–1.1 seed cm<sup>-3</sup>). Section A contains a relatively large concentration of *Andromeda polifolia* leaves (2.2 leaves cm<sup>-3</sup>) and *Dryas punctata* leaves

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(2.4 leaves cm<sup>-3</sup>). No identifiable macrofossils were found in Core C, but this may have been due to the small quantity of sediment analysed.

The relative proportion of total vascular plant to moss remains (the VPM ratio) was examined for all samples from Section A (Fig. A4), Section B (Fig. A3) and Core C (Fig. 6). Most of the samples from Core C were dominated by mosses, but the VPM ratio is relatively high at 24.5 cm (~ 0.16) and at 14.5 cm (0.11) core length, and also in the uppermost samples (e.g. at 2.5 cm, where it is 0.98). A similar pattern was found for Section B: high moss contents were determined for most of the lower horizons (except at 35.7 cm depth) and mosses dominate in most samples from above 24.2 cm depth, with a maximum VPM ratio of 0.8 at 10.4 cm and 8.1 cm depth. Mosses are particularly dominant (VPM ratio 0.3) in the uppermost part of Section B (above 1.2 cm depth). The VPM ratios for the lowermost and uppermost samples from Section A were 0.12 and 0.17, respectively.

Pollen analyses were performed on Core C. In total, 39 pollen taxa and 26 non-pollen palynomorphs were identified (Fig. 7). *Alnus fruticosa*-type, *Betula nana*-type and Poaceae are the most abundant pollen taxa. *Betula nana*-type remains quite stable throughout the core, while the abundances of Poaceae and *Alnus fruticosa*-type show opposing trends. *Larix* pollen occurs in low but stable abundances. Except at 18.5 cm core length the Cyperaceae (excluded from the pollen sum) have higher abundances than the terrestrial plants. We found relatively high quantities of the non-pollen palynomorphs *Botryococcus* and *Geumannomyces*.

Well-preserved diatom valves were found in all samples from Core C: 104 species were identified belonging to 35 genera. The most important genera were *Navicula* (11) *Eunotia* (11) *Pinnularia* (11), *Cymbella* (8 species), *Gomphonema* (7), *Fragilaria* (4) and *Stauroneis* (4) (Fig. 8). Most genera (63 % of the total) were represented by only one or two species, which is a regional characteristic of diatom communities in arctic waters. Most of the diatom species were rare: thus, for the statistical analyses we only used the 32 species that met our criteria (> 3 % over the whole core). Most samples were dominated by either benthic diatoms (e.g. *Navicula vulpina*, *Hantzschia amphioxys*,

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*Sellaphora pupula*) or epiphytic diatoms (e.g. *Gomphonema lagerheimii*, genus *Eunotia*, *Tabellaria flocculosa*), while planktonic taxa (*Aulacoseira subarctica*, *Stephanodiscus minutulus*, *Cyclotella ocellata*) were only dominant at 21.5 and 20.5 cm length in Core C. A second peak in *Stephanodiscus minutulus* was found at 7 cm length. The NMDS analysis (NMDS plot in Fig. 9) yielded a stress value of 17%. Samples from core lengths of 21.5 and 20.5 cm were excluded from the NMDS analysis due to their completely different diatom spectra. Most epiphytic taxa were located in the lower left part of the plot and most benthic taxa in the upper right part.

#### 4.5 Characterization of two different types of polygon pond sediment

Two sedimentary units that were only found in the polygon centre (Core C) and the rim-to-pond transition zone (Section B) can be distinguished on the basis of the various parameters described above, while the rim section (Section A) has a different composition. We defined the boundary between these units (Fig. 4) on the basis of the diatom NMDS analysis (Fig. 9). Meaningful sediment parameters (EMs, moss taxa, pollen taxa, non-pollen palynomorphs) were superimposed on the NMDS plot. An obvious relationship exists between diatom assemblages and these superimposed parameters, indicating that the diatom spectra are related to the end-member scores and to the relative abundances of *Scorpidium scorpioides*, *Meesia triquetra*, *Hamatocaulis vernicosus*, the green algae *Botryococcus*, and the *Alnus fruticosa* type pollen.

Unit 1 (Subunit 1.1: Core C, 27–24 cm core length and Section B 50.6–43.7 cm depth Subunit 1.2: Core C 14–9 cm core length and Section B, 27.6–16.1 cm depth. Subunit 1.3: Core C 3–0 cm core length and Section B 9.2–0 cm depth) is characterized by high scores for the residual member (RM) indicating a mixture of grain-size fractions. Both the TOC content and the TOC/TN ratio are quite high in these units. *Scorpidium* and some *Drepanocladus* species dominate the moss spectrum. In contrast, *Meesia triquetra* is only found in small quantities. Furthermore, Unit 1 is characterized by a dominance of epiphytic diatoms such as *Tabellaria flocculosa*, *Achnanthydium minutissimum*, and several *Eunotia* taxa (Figs. 8, 9). *Alnus fruticosa*-type pollen is abundant.

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Despite of these common features, each subunit (1.1, 1.2, and 1.3) has some specific characteristics that occur in both Core C and Section B. *Drepanocladus cossonii* is abundant in the samples from Subunit 1.1. They also contain large quantities of non-determined mosses (approximately 45 % of the total moos remains), which could also belong to the *Drepanocladus* genus. Subunit 1.2 contains a large proportion of *Scorpidium scorpioides* (40 %), but low abundances of *Meesia triquetra*. Subunit 1.3 is dominated by *Scorpidium* (95–100 %) in Core C, but the *Scorpidium* content in Section B is rather low (ranging from 86 % at 8 cm, to 4 % 1.2 cm) while other *Amblystegiaceae* were more abundant.

Unit 2 (Subunit 2.1: Core C 24–14 cm length and Section B 43.7–27.6 cm depth Subunit 2.2: Core C 9–3 cm length and Section B 16.1–9.2 cm depth) is characterized by high scores for EM1, indicating a dominance of well-sorted medium grained sand. The TOC values and TOC/TN ratio are mostly low. *Meesia triquetra* is present in most samples and often abundant. Most samples are dominated by benthic diatoms (e.g. *Navicula vulpina*, *Stauroneis phoenicenteron*, *Sellaphora pupula*). Furthermore, Poaceae are well represented in the pollen spectra. In addition, Ranunculaceae are abundant in some sections of Unit 2. Despite of these common characteristics subunits 2.1 and 2.2 have some specific characteristics. Subunit 2.1 contains abundant *Hamatoaulis vernicosus* (25–50 %) as well as abundant planktonic diatoms; it also has a low Cyperaceae content and generally a very low pollen concentration. The *Botryococcus* content shows a major peak in Subunit 2.2.

## 5 Discussion

### 5.1 Small-scale spatial structure of polygons

We found a strong correlation between thaw depth beneath the polygon and its surface elevation (Fig. 2a), i.e. shallow thaw depths beneath the elevated rims and greater thaw depths below the pond. It is assumed that the thick moss layer and the tall, dry



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vegetation on the polygon rims insulate the permafrost layer from the atmospheric temperatures (Blok et al., 2010; Luthin and Guymon, 1974). In contrast, the large thaw depth below the pond results from the high heat capacity of water an effect that is even enhanced by the positive feedback effect from thaw water (Woo and Guan, 2006). We observed a relatively homogenous distribution of neutral to slightly acidic pH in the water across the polygon which is in agreement with other studies on brown moss mires in arctic peatlands (Vanderpuye et al., 2002; Vitt, 1995; Wetterich et al., 2008).

The polygon is located in the Anabar–Lena subarctic zone (Isaev et al., 2010), with low summer temperatures and a short growing season. The present-day vegetation covering the investigated 07-SA-LY polygon (Fig. 3) is therefore dominated by typical elements of the arctic and boreal floral zone (Abaimov, 2010; Isaev et al., 2010; Jäger, 2005) such as *Andromeda polifolia* and *Vaccinium* sp. In these zones, sites with water-saturated soils are dominated by polygonal tundra or polygonal forest-tundra that typically show a well-developed microstructure of vegetation types tracing the hydrological microhabitats. We distinguished three different vegetation types corresponding to the dry polygon rim, the rim-to-pond transition zone, and the water-filled pond.

The vegetation on the rim contains taxa that prefer dry soils or at least avoid permanent water saturation (Höfle et al., 2013; Matveyeva, 1994; Minke et al., 2009). We observed three layers in the vegetation community of the polygon rim: (1) a tree and shrub layer composed of *Larix gmelinii*, *Betula nana*, *Salix* spp., and Ericaceae such as *Ledum palustre* and *Vaccinium* sp., (2) a dwarf-shrub and herb layer composed of *Polygonum* sp., *Pedicularis* sp., and Ericaceae and (3) a moss layer with xerophilous mosses (*Polytrichum* sp. and *Dicranum* sp.). The low resolution of environmental parameter measurement does not allow identification of structural parameters for the microhabitats on the rim. However previous investigations have shown that vascular plants and bryophytes trace small-scaled patterns related to the water table, thaw depth or pH (Ellis and Rochefort, 2004; Vitt, 1995; Walker et al., 2011).

The hydrological conditions in the rim-to-pond transition zone vary markedly during the short growing season (Økland, 1990). The assemblage comprises a low

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dwarf-shrub canopy of *Betula nana*, *Salix* type B, and *Andromeda polifolia* similar to the rim vegetation but containing a higher proportion of plants that have a greater water tolerance or occur commonly in raised bogs such as *Carex*. In contrast to our study, Minke et al. (2009) and Donner (2007) carried out comparable vegetation analyses in the northern Indigirka Lowland and identified *Comarum palustre*, *Drepanocladus revolvens* and *Sphagnum squarrosum* as common elements of vegetation in the rim-to-pond transition zone (in addition to *Carex* and *Andromeda polifolia*; de Klerk, 2008).

The water depth, and thus the position of the pond margin, depend on the water supply (e.g. from rainfall) and the evapotranspiration during summer. Since our field investigations took place in mid-August towards the end of the arctic summer we assume that the water level (17 cm at the deepest site) was the lowest of the summer season and that the pond at that time represented an area of permanent water (Boike et al., 2012). The transitional vegetation varies with the water depth with *Carex* spp. present in the shallow parts of the pond (as also reported by Tyler, 1981) The commonly submerged moss *Scorpidium scorpioides* that dominates (80–100 %) vegetation within the pond belongs to the Amblystegiaceae family which can assimilate under a thin ice cover and in low light conditions (Frahm, 2001). That special characteristic enables them to extend the short growing season and means that they are well-adapted to the harsh growing conditions in arctic polygons. The *Scorpidium scorpioides* community seems to represent a characteristic and widespread vegetation type as it has also been described by Liebner et al. (2011) in polygons of the Lena-Delta, Donner et al. (2007) in Siberian polygons and by Nebel and Philippi (2000) in mires of central Europe.

### 5.2 Age–depth relationships

Different ages have been inferred from different sites for when polygon mires first appeared in periglacial landscapes, sometime during the late Holocene. Botch et al. (1995) estimated ages of 2000 to 3000 yr for various mire zones in the European and Western Siberian parts of the Arctic. Other estimates have ranged from ~ 3600 yr cal BP in Canada (Fortier and Allard, 2004) to ~ 1200 yr cal BP for the Indigirka Lowland

in Yakutia (De Klerk et al., 2011) the latter age being roughly similar to our basal age from the rim and rim-to-pond transition sections. However, since our samples were only collected down to the permafrost table we cannot be sure that this is when the polygon sediments first started to accumulate. The radiocarbon dating of the basal samples indicates mean growth rates ranging between  $0.27 \text{ mm yr}^{-1}$  (Section A) and  $0.31 \text{ mm yr}^{-1}$  (Section B). These high sedimentation rates probably result from the dense bulk of moss, which provides a trap for minerogenic material. Furthermore, arctic mosses are known to be rapid biomass producers, according to Frahm (2001) producing up to  $100 \text{ gm}^{-2} \text{ yr}^{-1}$  depending on the local humidity. The age–depth relationship is thus rather uncertain for Core C, which makes direct comparisons with other palaeoclimatic records impossible.

### 5.3 Proxy value of the analysed parameters

A range of biotic and non-biotic sediment properties have been investigated and have contributed in various ways and to different extents to our reconstruction of polygon development. The most relevant proxies used to summarize the development of the pond and rim-to-pond transition zone in polygon 07-SA-LY are shown in Fig. 10. End-member scores summarize the granulometrical data in a meaningful way. The modelling was applied to separate sedimentary sections that differ in their main sources of minerogenic material (Weltje and Prins, 2003). In our case the end member EM1, which represents the fine to coarse sand fractions ( $180\text{--}720 \mu\text{m}$ ), probably represents the riverine transported sediment fraction Walling and Moorehead (1989) have shown that rivers in tundra lowlands transport sediments with a mean grain size (diameter) of  $150 \mu\text{m}$  (fine sand) during periods of normal water levels. The larger grain sizes in our record may indicate that a proportion of the minerogenic input may have accumulated during river high stands characterized by greater sediment transportability. We assume that the RM dominated by the silt fraction has various sources other than riverine sediments imported by lateral inflow of riverine water or sediments of aeolian origin.

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The general pollen composition is comparable to that of other late Holocene records from northern Siberia (Andreev et al., 2002, 2011), but is very rich in Cyperaceae compared to lake sediments (Klemm et al., 2013; Müller et al., 2010). Two pollen taxa may be used as climate proxies. We consider the percentages of Poaceae which are typically of tundra pollen spectra (Andreev et al., 2002; Klemm et al., 2013), to indicate cold conditions. In contrast, we consider high percentages of thermophilous *Alnus* to be an indicator of more temperate conditions (according to Salonen et al., 2011 for northern Europe and Klemm et al., 2013, for Siberian lowlands) We assume that a large proportion of the pollen supply to the polygon originates from more distant polygons and from the wider surroundings. The pollen signal thus reflects extra-local to regional vegetation changes rather than local changes within the studied polygon. In contrast, the non-pollen palynomorphs are mainly remains of aquatic organisms and thus represent a local signal (Birks and Birks, 1980).

Diatoms are abundant in arctic freshwater systems. We observed the most marked changes in the diatom record with respect to habitat preference. As reported in other studies from Yakutia (e.g. Pestryakova et al., 2012), the spectra are dominated by cosmopolitan taxa, with minor contributions of arcto-alpine and boreal forms. The most common species found by Patrick and Reimer (1966) in Alaska, by Moore (1975) in the southern part of Baffin Island, and by Komarenko and Vasilyeva (1975), Pestryakova (2008), and Palagushkina et al. (2012) in northern Yakutia (e.g. *Eunotia praeurupta*, *Tabellaria flocculosa*, *Achnantheidium minutissimum*) were also identified in our polygon and, as in these other studies, our shallow waterbody was dominated by benthic and epiphytic taxa. A previous study revealed that the diatom species *Tabellaria flocculosa* indicates relatively stable environmental conditions, while *Navicula vulpina* (benthic diatom species) prefers unstable slightly alkaline and oligotrophic phases. The proportion of *Navicula vulpina* was found to have increased in Unit 2.

We interpret the mosses *Meesia triquetra* and *Scorpidium scorpioides* to represent changing environmental dynamics in the water-logged polygon depression and the adjacent rim-to-pond transition zone. *Scorpidium scorpioides* (which is abundant in Unit

1) indicates a relatively stable pond with a steady water level, while *Meesia triquetra* (which is abundant in Unit 2) can grow under more unstable conditions with variable water level and prefers more terrestrial habitats than *Scorpidium scorpioides* (Nebel and Philippi, 2000).

5 The  $\delta^{13}\text{C}$  values from bulk-organic material show similar general fluctuations to the EM1 scores and the moss record. Changes in the composition of the vegetation and source water are possible reasons for variations in the  $\delta^{13}\text{C}$  record. Our analyses indicated that the bulk-organic material is dominated by moss remains. Mosses discriminate against heavy carbon isotopes during photosynthesis (although not as strong as vascular plants) due to the absence of any epidermis or stomata (Farquhar et al., 10 1989; Peisker, 1984). A high hydrogen carbonate content in the source water, probably contributed by riverine waters, may also lead to high isotope values.

We interpret the wt % of bulk organic material to be an indication of the minerogenic input (dilution effect) rather than of biomass productivity. However, a high minerogenic 15 input and a temporary low water table may also have promoted oxygen infiltration and thus produced aerobic conditions, which may in turn have reduced the organic matter content due to higher decomposition rates, as is also suggested by high TOC/TN ratios (Schirrmeister et al., 2011).

## 5.4 The general polygon development

20 We observed marked variations in the sedimentation record of the polygon. The grain-size data indicate that various sources contributed to the minerogenic input in Subunit 1.1 and that the influence of riverine flooding was low, which is supported by the relatively high TOC values and TOC/TN ratios. The vegetation was dominated by *Drepanocladus cossonii* which was found by Peterson and Billings (1980) to be characteristic of wet environments; de Klerk et al. (2011), also found only sparse distributions of this 25 species on dry rims. Thus the present-day polygon centre was a wet depression, even during the earliest recorded phase of development of the polygon. The diatom assemblage is dominated by epiphytes, and epiphytes that are specialized on moss require

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a particularly long ice-free period for their development (Moser et al., 2000) providing an indication that favourable growing conditions must have existed. According to Palagushkina et al. (2012), the presence of *Tabellaria flocculosa* indicates stable, slightly acidic, environmental conditions. The lower part of Subunit 1.1 was characterized by a relatively low water level and the upper part by a relatively high pH. Relatively warm conditions during that period can be inferred from the high abundances of *Alnus fruticosa*-type.

Repeated riverine flooding is indicated by the sandy organic-poor sediments of Subunit 2.1. The high rate of sediment supply probably led to a turnover in the moss coverage. Apart from the floating Amblystegiaceae (*Hamatocaulis vernicosus* and *Drepanocladus exannulatus*), there is an abundance of *Meesia triquetra* which is known as a pioneer species in young, post-glacial mires (Nebel and Philippi, 2000) characterized by minerogenic sediments. Furthermore, the greatest moss diversity was observed in this section, an additional indication of unstable growing conditions. Similarly, the diatom record was dominated by benthic diatoms (often cosmopolitan species with a large ecological amplitude) with preferences for disturbed (*Navicula vulpina*) or more aerophilic (*Hantzschia amphioxys*) habitats. Compared to Subunit 1.1, the diatoms in Subunit 2.1 reflect an increase in the pH (Dam et al., 1994) probably because the polygon's internal pH adjustment was repeatedly disturbed by fresh inputs of sediment and water. A sudden change to a short-period of dominance of planktonic diatoms indicates that the polygon was affected by an extreme flooding event (also indicated by an input of riverine *Pinus* pollen), and that the water level remained high for that period of time. A study by Moser et al. (2000) showed that in Canadian and Finnish lakes with depths of less than 5 m, planktonic diatoms never exceeded 20 % of the total diatom community. The vegetation on the polygon rim and in the uplands of the wider area surrounding the polygon contained relatively high proportions of Poaceae which is indicating cold conditions (Andreev et al., 2002). Early summer temperatures may have been decreasing at the time, resulting in reduced evapotranspiration within the polygon and an increase in the soil-moisture (Seppä and Birks, 2001), which may in

turn have led to a lower water absorbance capacity and eventually to more extreme flooding along the banks of the Anabar River.

The riverine sediment input was low during the accumulation of Subunit 1.2 The polygon was fully covered by moss as indicated by both the high VPM ratio and the high proportion of specialized epiphytic diatoms growing on mossy habitats (*Gomphonema lagerheimii*). The diatoms indicate a shallow water level between swamp mosses. Despite of these indications of rather stable conditions within the polygon, *Scorpidium scorpioides* which often prefers submerged habitats (Miller, 1980), suddenly became established in both Core C and Section B in the middle part of this subunit. Either the conditions within Subunit 1.2 changed from a temporarily water-filled depression to a permanently established pond or the hydrological conditions did not change but *Scorpidium scorpioides* just invaded the area at that time. Favourable growing conditions can be inferred from the high proportions of *Alnus fruticosa*-type and *Larix* in the polygon rim and upland vegetation, and the reduced proportion of Poaceae.

The grain-size data indicate that riverine flooding again became more common during Subunit 2.2, but not to the same extent as in Subunit 2.1. Interestingly, organisms with mutually exclusive habitat requirements such as *Navicula vulpina* and *Meesia triquetra* (indicators of unstable and disturbed conditions) and *Tabellaria flocculosa* and *Scorpidium scorpioides* (indicators of relatively stable conditions in an established environment) co-occur in one and the same sample which may indicate that the temporal resolution of the samples was not high enough to differentiate between individual subunits. However, the dominance of *Scorpidium scorpioides* (and *Gomphonema lagerheimii*) indicates that a permanent water-filled pond existed for at least some periods within this subunit. As was the case for Subunit 2.1, this polygon stage co-occurred with an expansion of tundra elements in the surrounding area.

The accumulation of Subunit 1.3 saw the start of conditions similar to those of the present day. The minerogenic influx is dominated by a fine silt fraction which we assume to be a result of lateral water flow through the polygon body. A stable stage is indicated by the dominance of *Tabellaria flocculosa* and by the monospecific *Scorpidium*

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*scorpioides* vegetation occupying the permanent waterbody. A decrease in *Scorpidium scorpioides* toward the top of the section can be recognized in Section B, which may indicate a slight lowering of the water level as a result of either an increase in evaporation or a decrease in water influx during recent times. The vegetation in the surrounding area was characterized by the highest *Alnus fruticosa*-type percentages and the lowest Poaceae pollen percentages of the entire section, particularly in the upper part of Subunit 1.3, indicating the most favourable growing conditions in the entire record.

## 5.5 Polygon development as a function of external controls and internal adjustment mechanisms

The studied polygon is part of a large polygonal field and has the typical form of a low-centred polygon with rims between the adjacent polygons (Chernov and Matveyeva, 1997; Meyer, 2003). We therefore assume that it represents a model polygon for the North Siberian Lowland and that inference concerning polygon development can, with caution, also be applied to other polygons in the same area. We hypothesize, on the basis of the spatial and temporal analyses of this polygon on the Anabar River lowlands of north-eastern Siberia, that polygon development is driven by both external controls and internal adjustment processes. Factors acting on a (supra-) regional scale such as climate, overall soil substrate, or river flooding exert pulses of external influence that disturb the dynamic equilibrium within the polygon that is otherwise maintained by local regulatory mechanisms such as plant colonisation (Epstein et al., 2012; Shaver and Chapin III, 1991), moisture, summer heating and soil thawing (Peterson and Billings, 1980; Vitt, 1995; Woo and Guan, 2006), and ice wedge growth or peat accumulation (Chernov and Matveyeva, 1997). Through these internal adjustments the equilibrium in the biotope is reconfigured and feedbacks can occur that either reduce or amplify the impact of the external controls (Harding et al., 2002; Oechel and Vourlitis, 1994). Climatic factors and a loose soil substrate may represent the primary external controls in the initiation and development of polygons. Rapidly falling temperatures in autumn lead to an increase in tensile elongation within the soil substrate, which then begins to

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crack (French, 2007). Meltwater or river water fills these cracks during the subsequent spring, and then freezes forming ice-veins which depending on their length thaw during the following summer and either fill with sediment or initiate perennial ice-wedges (Romanovsky, 1985). The annual growth of the wedges (approximately  $1 \text{ mm yr}^{-1}$ : Oven-

5 den, 1990) lifts the ambient soil substrate and forms elevated polygonal rims. As a consequence of the rim uplift a depression then develops in the polygon centre. We interpret an initial depression stage with waterlogged conditions reflected by deposition of the first unit followed by successive deepening of the depression during subsequent time periods (Chernov and Matveyeva, 1997).

10 It is assumed that the crackinitiated shape of a polygon is the result of interactions between its position within the landscape and the nature of the soil substrate (French, 2007; Minke et al., 2007). Our polygon is situated within an old meander belt of the Anabar River.

15 Polygons in Canada have been found to develop on sediments of various glacial and fluvial origins (Burn, 1990; Fortier and Allard, 2004). However, the original substrate for the polygon field in which our studied polygon is located probably consisted of river sands from the Anabar River, but these may have already had a history of soil development. Although polygons in Siberian lowlands typically occur along river banks, the rivers themselves have seldom been considered to be drivers of polygon develop-

20 ment, which is what we propose for our polygon record from the Anabar River lowlands. We assume that the polygon's development was affected by slow lateral water inflow through the polygon during the accumulation of the sediments of Unit 1. In contrast, during the accumulation of Unit 2 the polygon was repeatedly flooded and the water velocity was high enough to transport medium-grained, and even coarse-grained,

25 sand. However, the flooding would only have been seasonal as the vegetation (represented by the moss and diatom record) does not indicate a permanently high water level during the major growing season. A long-term flood probably did occur during the deposition of the sediments at 20.5 cm core length in Core C, which are characterized by a high planktonic diatom component. The development of the polygon was

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interrupted by the impact of this new control, but subsequently returned to the previous level of development. We assume that a relatively stable stage was established during the accumulation of Unit 1, as suggested by the relatively low diversity in the structure of the plant cover within the pond, the dominance of organisms with a narrow range of tolerance, the complex plant-to-plant interactions (e.g. a high number of epiphytic diatoms), and a generally high bioproductivity. In contrast, some phases during the accumulation of Unit 2 represent long-term disturbance and instability in the sedimentary succession within the low-centred polygon initiated by repeated phases of sediment input as indicated for example by the large number of benthic taxa (which are often pioneer taxa) and the high diversity of mosses present (as concurrence may not yet have led to species exclusion).

Our records suggest that the controlling factors discussed above may be interdependent with regards to their timing and strength. Flooding, for example, was found to be more extreme during the colder climatic periods. Thus the response of a polygon to climate change is complex and difficult to predict. We tentatively infer from our records that higher temperatures and reduced flooding would result in the polygon communities being characterized by high moss productivity (with a high moss to minerogenic sediment ratio), a low pH, and a specialized diatom community that is epiphytic on moss.

## 6 Conclusions

The low-centred polygon in this study is at an advanced stage of succession in space and time, with distinctive vegetation patterns. The main conclusions drawn from our research are as follows:

(1) The present-day spatial vegetation patterns are closely related to abiotic factors: we observed the elevation above or below the water level to be the main controlling factor. Rim vegetation (dominated by *Larix*, *Vaccinium*, *Dryas* or *Poaceae*) can be differentiated from vegetation in the rim-to-pond transition zone (*Andromeda*, *Pedicularis*

or *Carex*) and in the pond (comprised solely of *Scorpidium scorpioides* indicating that *Carex* is limited by a higher water level in the pond).

(2) The peat composition differs depending on the stability of the environmental conditions. Stable conditions occurred during periods with little or no riverine flooding, when submerged moss communities overgrown by epiphytic diatoms dominated the polygon centre. During times of repeated riverine flooding (indicated by medium to coarse grained sand within the deposits), a community became established that was adapted to instability and sediment supply (i.e. a community in which *Meesia triquetra* is abundant in the moss spectra, and benthic taxa in the diatom record). However, despite changes in the composition of the pond community, the general position of the pond and rim has not changed over the last ~ 1500 yr of recorded peat accumulation.

(3) The development of polygons in flood plains is not straightforward and is influenced by the river, which may in turn be related to climatic conditions. However, various internal regulating factors within the polygon react to these changes and affect succession after such pulses of external influence, readjusting the system to a dynamic equilibrium under the new environmental conditions.

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**Table 1.** Results of AMS dating indicating an age of  $\sim 1600$  yr BP for the bottom samples from sections A and B, and relatively homogeneous ages for Core C (core length: 27 cm).

Section/ core	Depth [cm]	Radiocarbon age [yr BP]	[cal yr AD] ( $2\sigma$ range)	Fraction (C content)	Sample ID Lab.
Section A	19.6	$426 \pm 27$	1426–1495	Plant seed (1.9 mg)	KIA39094
	40.3	$1587 \pm 24$	420–539	Plant seed, wood (2.3 mg)	KIA38752
Section B	49.5	$1566 \pm 35$	417–568	Plant seed (0.7 mg)	KIA38753
Core C	3.5	$342 \pm 25$	1536–1635	Mosses ( <i>Amblystegiaceae</i> ), (0.97 mg)	KIA46253
	5.5	$791 \pm 25$	1213–1275	Mosses ( <i>Amblystegiaceae</i> ), (1.22 mg)	KIA46254
	6.5	$544 \pm 23$	1391–1431	Plants (2.47 mg)	KIA46255
	8.5	$535 \pm 25$	1392–1436	Mosses (unidentified species) (1.1 mg)	KIA44742
	12.5	$435 \pm 20$	1430–1471	Mosses (unidentified species) (2.5 mg)	KIA44743
	15.5	$568 \pm 23$	1387–1419	Mosses ( <i>Amblystegiaceae</i> ) (2.47 mg)	KIA46256
	20.5	$699 \pm 22$	1368–1382	Mosses ( <i>Amblystegiaceae</i> ) (2.3 mg)	KIA44744
	24.5	$518 \pm 22$	1398–1440	Mosses ( <i>Amblystegiaceae</i> ) (1.53 mg)	KIA46257
	25.5	$407 \pm 26$	1436–1516	Plants (3.5 mg)	KIA39095

**Table A1.** Legend for the vegetation types (Fig. 1c) according to the agriculture map of Yakutia (Matveev, 1989).

<b>Subarctic tundra vegetation</b>	
1:	Widely spaced dwarf shrub tundra ( <i>Betula exilis</i> , <i>Salix pulchra</i> ), dwarf shrub tundra ( <i>Vaccinium vitis-idaea</i> , <i>V. uliginosum</i> , <i>Ledum palustre</i> ), lichens and mosses ( <i>Aulacomnium turgidum</i> , <i>Hylocomium splendens</i> var. <i>alaskanum</i> , <i>Cetraria cucullata</i> ), humps together with tundra bogs
2:	Widely spaced dwarf shrub tundra ( <i>Betula exilis</i> , <i>Salix pulchra</i> ), dwarf shrub tundra ( <i>Vaccinium vitis-idaea</i> , <i>V. uliginosum</i> , <i>Ledum palustre</i> ), lichens and mosses ( <i>Aulacomnium turgidum</i> , <i>Hylocomium splendens</i> var. <i>alaskanum</i> , <i>Cetraria cucullata</i> ), humps mixed with sections of widely spaced larch forests ( <i>Larix gmelinii</i> ) with lichens and mosses with <i>Ledum</i> and stone bramble
3:	Widely spaced dwarf shrub tundra ( <i>Betula exilis</i> , <i>Salix pulchra</i> ), dwarf shrub tundra ( <i>Cetraria cucullata</i> together with <i>Alectoria ochroleuca</i> , <i>Coelocaulon divergens</i> ) and tundra bogs
<b>Rock desert and mountain tundra</b>	
4:	Lichens ( <i>Alectoria ochroleuca</i> , <i>Coelocaulon divergens</i> ) and dwarf shrubmountain tundra ( <i>Dryas punctata</i> , <i>Cassiope tetragona</i> )
<b>Vegetation of riparian areas, valleys and coasts</b>	
5:	Polygonal peat tundra with thickets of <i>Duschekia fruticosa</i> , willow ( <i>Salix pulchra</i> , <i>S. glauca</i> ), <i>Eriophorum vaginatum</i> and <i>Carex stans</i> on humps and depressional grass bogs together with various types of willow ( <i>Salix pulchra</i> , <i>S. alaxensis</i> ), grass bogs ( <i>Eriophorum polystachyon</i> , <i>Arctophila fulva</i> ) and valley meadows from horsetail and grass ( <i>Equisetum arvense</i> , <i>Poa alpigena</i> )
<b>Vegetation in the river valley with dominant larch forests</b>	
6:	Leaves mosses ( <i>Aulacomnium turgidum</i> , <i>Hylocomium splendens</i> ) and <i>Sphagnum</i> mosses ( <i>Sphagnum balticum</i> , <i>Sphagnum warnstorffii</i> ) with tundra bogs

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**Table A2.** Summary list of determined diatom taxa (Core C) with life form information.

Taxon code	Taxon name	Author	Life form
AchMin	<i>Achnantheidium minutissimum</i>	(Kützing) Czarnecki	Epiphytic
AchSp.	<i>Achnanthes</i> spp.		Epiphytic
NavBry	<i>Adlafia bryophila</i>	Peterson	Benthic
Amplna	<i>Amphora inariensis</i>	Krammer	Benthic
AulGra	<i>Aulacoseira granulata</i>	(Ehr.) Simonsen	Planktonic
AulSp.	<i>Aulacoseira</i> spp.		Planktonic
AulSub	<i>Aulacoseira subarctica</i>	(Müller) Haworth	Planktonic
CalSp.	<i>Caloneis</i> spp.		Benthic
CalThe	<i>Caloneis thermalis</i>	(Grun.) Krammer	Benthic
CocPla	<i>Cocconeis placentula</i>	Ehr.	Epiphytic
CycMen	<i>Cyclotella meneghiniana</i>	Kützing	Planktonic
CycOce	<i>Cyclotella ocellata</i>	Pant.	Planktonic
CycSp.	<i>Cyclotella</i> spp.		Planktonic
CymGra	<i>Cymbella gracilis</i>	(Ehr.) Kützing	Epiphytic
CymHet	<i>Cymbella heteropleura</i>	(Ehr.) Kützing	Epiphytic
CymInc	<i>Cymbella incerta</i>	(Grun.) Cleve	Epiphytic
CymMes	<i>Cymbella mesiana</i>	Cholnoky	Epiphytic
CymPau	<i>Cymbella paucistriata</i>	Cleve-Euler	Epiphytic
CymPrx	<i>Cymbella proxima</i>	Reimer.	Epiphytic
Cym	<i>Cymbella</i> spp.		Epiphytic
CymTyn	<i>Cymbella tynnii</i>	Krammer	Epiphytic
CymCus	<i>Cymbopleura cuspidata</i>	Kützing	Epiphytic
CymHub	<i>Cymbopleura hybrida</i>	(Grunow) Krammer	Epiphytic
CymEhr	<i>Cymbopleura inaequalis</i>	(Ehrenberg) Krammer	Epiphytic
DipSp.	<i>Diploneis</i> spp.		Benthic
CymCes	<i>Encyonema cesatii</i>	(Rabenh.) Grun.ex A.S.	Epiphytic
CymSil	<i>Encyonema silesiacum</i>	(Bleisch) Mann	Epiphytic

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**Table A2.** (Continued.)

Taxon code	Taxon name	Author	Life form
EpiAdn	<i>Epithemia adnata</i>	(Kutz.) Breb.	Epiphytic
EpiSor	<i>Epithemia sorex</i>	Kützing	Epiphytic
EpiSp.	<i>Epithemia</i> spp.		Epiphytic
EunArc	<i>Eunotia arcus</i>	Ehr.	Epiphytic
EunBil	<i>Eunotia bilunaris</i>	(Ehr.) Mills.	Epiphytic
EunFab	<i>Eunotia faba</i>	(Ehr.) Grun.	Epiphytic
EunFor	<i>Eunotia formica</i>	Ehr.	Epiphytic
EunLun	<i>Eunotia lunaris</i>	Brevisson	Epiphytic
EunMin	<i>Eunotia minor</i>	(Kutz.) Grun.	Epiphytic
EunMon	<i>Eunotia monodon</i>	Ehr.	Epiphytic
EunPec	<i>Eunotia pectinalis</i>	(Dillw.) Rabenh.	Epiphytic
EunPra	<i>Eunotia praerupta</i>	Ehr.	Epiphytic
EunPse	<i>Eunotia pseudopectinalis</i>	Hust.	Epiphytic
EunSp.	<i>Eunotia</i> spp.		Epiphytic
FraCap	<i>Fragilaria capucina</i>	Desm.	Planktonic
FraCon	<i>Fragilaria construens</i>	Hust.	Epiphytic
FraSp.	<i>Fragilaria</i> spp.		Epiphytic
FraUln	<i>Fragilaria ulna</i>	(Nitzsch.) Lange-Bertalot	Epiphytic
GomAcu	<i>Gomphonema acuminatum</i>	Ehr.	Epiphytic
GomCla	<i>Gomphonema clavatum</i>	Ehr.	Epiphytic
GomGra	<i>Gomphonema gracile</i>	Ehr.	Epiphytic
GomLag	<i>Gomphonema lagerheimii</i>	Cleve	Epiphytic
GomPar	<i>Gomphonema parvulum</i>	Kützing	Epiphytic
GomSp.	<i>Gomphonema</i> spp.		Epiphytic
GomSut	<i>Gomphonema subtile</i>	Ehr.	Epiphytic
HanAmp	<i>Hantzschia amphioxys</i>	(Ehr.) Grun.	Benthic
HanElo	<i>Hantzschia elongata</i>	(Hantzsch.) Grun.	Benthic
HanSp.	<i>Hantzschia</i> spp.		Benthic
MelVar	<i>Melosira varians</i>	Ag.	Planktonic

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**Table A2.** (Continued.)

Taxon code	Taxon name	Author	Life form
NavAmp	<i>Navicula amphibola</i>	Cleve	Benthic
NavCry	<i>Navicula cryptocephala</i>	Kützing	Benthic
NavIgn	<i>Navicula ignota</i>	Krasske	Benthic
NavPss	<i>Navicula pseudotuscula</i>	Hust.	Benthic
NavRad	<i>Navicula radiosa</i>	Kützing	Benthic
NavRhy	<i>Navicula rhynchocephala</i>	Kützing	Benthic
NavSem	<i>Navicula semen</i>	Ehr.	Benthic
NavSp.	<i>Navicula</i> spp.		Benthic
NavLan	<i>Navicula trivialis</i>	Lange-Bertalot	Benthic
NavVul	<i>Navicula vulpina</i>	Kützing	Benthic
NavJaa	<i>Navicula jaagii</i>	(Krammer and Lange-Bertalot)	Benthic
NesSp	<i>Neidiodopsis</i> spp.		Benthic
NeiAmp	<i>Neidium ampliatum</i>	(Ehr.) Krammer	Benthic
NeiDil	<i>Neidium dilatatum</i> var. <i>jacuticum</i>	I.Kiss.	Benthic
NeiHit	<i>Neidium hitchcockii</i>	(Ehr.) Cleve	Benthic
NeiIri	<i>Neidium iridis</i>	(Ehr.) Cleve	Benthic
NeiSp.	<i>Neidium</i> spp.		Benthic
NitAmp	<i>Nitzschia amphibia</i>	Grun.	Benthic
NitSp.	<i>Nitzschia</i> spp.		Benthic
PinBor	<i>Pinnularia borealis</i>	Ehr.	Benthic
PinBrc	<i>Pinnularia brevicostata</i>	Cleve	Benthic
Pingib	<i>Pinnularia gibba</i>	Ehr.	Benthic
PinInr	<i>Pinnularia intermedia</i>	(Lagerstedt) Cleve	Benthic
PinInt	<i>Pinnularia interrupta</i>	W.Sm.	Benthic
PinMic	<i>Pinnularia microstauron</i>	(Ehr.) Cl.	Benthic
PinSp.	<i>Pinnularia</i> spp.		Benthic
PinSth	<i>Pinnularia stauroptera</i>	(Grun) Cleve	Benthic
PinStr	<i>Pinnularia streptoraphe</i>	Cleve	Benthic

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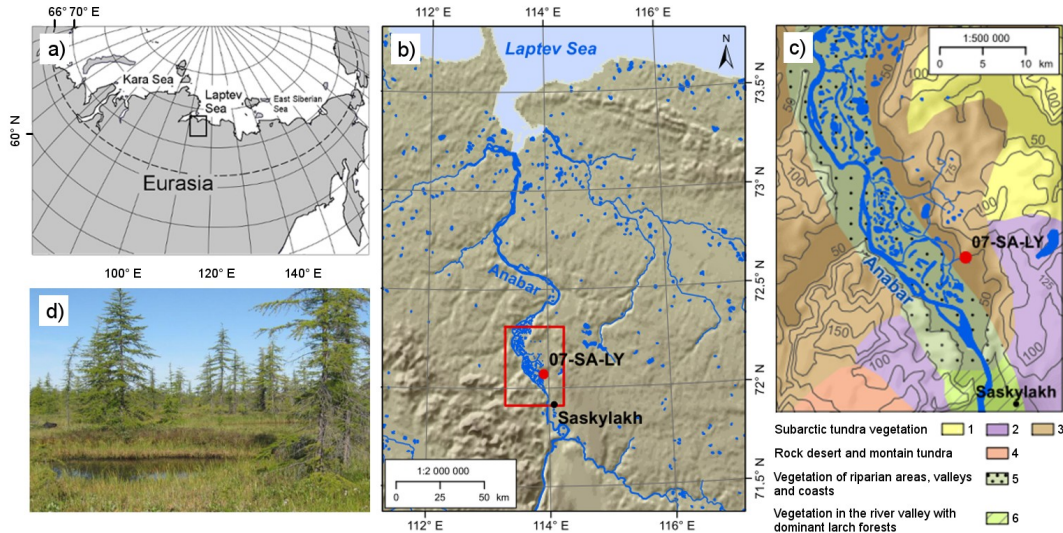
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**Table A2.** (Continued.)

Taxon code	Taxon name	Author	Life form
PinSgi	<i>Pinnularia subgibba</i>	Krammer	Benthic
PinVir	<i>Pinnularia viridis</i>	(Nitzsch.) Ehr.	Benthic
NavGas	<i>Placoneis gastrum</i>	(Ehr.) Meresch.	Benthic
FraBre	<i>Pseudostaurosira brevistriata</i>	(Grunow) Williams and Round	Epiphytic
CymSin	<i>Reimeria sinuata</i>	(Gregory) Kociolek and Stoermer	Epiphytic
RhoCur	<i>Rhoicosphenia abbreviata</i>	(Ag.) L.-B.	Epiphytic
RhoGib	<i>Rhopalodia gibba</i>	(Ehr.) O. Mull.	Epiphytic
SelLae	<i>Sellaphora laevisissima</i>	(Kütz.) D. G. Mann	Benthic
SelPup	<i>Sellaphora pupula</i>	Kutz.	Benthic
StaAgr	<i>Stauroneis agrestis</i>	Peterson	Benthic
StaAncGr	<i>Stauroneis anceps et f. gracilis</i>	Rabh.	Benthic
StaPho	<i>Stauroneis phoenicenteron</i>	(Nitzsch.) Ehr.	Benthic
StaSp.	<i>Stauroneis</i> spp.		Benthic
FraConVe	<i>Staurosira venter</i>	(Ehr.) Cleve and Möller	Epiphytic
FraPin	<i>Staurosirella pinnata</i>	Ehr	Epiphytic
SteMin	<i>Stephanodiscus minutulus</i>	(Kütz.) Cleve and Möller	Planktonic
SteSp.	<i>Stephanodiscus</i> spp.		Planktonic
TabFen	<i>Tabellaria fenestrata</i>	(Lungb.) Kutz.	Planktonic
TabFlo	<i>Tabellaria flocculosa</i>	(Roth.) Kutz.	Epiphytic
TetDla	<i>Tetracyclus glans</i>	(Ehr.) Mills	Epiphytic

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**Fig. 1.** Location of the study area. The studied polygon (07-SA-LY) is located south of the Taymyr Peninsula (a) and 160 km south of the Laptev Sea coastline in the vicinity of the Anabar River North Siberian Lowland (b), within an abandoned meander on the river's floodplain (c) *Larix* is sparsely distributed throughout the area (d) and the land surface is covered with a shrub-moss community (Photo: U. Herzsuh, August 2007).

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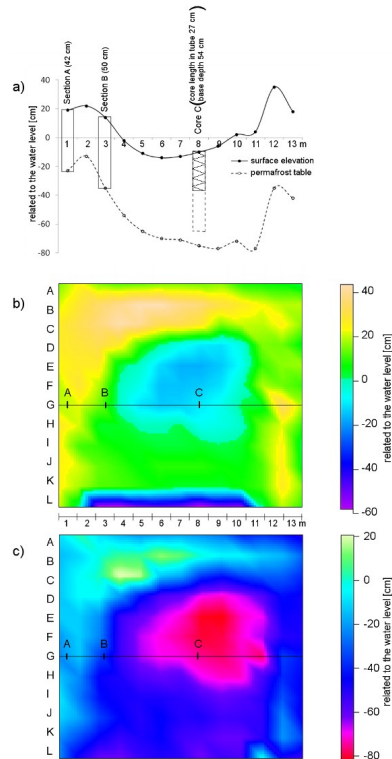
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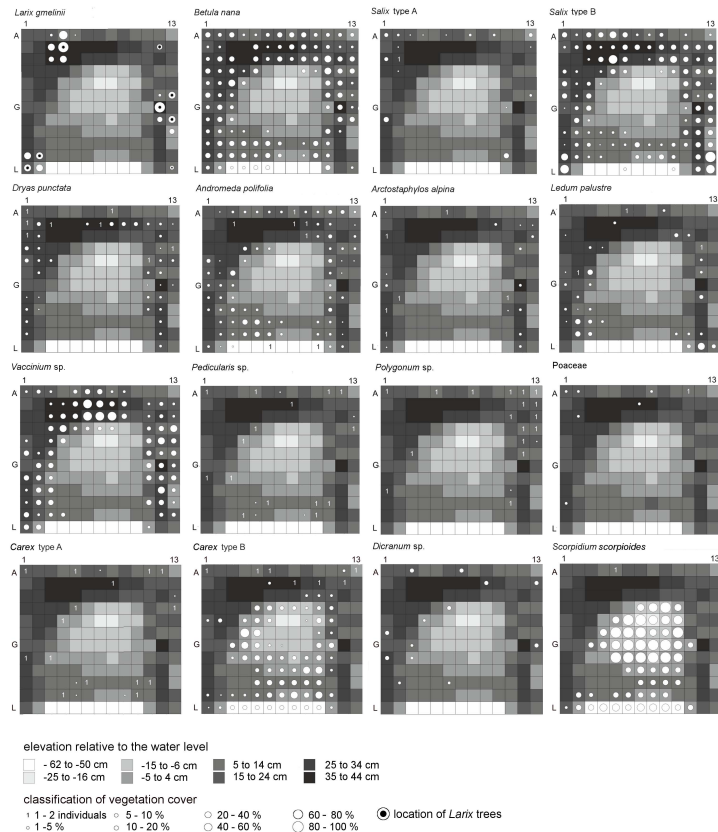




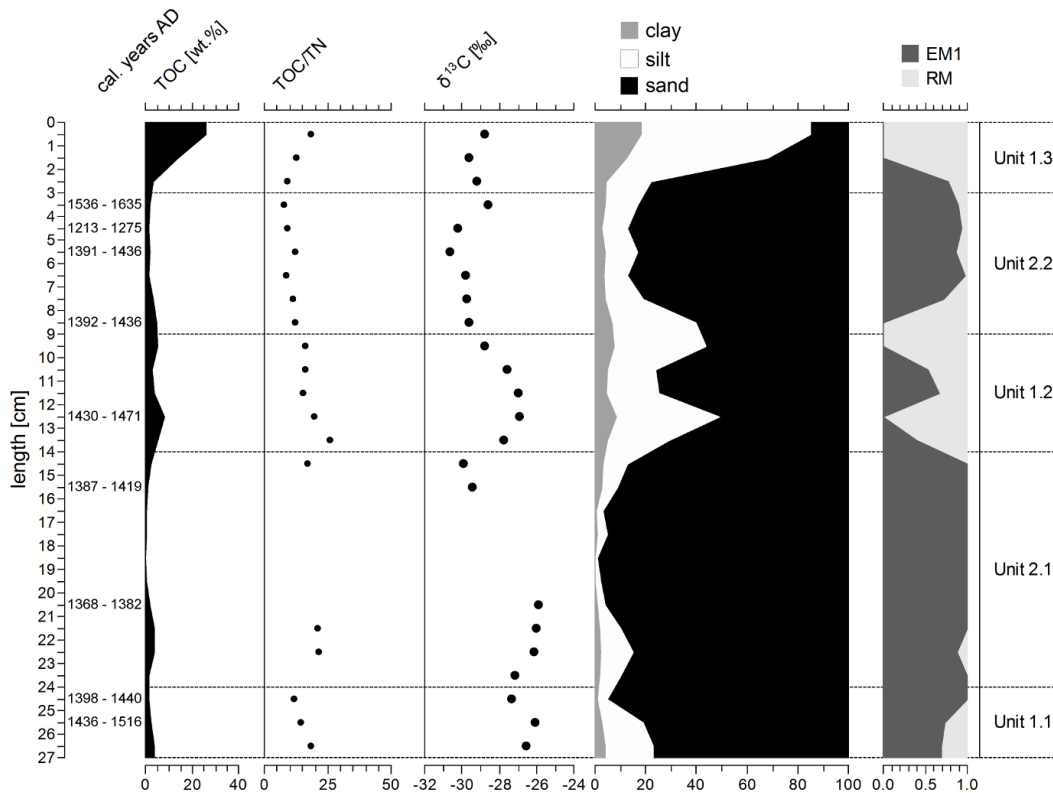
**Fig. 2.** The 07-SA-LY polygon was divided into a 12 m (A–L) by 13 m (1–13) grid of square metre plots. A surface transect from which all three profiles were taken, was positioned in the grid along line G. **(a)** Surface elevation and thaw depth relative to the water level in the polygon pond along the surface transect (Transect G), illustrating the differences between a shallow thawing depth below the rims in contrast to conditions beneath the pond. (Note: Core C was highly compressed, being reduced from 54 to 27 cm during sampling.) **(b)** The elevation of the ground surface relative to the water level in the pond. **(c)** The elevation of the permafrost table, relative to the water level in the pond.

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**Fig. 3.** Distribution of plant taxa presented in vegetation abundance/cover classes (after Braun-Blanquet, 1964) represented by circles and related to the elevation (in cm, relative to the water level in the pond). Double circles indicate the locations of *Larix* trees (after Wolter, 2010).



**Fig. 4.** Organic matter (TOC, TOC/TN,  $\delta^{13}\text{C}$ ) and granulometrical characteristics of Core C.

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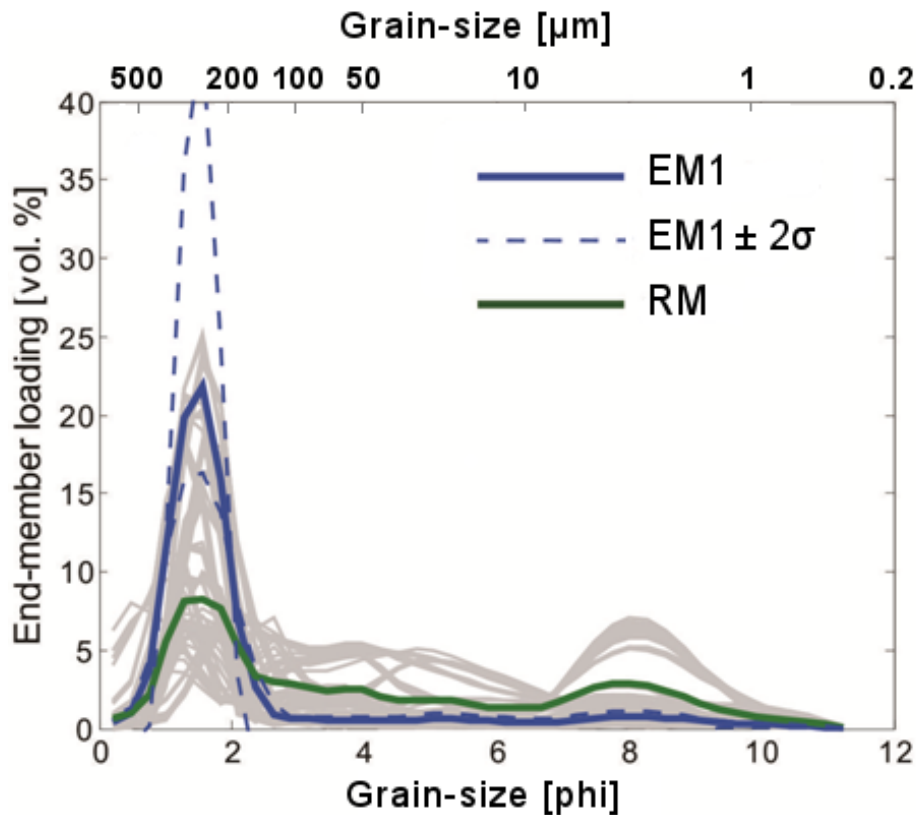
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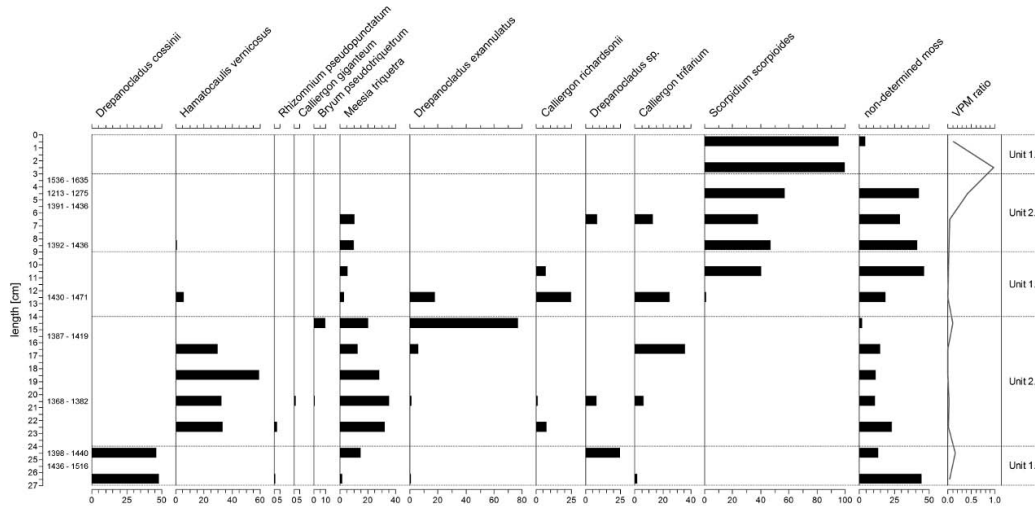




**Fig. 5.** End-member modelling for samples from sections A, B and Core C. Robust end-member scores give the proportion of variance explained by the robust end-members (EM1) and the residual member (RM) for each time slice. The solid grey lines are all end-members from the 16 end-member modelling versions. The blue dotted line is the range of confidence ( $\pm 2\sigma$ ) of the EM1.

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**Fig. 6.** The moss analyses revealed a dominance of Amblystegiaceae in Core C. The different species of this family vary along the units with a recent dominance of *Scorpidium scorpioides*. *Meesia triquetra* is abundant as species of the Amblystegiaceae in Subunits 2.1 and 2.2.

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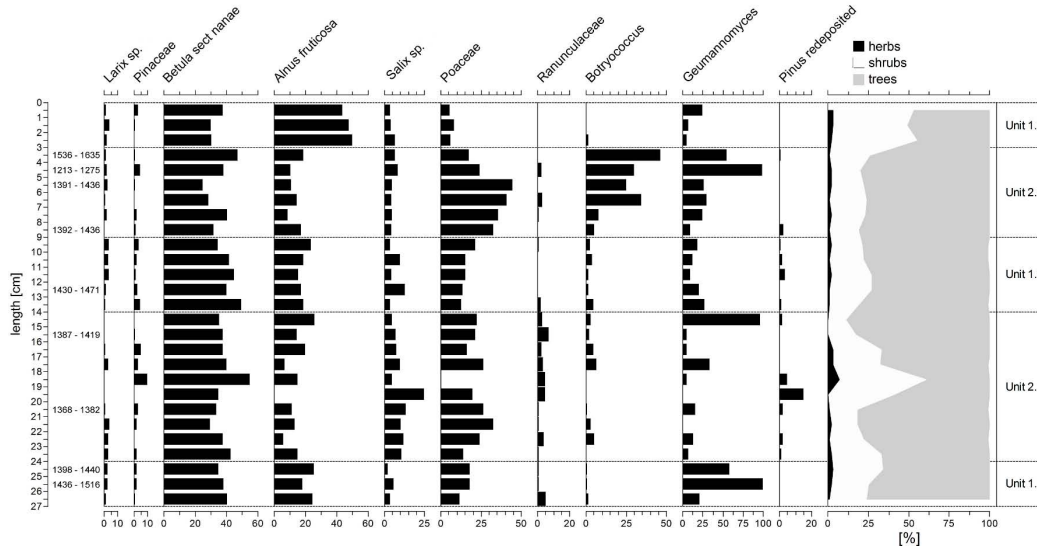
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**Fig. 7.** Pollen percentage plot showing the important palynomorphs and non-palynomorphs (*Botryococcus* and *Geumannomyces*) from Core C.

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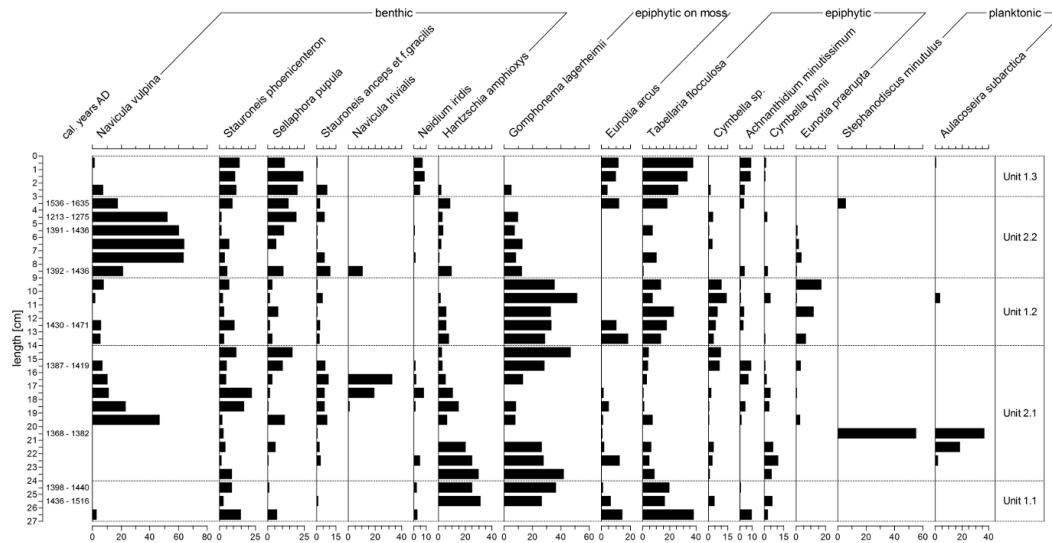
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**Fig. 8.** Four ecological sections showing the important species in the diatom assemblage from Core C. The occurrence of epiphytic diatoms correlates mainly with Unit 1, while Unit 2 is dominated by benthic diatoms. Planktonic diatoms are abundant between 22 and 20 cm core depth.

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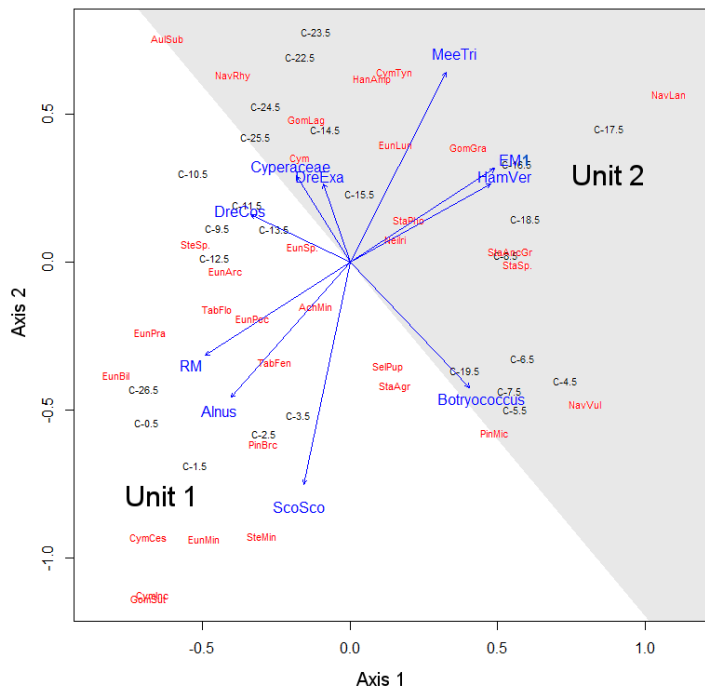
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**Fig. 9.** NMDS plot of the diatom community (excluding the samples at 21.5 and 19.5 cm length) overlain by the calculated robust end-member (EM1) and the residual member (RM) together with data on important species of moss, pollen and non-palynomorphs. The EM1, the RM, the mosses *Scorpidium scorpioides* (ScoSco), *Meesia triquetra* (MeeTri), *Hamatocaulis vernicosus* (HamVer), the palynomorph *Alnus fruticosa* and the non-palynomorph *Botryococcus* divide the diatom spectra into our assumed units (Unit 1 and Unit 2). Other parameters were tested (DreCos – *Drepanocladus cossonii*; DreExa – *Drepanocladus exannulatus*), but were not significant. The sample scores labelled within Core C the length position of the sample. (For abbreviations of diatom species see Table A1.)

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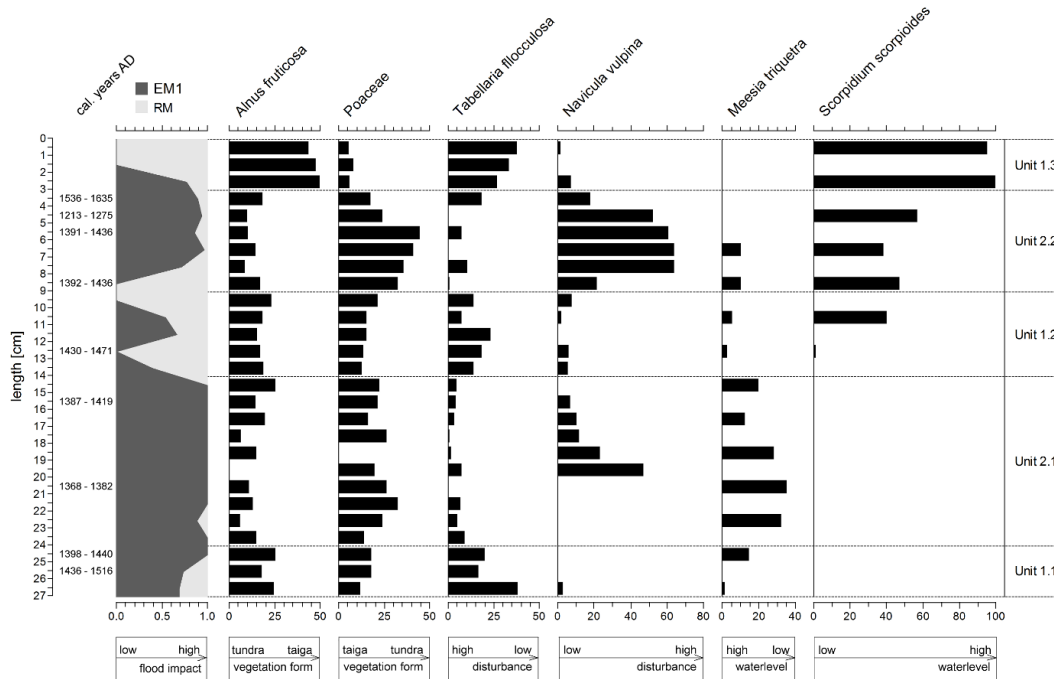
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**Fig. 10.** Summary plot of important indicator records. The variations in end-member, pollen and diatom data indicate the boundaries between Unit 1 and Unit 2, while the two moss species reflect the local hydrology in plot G8 (Core C) ranging from a water-saturated to a permanently water-filled location.

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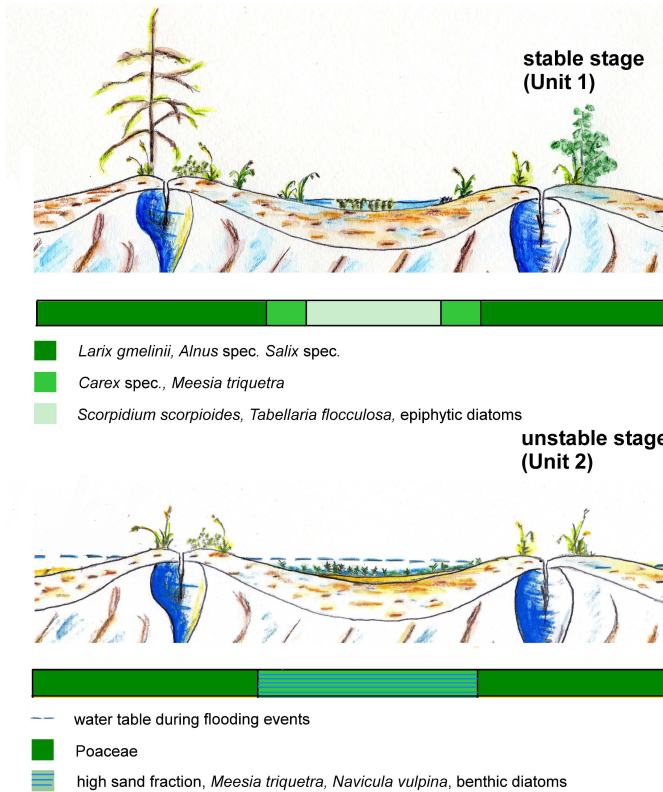
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**Fig. 11.** Presentation showing vegetation cover during stable (Unit 1) and unstable (Unit 2) stages. For Unit 1 we detected a shrub-tussock tundra with scattered occurrences of *Larix* while Unit 2 had a high proportion of sand which affected plant communities on the rim and in the water-saturated depression (with *Meesia triquetra*).

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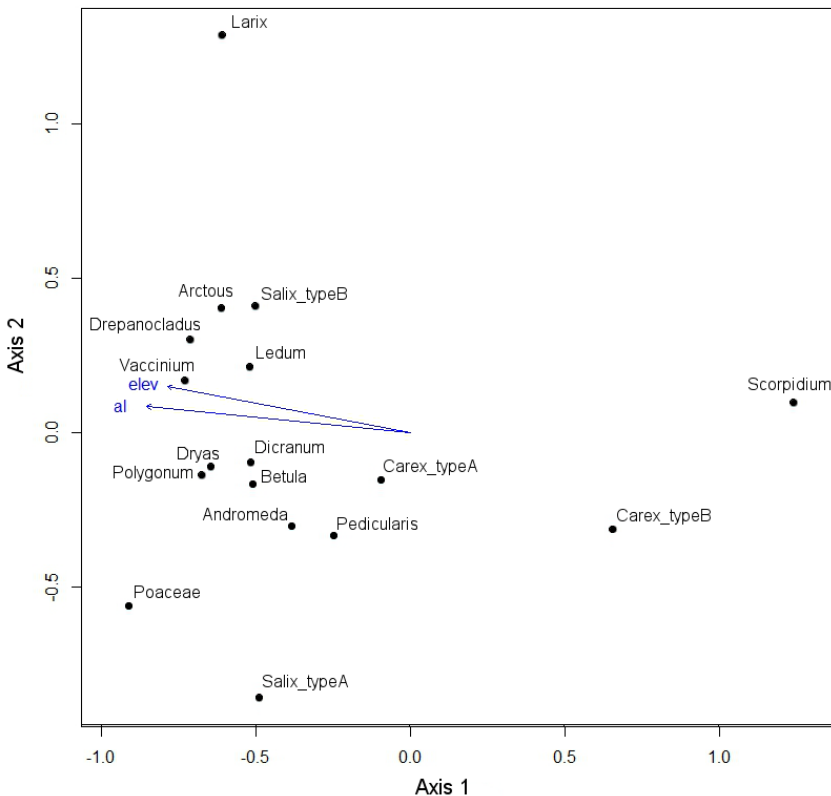
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**Fig. A1.** NMDS plot for the surface vegetation from the 07-SA-LY polygon. Axis 1 clearly shows the distribution of the vegetation along a water depth gradient. Thus *Scorpidium scorpioides* grows under water in the pond and comprises the main taxa on the rim, while the two *Carex* species are distributed between the pond and the rims. The overlaid environmental factors active layer (al) and elevation (elev) only correlate with taxa above the water surface because the water depth within the pond affects the vegetation. The distribution of the vegetation along Axis 2 is not clear from the available data set.

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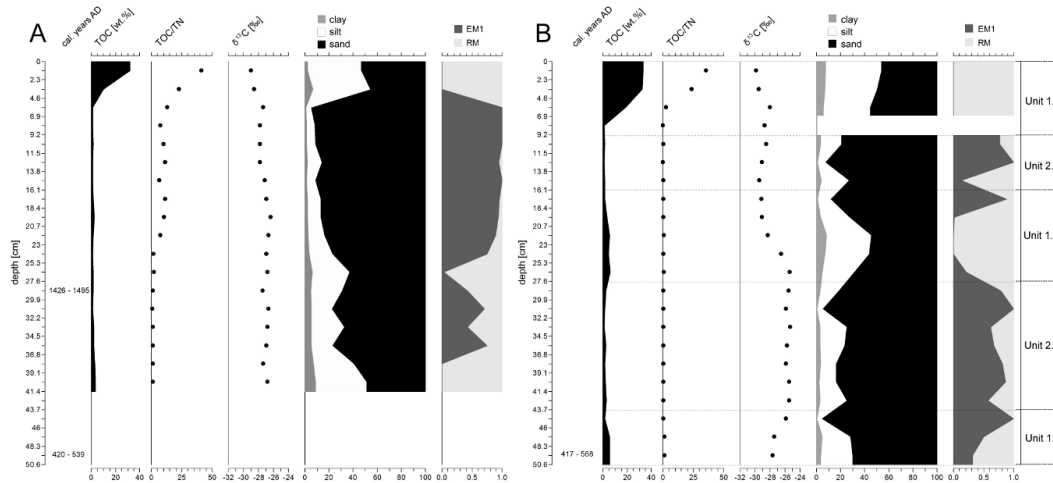
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**Fig. A2.** Summary plot of organic matter (TOC, TOC/TN) and granulometric characteristics (end-member scores and grain-size distribution) from sections A and B. The granulometric parameters for Section B 9.2 cm to 6.9 cm were not measured.

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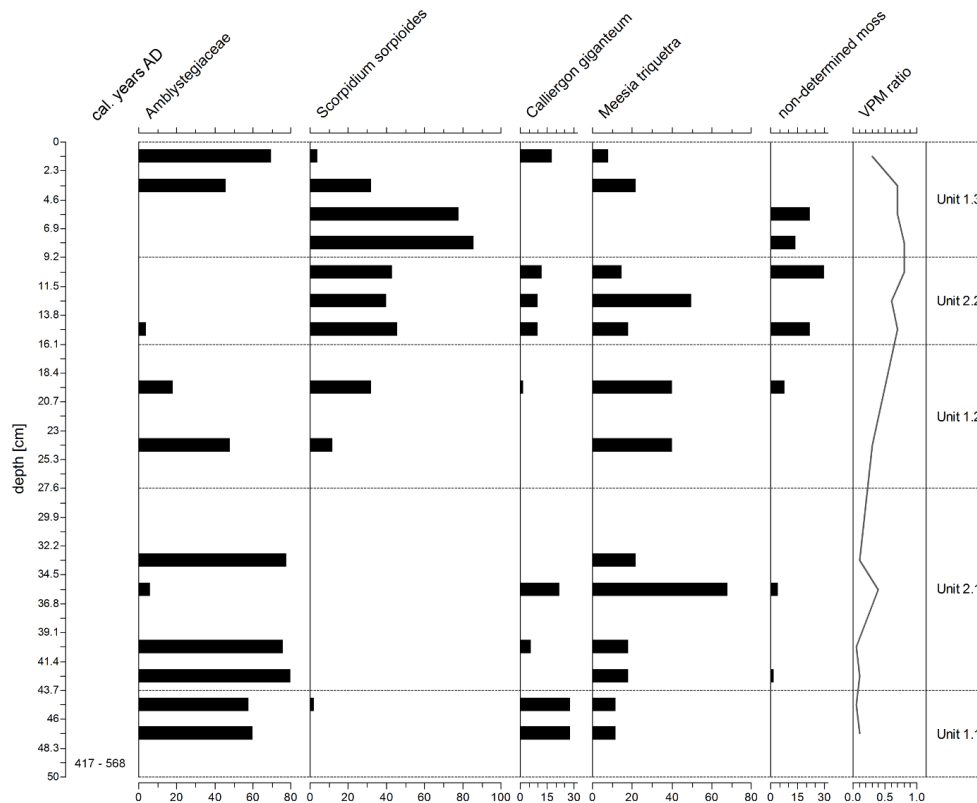
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**Fig. A3.** Moss spectrum for Section B. The figure does not include the complete moss spectrum as only several Amblystegiaceae taxa were analyzed in this section, especially *Scorpidium scopioides*, *Calliergon giganteum*, and *Meesia triquetra* (after Schilling 2012). The constant presence of *Meesia* represents transitional conditions with fluctuating water level.

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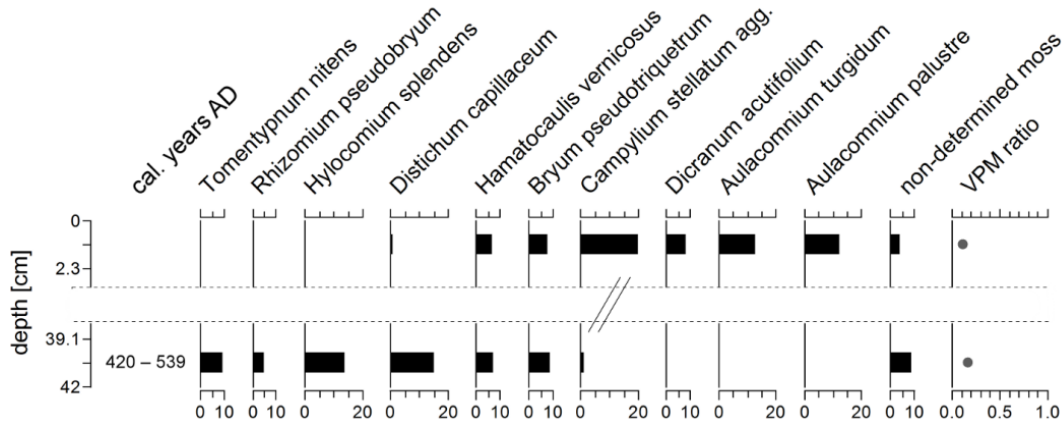
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**Fig. A4.** Moss spectrum for Section A. Only the surface and bottom samples from this section were analysed. An inversion was recognized from a typical polygonim moss community to a community with a preference for wetter conditions.

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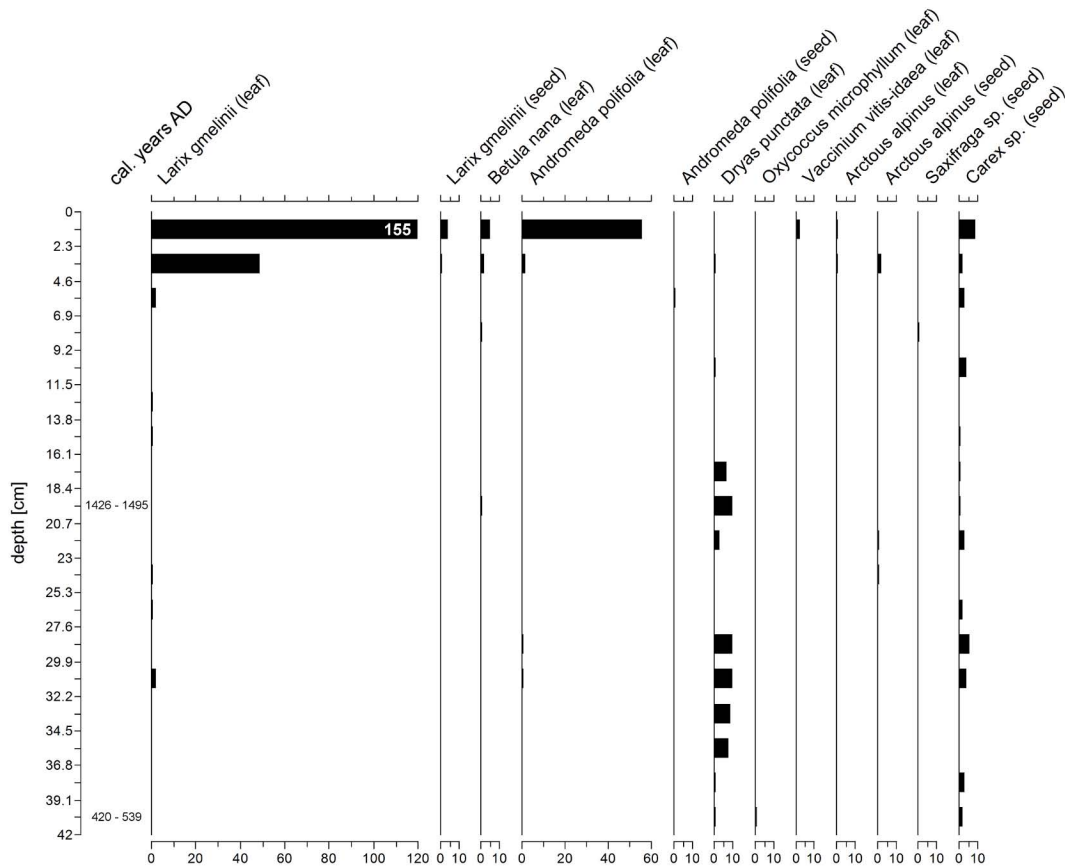
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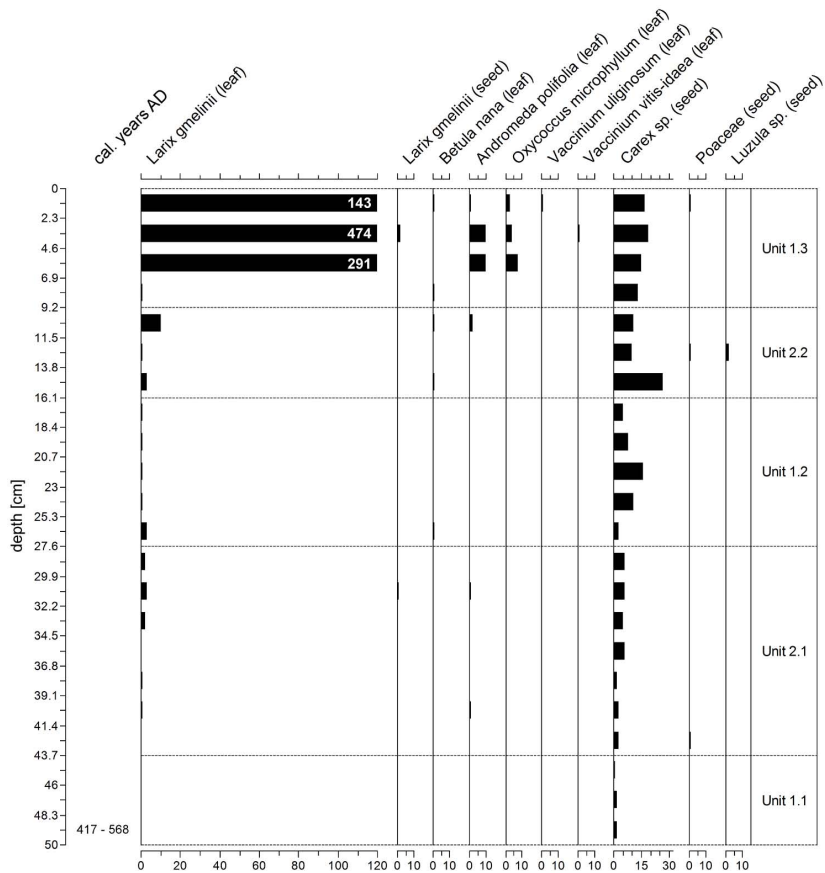
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**Fig. A5.** Results of the macrofossil analysis for Section A showing a dominance of *Carex* and *Dryas* in the lowermost samples with *Larix gmelinii* only becoming established in the uppermost samples from 4.8 cm to the surface.



**Fig. A6.** The macrofossil spectrum for Section B showing wetter vegetation with a dominance of *Carex* taxa and *Andromeda polifolia*. The peak of *Larix* leaves in the uppermost samples is due to the presence of trees in the vicinity of the Section B sampling location (see Fig. 3).

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