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Authors: Perren, Bianca B., Bradley, Raymond S., and Francus, Pierre

Source: Arctic, Antarctic, and Alpine Research, 35(3) : 271-278

Published By: Institute of Arctic and Alpine Research (INSTAAR), University of Colorado

Rapid Lacustrine Response to Recent High Arctic Warming: A Diatom Record from Sawtooth Lake, Ellesmere Island, Nunavut

Bianca B. Perren, Raymond S. Bradley, and Pierre Francus

Climate System Research Center, Department of Geosciences, University of Massachusetts, Amherst, MA 01003, U.S.A.
Current address: Bianca Perren
Department of Geology,
University of Toronto, 22 Russell St.,
Toronto, Ontario M5S 3B1, Canada.
perren@geology.utoronto.ca

Abstract

Diatoms from Sawtooth Lake (79°20'N, 81°51'W) on the Fosheim Peninsula in Central Ellesmere Island, Canada were analyzed to assess the temporal extent and magnitude of climatic change in the High Arctic during the late Holocene. Diatom results from the sediment cores show an absence of diatoms throughout the last ~2.5 ka (4.6 m) until the 1920s. However, ca. 1926 (5.3-cm depth), a rapid colonization of diatoms in the lake occurred. Within the uppermost section of the core (~1920 to ~1997), the diatom flora shift from a small Fragilaria-dominated assemblage to a more diverse assemblage that is dominated by large planktonic taxa (e.g., Cyclotella bodanica) and large raphid benthic species. The postglacial nature of this assemblage suggests a decrease in ice cover and a concomitant increase in light and nutrient availability for diatom growth over the last ~75 yr. Of particular significance is this absence of diatoms prior to the ~1920s, which indicates that environmental conditions of the last ~75 yr are unlike any of the previous ~2500 yr.

Introduction

Holocene climate fluctuations and recent warming have been reconstructed from a number of proxy climate data sets in the Northern Hemisphere. These reconstructions show pronounced warming in the last 50 yr, which is unprecedented in relation to the last 1000 yr (Mann et al., 1999). While these reconstructions provide much-needed high-resolution data for climate variability before the availability of instrumental data, it is necessary to push this application further back in time to expand our knowledge of natural climate variability throughout the Holocene, thereby placing the magnitude and rate of change in recent years within a longer perspective. This need is especially true in the High Arctic, where global circulation models (GCMs) predict an amplified response to greenhouse gas-induced warming (Zwiers, 2002), yet few long-term high-resolution climate data sets exist.

In the Arctic, recent environmental changes have been documented from many paleoclimatic records. These changes are manifested in the increased thickness of lacustrine varves (Smith, 1997; Hughen et al., 2000), in recent shifts in fossil algal assemblages preserved in lacustrine sediments (Douglas et al., 1994; Doubleday et al., 1995; Gajewski et al., 1997; Wolfe, 2000; Wolfe and Perren, 2001; Sorvari et al., 2002), and from other high-resolution paleoclimatic proxy records (reviewed in Overpeck et al., 1997). These changes are thought to represent recent warming in the Arctic that is unprecedented in the context of recent centuries. The warming is consistent with GCM predictions of climate expected under higher CO2 levels, suggesting that it is, at least in part, the result of anthropogenic influences on global climate.

In this paper, we present a late Holocene paleolimnological record from the sediments of Sawtooth Lake, Ellesmere Island, Nunavut, Canada. The purpose of this study is to determine the nature of recent environmental change from the biological record contained within the sediments by a detailed, high-resolution analysis of the fossil diatoms.

ECOLOGICAL RESPONSE TO CLIMATE CHANGE IN THE HIGH ARCTIC

The development of calibration sets for circumarctic sites (Pienitz et al., 1995; Lim et al., 2001; Weckström and Korhola, 2001; Joynt and Wolfe, 2001) is rapidly changing the applicability of quantitative models to both high and low arctic sites; however, because of limited autecological information in the High Arctic, hindcasting climatic trends using diatoms has relied historically on qualitative models of their inferred responses to climatic change (Smol, 1983). Because high arctic lacustrine environments are dominated by low temperatures, thick and persistent ice cover, limited light, and a short (often less than 8 wk) growing season, a reduction in ice cover is expected to be the major response of arctic lakes to climatic warming (Douglas and Smol, 1999). A reduction in ice cover has many implications for the aquatic ecology of arctic lakes (Rouse et al., 1997; Douglas and Smol, 1999, Battarbee, 2000). It promotes an increased diversity of habitats for autotrophic species, such as diatoms, by increasing light in the euphotic zone for plankton growth, and allows the littoral zone to be colonized by mosses and thus epiphytic species. A longer growing season allows these species to establish themselves and reproduce for longer periods of time, thereby increasing productivity and ecological complexity. In addition, warmer climates accelerate weathering in the watershed. This increased buffering releases more nutrients and dissolved solids into the lake, which augment primary productivity (Douglas and Smol, 1999).

ICE COVER MODEL

The main model used in the Arctic relates diatom species assemblages to changes in ice cover (Smol, 1983). As warming occurs, the ice pan on the lake shrinks, thereby allowing more photosynthetically active radiation to penetrate the lake. The moat (the area between the shore and the ice pan) becomes wider, allowing for the proliferation of deeper water and epiphytic taxa. Convective circulation in the ice-free areas and increased nutrients allow for larger phytoplankton to thrive in the euphotic zone and for larger epiphytic taxa to colonize the littoral zone of the lake. Thus, with warming, a shift from small shallow-water taxa (e.g., Fragilaria pinnata) to larger epiphytic benthic species (e.g., Navicula spp.) and planktonic taxa (e.g., Cyclotella spp., Stephanodiscus spp.) should be apparent (Lotter and Bigler, 2000). Indeed, this species transition is also a noted sequence in the glacial/postglacial ontogeny of lakes at lower latitudes (reviewed in Round, 1981).

While this model provides the means by which diatom records can be interpreted, it cannot be directly tied to any single meteorological
variable, which is necessary for the use of a transfer function. Ice cover does relate to summer warmth (Doran et al., 1996; Schindler et al., 1996), but not linearly. Several factors such as winter temperature, windiness, and internal lacustrine thermal feedbacks complicate the relationship (Doran et al., 1996), as do changes in snow depth, which affect sub-ice circulation, diatom suspension, and photosynthesis (Kelley, 1997). In addition, caution must be used in ascribing changes solely to ice cover. A reduction in ice cover is merely one factor among others, such as the increases in nutrient supply, light, and habitat availability that result from climatic warming.

Study Area

Sawtooth Lake (79°20′N, 83°51′W, 280 m a.s.l.) is the highest of a chain of three lakes situated within a col in the Sawtooth Mountain Range on the Fosheim Peninsula in central Ellesmere Island (Fig. 1). The surrounding unglacierized landscape is considerably high in relief, with the steep-sided, east-west-trending valley walls reaching 1000 m in elevation. It is a large (260 ha), deep (>90 m) lake that retains considerable ice cover during the summer months and is rarely ice free. The prevalence of lake ice sharply contrasts with the relatively warm climatic and vegetational characteristics of this intermontane region. Summer temperatures (June, July, and August) average 5.5°C in nearby Eureka (60 km to the north). Mean annual temperature averages −20°C. The area is an extreme polar desert where annual precipitation, which falls mainly as snow, averages 68 mm, making it the driest place in Canada (Environment Canada, 2003). The catchment is largely unvegetated: the silt and clay-rich lowlands support an enriched prostrate shrub flora (Edlund and Alt, 1989), whereas the felsenmeer highlands, predominantly composed of the highly erodable carbonate bedrock of the Eureka Sound Group (Miall, 1991), support only a sparse saxifrage-dominated community.

The lake is situated well above Holocene marine limit (<140 m a.s.l.; Bell, 1996) and is slightly alkaline (pH 8) and dilute with concentrations of major cations ranging from ~0–4.5 mg L⁻¹ for Na⁺, K⁺, Mg²⁺, and Si⁴⁺, with higher concentrations of Ca²⁺ (~20 mg L⁻¹) derived from the surrounding bedrock. Lake water temperatures range between 0 and 2°C, and oxygen levels range between over 15 mg L⁻¹ at the surface to 4 mg L⁻¹ at depth in the distal basin (see Table 1).

The mixing regimen of Sawtooth Lake is complicated. The predominance of ice on the lake prevents classic cold monomictic overturn. However, the inflowing streams from the eastern end of the watershed deliver considerable sediment to the lake in the form of overflows, interflows, and underflows (Patridge, 1999) and effectively mix the lake. A sill (60-m depth) in the middle of the lake acts as a barrier to communication between the two basins and creates an
isolated trough in the distal basin. This trough is depleted in oxygen, has relatively higher concentrations of hypolimnetic cations, and allows for the preservation of laminae from which a high-resolution record may be obtained.

**Methods**

**CORING AND DIATOM PREPARATION**

A suite of gravity and vibracores was collected from the distal basin of Sawtooth Lake in the summer of 1999. Diatoms were sampled from the 36-cm-long gravity core (10–5) at 1–2-mm intervals to approximate an annual record in the uppermost 6 cm and at 1-cm intervals thereafter to 36 cm. The vibracore (10–6B) was sampled at 10-cm intervals to 4 m; thereafter it was sampled at 5-cm intervals to the base at 4.6 m.

Diatom samples from the short core were freeze-dried and then digested in 30% hydrogen peroxide following standard oxidative techniques (Battarbee, 1986). They were repeatedly rinsed, and then aliquots were evaporated at room temperature on round coverslips in evaporation trays. Coverslips were permanently mounted on plain glass slides using Naphrax® mounting medium (RI = 1.74). At least 300 diatoms were enumerated from the uppermost samples using an Olympus HB microscope equipped with phase optics. Where diatom concentrations declined, at least 5 transects of the slide were counted. For several of these (below 5-cm depth), total diatom counts were less than 5. Diatom species are expressed as relative percentages and as estimated concentrations within the sediment based upon the known weight of the original sample, the volume of the subsequent aliquots taken, and the area of the slide surface counted (Battarbee, 1973). Taxonomic assignations were made using reference literature (Krammer and Lange-Bertalot, 1986–1991).

**CHRONOLOGY**

The chronology of Sawtooth Lake is complicated. An additional surface lamina recovered in each successive year of coring suggests that the sediments are annually laminated (varved). However, a large erosive turbidite at 2-cm depth and several sections where laminae are disturbed preclude a definitive varve chronology for the uppermost section. Two trends in the diatom stratigraphy are apparent over the last 80 yr. In some sections, the diatom assemblages, as we will discuss further, can be constrained within a decade. Diatom samples were taken from individual laminae in an effort to approximate an annual record and were assigned ages based on linear interpolation between 210Pb-generated dates.

**STATISTICAL METHODS**

Diatom species >1% abundance in any one sample were employed in statistical analyses. Gradient lengths were assessed using detrended correspondence analysis (DCA) on CANOCO software (ter Braak, 1990). Principle Components Analysis (PCA) was chosen for further analysis due to the linear response of the data, indicated by the short gradient length (1.896) of the DCA 1 axis. The PCA was run using a centered covariance data matrix and then plotted as both a species scatter (Fig. 4) and as downcore first-axis scores (Fig. 5).

**BIOVOLUME**

Diatom biovolume (μm³ g⁻¹ dry sediment) was estimated for diatom species using the BIOVOL program (Kirschelt, 1996). Size ranges were estimated from reference photographs from Sawtooth Lake and other girdle views from other high arctic lakes (usually smaller than temperate-cell size ranges). In cases where girdle views were unavailable, volumes were estimated from similarly sized species. Due to the inherent uncertainty in estimating cell volumes, the values are used to provide a rough estimate of changes in diatom production over time rather than absolute numbers. Despite these uncertainties, volumes calculated here were comparable to other, published diatom biovolumes (compare Joynt and Wolfe, 2001).

**Results**

**GENERAL DIATOM STRATIGRAPHY**

Diatoms are absent from the entire retrieved sedimentary record of Sawtooth Lake below 5.3-cm depth (~1926). However, in the uppermost 5.3 cm, diatom concentrations increase rapidly and oscillate between 0 and 3.5 x 10³ valves g⁻¹ dry sediment. Of a total of 66 species seen in the lake sediments, only 21 diatom species were recorded in >2% relative abundance (Table 2). The siliceous remains of chrysophytes are equally variable and appear periodically throughout the core, but are most abundant at the top, where they range from 0 to 1 x 10⁷ cysts g⁻¹ dry sediment (Fig. 2) and below 4-m depth, where they were inconsistently observed.

A trend in the diatom stratigraphy is apparent over the last 80 yr (Fig. 3). In general, diatoms increase in both size and diversity toward the surface: from the early dominance of small Fragilaria species to larger planktonic Cyclotella species and larger benthic taxa (Navicula, Cymbella, etc.) in recent decades. This general trend is superimposed upon the higher-frequency oscillations in species dominance, which punctuate the record.

When diatoms first appear in the lake ca. 1926, it is as a low concentration and low biovolume population of Fragilaria pinnata. Fragilaria species diversify quickly and are replaced by larger taxa, both benthic and planktonic, which reach a maximum circa 1980. In...
The early 1990s, concomitant with a large turbidite, diatom species diversity and concentration within the sediments plummet. The mid-to late 1990s are marked by a return to high species diversity and concentration.

ORDINATION

The PCA of diatom species highlights trends within the data (Fig. 4). Negative values on the first axis ($\lambda = 64\%$) represent "warmer"-condition taxa: large, high-biomass, centric diatoms (Cyclotella bodanica and Stephanodiscus alpinus) and larger benthic, epiphytic species belonging to the genera Achnanthes, Cocconeis, Cymbella, Denticula Diploneis, Navicula, Pinnularia, and Nitzschia. The positive, "colder" side of the axis is dominated by the presence of small Fragilaria pinnata (and the occasional Campylodiscus noricus), which is typically associated with late glacial lakes at more temperate latitudes (Haworth, 1976; Stabell, 1985; Marciniak, 1986). The second axis ($\lambda = 17\%$) shows intermediate species: other Fragilaria and small benthic taxa (e.g., Amphora pediculus, Caloneis spp.). These groupings can therefore be seen as a gradient from "cold" to "warm." The downcore diatoms appear to follow this "ice-cover" gradient with the transition from cold, ice-covered conditions to warmer conditions in the last few decades (see Fig. 5). Indeed, this trend echoes the climate trends of the region where 6 of the warmest years on record were in the last decade (Environment Canada, 2003).

BIOMASS

Two main features are apparent in the diatom stratigraphy of Sawtooth Lake. Years with low diatom concentrations are dominated by the small diatom Fragilaria pinnata. Those years with high diatom concentrations have higher diatom diversity and a larger relative abundance of large diatom taxa. These features are evident in the stratigraphic changes in diatom biovolume (see the right column on Fig. 3). Cyclotella bodanica is roughly 10–15 times the volume of small Fragilaria pinnata and thus requires considerably more resources for proliferation. This change in biomass, from a few small diatoms to a lot of larger diatoms, illustrates the degree to which the diatom production in Sawtooth Lake has exponentially increased within the last 100 yr.

**TABLE 2**

<table>
<thead>
<tr>
<th>Diatom species &gt;2% abundance in any sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Achnanthes marginulata Grun. in Cleve &amp; Grun. 1880</td>
</tr>
<tr>
<td>Achnanthes minutissima Kutz. 1833</td>
</tr>
<tr>
<td>Amphora maritima Krammer</td>
</tr>
<tr>
<td>Amphora hyalina Ehrl.</td>
</tr>
<tr>
<td>Amphora pediculus (Kutz.) Grun.</td>
</tr>
<tr>
<td>Campylodiscus noricus Ehrenb. ex Kutz. 1844</td>
</tr>
<tr>
<td>Cyclotella bodanica var. lemanica (O. Muller ex Schroter) Bachmann 1903</td>
</tr>
<tr>
<td>Cyclotella comensis Grun. in Van Heurck 1882</td>
</tr>
<tr>
<td>Cyclotella pseudostelligera Hust. 1939</td>
</tr>
<tr>
<td>Denticula tenuis Kutz. 1844</td>
</tr>
<tr>
<td>Diploneis cf. parva Cleve 1891</td>
</tr>
<tr>
<td>Fragilaria capucina var. vaucheri (Kutz.) Lange-Bertalot 1980</td>
</tr>
<tr>
<td>Fragilaria construens var. venter (Ehrenb.) Grun. in Van Heurck 1881</td>
</tr>
<tr>
<td>Fragilaria pinnata var. intercedens (Grun.) Hust. 1931</td>
</tr>
<tr>
<td>Fragilaria pinnata var. pinnata Ehrenb. 1843</td>
</tr>
<tr>
<td>Fragilaria pseudococonus Marciniak. 1982</td>
</tr>
<tr>
<td>Hansaea arcus (Ehrenb.) Patr. in Patr. and Reimer 1966</td>
</tr>
<tr>
<td>Navicula cincta (Ehrenb.) Rafls in Pritch. 1861</td>
</tr>
<tr>
<td>Navicula cryptocephala Kutz. 1844</td>
</tr>
<tr>
<td>Nitzschia perminuta (Grun.) M. Perag. 1903</td>
</tr>
<tr>
<td>Stephanodiscus alpinus Hust.</td>
</tr>
</tbody>
</table>

**FIGURE 2.** Concentration of (a) diatoms and (b) chrysophyte cysts within core 10–8. Diatoms are absent below 6 cm. The gray band marks the presence of a turbidite.

**Discussion**

Due to the lack of pitting or etching on the valve surfaces, repetitions in laboratory preparations of diatoms, and the oscillations in diatom concentration at the top of the core, the sedimentary sequence is interpreted as recording an environmental rather than taphonomic signal. First-order changes in both the relative abundance and concentration of diatoms suggest that major changes have occurred within the lake environment over the last 100 yr.

Furthermore, the record of annual median grain size at Sawtooth Lake, interpreted as the early-summer snowmelt intensity (Francus et al., 2002) shows a clear coarsening in the 20th century, from ~1900 until the 1960s. Unlike the record for diatoms, the coarsening recorded is not unprecedented in the last 400 yr. Nevertheless, both of these independent proxies, i.e., grain size and diatoms, from the same site clearly demonstrate a response to warming during the latter half of the 20th century.

**LAKE ONTOGENY**

The change in diatoms reflects the ontogeny of Sawtooth Lake and the evolution of diatom production within it. The small Fragilaria pinnata is considered a pioneering species, commonly found in late glacial sediments (Haworth, 1976; Stabell, 1985; Marciniak, 1986), whose adaptation to marginal environments allows it to outcompete larger species in marginal environments (Lotter et al., 1999; Lotter and Bigler, 2000). The autecology of other Fragilaria species is less well known. However, the diatom record from nearby Solstice Lake (Wolfe, 2000) suggests that F. pseudococonus may be viewed as an intermediate species on a gradient between "glacial" conditions represented by F. pinnata and interglacial conditions marked by larger diatom taxa in high arctic environments.

Indeed, the changes seen over the last 80 yr in Sawtooth Lake are similar in character to changes seen at lower latitudes in the transition
FIGURE 3. Percentage diagram for Sawtooth Lake diatoms and their concentration within the sediments. Sum of other benthic taxa includes Achnanthes, Denticula, Diploneis, Hannaea, Navicula, and Nitzschia as well as other less well-represented genera such as Caloneis, Cocconeis, Cymbella, Diatoma, and Pinnularia. (See Table 2 for taxa list.) The gray band marks the presence of a turbidite.

FIGURE 4. PCA biplot of species indicating “cold” and “warm” taxa.
REGIONAL DIATOM/LACUSTRINE CHANGE

The paleoclimate records from fossil diatoms across the Canadian High Arctic reveal strikingly similar changes in recent decades (Table 3). In general, all of these records depict a distinct transition from small Fragilaria species, which are indicative of low-productivity environments, to larger benthic, periphytic species. While the nature of these transitions is the same, the timing of the shift varies. Several records show the changes occurring in the late 1800s (Douglas et al., 1994; Wolfe, 2000), whereas others show it happening sometime within the last 50 yr (Gajewski et al., 1997; Doubleday et al., 1995).

Three diatom records from Cape Herschel on Ellesmere Island (Douglas et al., 1994) show a distinct transition from small Fragilaria species to larger epiphytic species (Pinnularia, Cymbella, Navicula) in the uppermost centimeters of the sediment cores. These records indicate that changes that have occurred in the last ~150 yr (dated by $^{210}$Pb) are unprecedented within the context of the last 6000–8000 yr ($^{14}$C dating). Similar records are found elsewhere in the Arctic. The record from Solstice Lake (Wolfe, 2000), a small lake on the Fosheim Peninsula, Ellesmere Island, shows the same timing of the change from small Fragilaria species to larger benthic taxa as the Cape Herschel records. The recent changes in diatoms in these two records show that the magnitude of recent warming since the Industrial Age is unprecedented in the context of the middle to late Holocene.

The record from DV09, a small lake with laminated sediments on northern Devon Island (Gajewski et al., 1997), shows that these changes happened in the last 60 yr. Staurosirella (=Fragilaria pinnata group) and other small taxa gave way to the dominance of larger raphid taxa in recent years. In addition, the total diatom concentration has increased dramatically since 1948. These changes are paralleled by an increase in summer melting on the Devon Ice Cap, as revealed by ice-core studies (Fisher and Koerner, 1994).

The record from Lower Dumbell Lake (Doubleday et al., 1995), near Alert military base on the north coast of Ellesmere Island, shows the same type of record and timing of change. Diatoms are absent below 1.5 cm in the core. In the uppermost 1 cm, the fossil assemblage shifts from small Fragilaria pinnata to larger epiphytic and planktonic taxa (Cyclotella spp.). Due to the thick ice cover in this northern locale, the change is interpreted to represent a reduction in ice cover over the last ~50 yr, with the chronology based on the coincident presence of black carbon spherules (combustion products from nearby Alert, which was established in 1950).

These records all suggest that diatoms in high arctic lakes and ponds, regardless of lake size, latitude, or altitude (Table 3), have most likely responded to a decline in ice cover and a concomitant increase in light availability, growing season length, epiphytic habitats, and increased nutrient supply, all of which have likely occurred as a result of recent (last ~100 yr) warming. Similar records from across the circumpolar region depict the same types of ecological change (Sorvari et al., 2002; Korhola et al., 2002). This inferred warming appears to be unique within the context of the late Holocene. Unfortunately, the resolution of these records and the resulting discrepancy in the timing of the shifts preclude any detailed, decadal, regional synthesis. The temporal inconsistency probably relates to differences in sampling and dating resolution, levels of bioturbation, and the size of the lakes, which affect their threshold of sensitivity to climatic perturbations.

**TABLE 3**

<table>
<thead>
<tr>
<th>Lake</th>
<th>Latitude</th>
<th>Longitude</th>
<th>Altitude</th>
<th>Depth</th>
<th>Length</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sawtooth L.</td>
<td>79°20’N</td>
<td>81°51’W</td>
<td>280 m</td>
<td>&gt;90 m</td>
<td>4000 m</td>
<td>this study</td>
</tr>
<tr>
<td>DV09</td>
<td>75°34’N</td>
<td>89°18’W</td>
<td>35 m</td>
<td>16 m</td>
<td>300 m</td>
<td>Gajewski et al., 1997</td>
</tr>
<tr>
<td>Col Pond</td>
<td>78°37’N</td>
<td>74°42’W</td>
<td>135 m</td>
<td>ephemeral</td>
<td>&gt;100 m</td>
<td>Douglas et al., 1994</td>
</tr>
<tr>
<td>Camp Pond</td>
<td>78°37’N</td>
<td>74°42’W</td>
<td>58 m</td>
<td>&lt;0.8 m</td>
<td>&gt;100 m</td>
<td>Douglas et al., 1994</td>
</tr>
<tr>
<td>Elison L.</td>
<td>78°37’N</td>
<td>74°42’W</td>
<td>17 m</td>
<td>&gt;1.5 m</td>
<td>~500 m</td>
<td>Douglas et al., 1994</td>
</tr>
<tr>
<td>Dumbell L.</td>
<td>82°29’N</td>
<td>62°29’W</td>
<td>83 m</td>
<td>&gt;30 m</td>
<td>1000 m</td>
<td>Doubleday et al., 1995</td>
</tr>
<tr>
<td>Solstice L.</td>
<td>79°25’N</td>
<td>84°07’W</td>
<td>305 m</td>
<td>6 m</td>
<td>600 m</td>
<td>Wolfe, 2000</td>
</tr>
</tbody>
</table>

**FIGURE 5.** PCA axis 1 scores showing inferred cold-warm transition in recent decades.
Lake highlights the magnitude of recent changes in the arctic environment. The mass balance of circum-arctic glaciers and recent climate change: evidence from small glaciers. Quaternary Research, 48: 1–14.


