Siliceous microfossil succession in the recent history of two basins in Lake Baikal, Siberia

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Abstract

As part of the international cooperative Baikal Drilling Project, siliceous microfossil assemblage succession was analyzed in two short (~ 30-cm) sediment cores from Lake Baikal. One core was recovered from the north basin (Core 324, 55°15'N, 109°30'E), a second from between the central and southern basins (Core 316, 52°28'N, 106°50'E). The northern core had higher amounts of biogenic silica (40 g SiO2 per 100 g dry weight sediment) compared to the southern core, and increased deposition in the more recent sediments. Weight percent biogenic silica was lower in the southern core, ranging from approximately 20-30 g SiO2 per 100 g dry weight sediment throughout the entire core. Trends in absolute microfossil abundance mirror those of biogenic silica, with generally greater abundance in the northern core (86-275 x 106 microfossils g⁻¹ dry sediment) compared to the southern core (94-163 x 106 microfossils g⁻¹ dry sediment).

Cluster analyses using relative abundance of the dominant diatom and chrysophyte taxa revealed four zones of microfossil succession in each core. Microfossil assemblage succession in the north basin may be reflecting shifts in nutrient supply and cycling driven by climatic changes. The most recent sediments in the northern basin (Zone 1, c. 1890’s–1991 A.D.) were characterized by an increased abundance of Aulacoseira baicalensis and Aulacoseira ‘spore’. Zone 3 (c. 1630’s–1830’s A.D.) was dominated by the endemic Cyclotella spp. and reduced abundance of the Aulacoseira spp. Zone 3 corresponds approximately to the Little Ice Age, a cooler climatic period. The microfossil assemblages between Zones 1 and 3 (Zone 2, c. 1830’s–1890’s A.D.) and below Zone 3 (Zone 4, c. 830’s–1430’s A.D.) are similar to one another suggesting they represent transitional intervals between warm and cold periods. Southern basin sediments record similar changes in the endemic taxa. However, the increased abundance of non-endemic planktonic taxa (e.g. Stephanodiscus binderanus, Synedra acus, Cyclostephanos dubius) during two periods in recent history (post World War II and late 1700’s) suggests evidence for anthropogenic induced changes in southern Lake Baikal.

Introduction

Lake Baikal is the largest (23 000 km²), deepest, and perhaps the oldest lake on Earth. Beneath its greater than 1.6 km depth, lies over 7 km of sediment, preserving a 25 million year Neogene history. The lake’s age, large size, and extremely high degree of organismal endemism has attracted much scientific interest (Kozhov, 1963) and Baikal’s great depths and geographic locale produce unique limnological conditions (Weiss et al., 1991; Shimaraev et al., 1993; Bradbury et al., 1994).

Lake Baikal represents a unique and immense natural resource for Russia. Unfortunately, urban and industrial development, especially surrounding the southern basin, have placed added burden on the Baikal system (Belt, 1992). Maatela et al. (1990) revealed impacts on Baikal’s benthic environment near southern industrial localities from effluent loadings. Galazii (1991) summarized some recent findings surrounding
environmental impacts on Baikal, ranging from local extirpations of endemics to biodiversity loss. Fortunately, there appears to be increased awareness at local, national, and global levels surrounding the future use, protection, and understanding of Lake Baikal.

One sign of this new awareness surrounding Lake Baikal has been the creation of a cooperative group of international scientists to form the Baikal Drilling Project. This interdisciplinary group was formed to study the immense Baikal sediment record, with goals of reconstructing the geological evolution of the Lake Baikal depression and resolving a detailed, quantitative Quaternary climatic record (Lake Baikal Paleoclimate Project Members, 1992; Kuzmin et al., 1993).

The 7.5 km Baikal sediment record is very suitable for paleoclimatic reconstruction. An apparently continuous sedimentary record has been deposited since the Miocene and glaciations in the basin have been limited high and Asiatic low (Lydolph, 1977; Bradbury et al., 1994). Lake Baikal’s high latitude location makes it sensitive to long term insolation patterns reflecting changes in the Earth’s orbital parameters (Kuzmin et al., 1993) and more short-term continental climate changes resulting from seasonal patterns associated with the shifts in the dominant Siberian high and Asiatic low (Lydolph, 1977; Bradbury et al., 1994). Changes in climatic conditions have affected Lake Baikal’s limnological patterns, particularly ice thickness and duration, mixing depth, nutrient loading and recirculation, which all subsequently affect the timing and levels of algal production.

Paleolimnological methods hold strong promise for resolving climatic changes in Baikal’s past. Bradbury & Dieterich-Rurup (1993) analyzed the varved sediment record in north temperate Elk Lake (Minnesota, USA) and reported 100 to 1000 year climatic patterns during the past 10 400 years. Even recent climatic events, such as the Little Ice Age (c. 1450–1850 A.D.), possibly characterized by cooler conditions and late springs, were resolved using the diatom record.

Earlier paleolimnological studies on Baikal have identified dramatic changes in the diatom communities over time. Chernyaeva (1970) studied diatom remains in a transect of seven cores across Baikal’s north basin. While high resolution sampling was not employed, Chernyaeva showed that, in addition to high benthic diversity and well preserved remains, past changes had occurred in just the upper meter of sediment. These changes ranged from extirpation of some taxa, to taxonomic shifts in quantitative abundance. Bradbury et al. (1994) compiled data from cores (up to 11 m deep) taken throughout the Baikal basin and demonstrated the utility of Baikal sediments for resolving climate change. One core (305) provided an especially detailed record of the post-glacial Holocene (15.0–1.3 ka). Bradbury et al. (1994) suggested that shifts in taxonomic composition of the microfossil assemblage reflected climatic shifts between cold-dry and warm-moist periods, as well as changes in nutrient dynamics associated with loadings and circulation. Unfortunately, both of these previous studies (Chernyaeva, 1970; Bradbury et al., 1994) had poor resolution of nearsurface changes in sedimentary signals because of either wide sampling intervals or loss of surficial sediments during coring.

While earlier paleolimnologic studies have not focused specifically on anthropogenic effects, there have been changes noted in Lake Baikal’s primary producer communities over the last 50 years (Bondarenko et al., 1993; Popovskaya, 1993; Flower, 1993) possibly associated with human impacts (Galazii, 1991). Changes include, most notably, a significant increase in pennate diatom contribution to the plankton by Nitzschia acicularis W. Sm. and Synedra acus Kütz. since the 1950’s, a decrease in relative abundance of some endemic taxa (e.g. Cyclotella baicalensis Skv. and C. ornata (Skv.) Flower), and recent appearance of nonendemic taxa. Changes of this nature often indicate changes in nutrient loading and a potential for increased productivity (Stoermer et al., 1993). However, while there has been increased nutrient loading to Baikal (Galazii, 1991), correlations to productivity increases have been difficult to make. Overall, increased productivity has not occurred, at least not at levels resolvable from annual variation. The system’s plankton response has been through increased numerical abundance, especially of the small-celled species (Popovskaya, 1993). Similar changes were also evident from a survey of surficial sediments in Lake Baikal (Stoermer et al., 1995). Increased abundance of nonendemic centric taxa and greater numbers of planktonic pennates characterized the southern basin. Northern basin surficial sediments, on the other hand, were still dominated by the endemic Aulacoseira and Cyclotella taxa.

Records of modern changes in Baikal phytoplankton assemblages are incomplete and difficult to interpret. Dramatic seasonal and, more importantly, annual differences in phytoplankton composition and abundance (Kozhov, 1963; Popovskaya, 1991) make interpretations and predictions of annual production difficult. However, the sedimentary record preserved in
the Baikal basin can provide answers to questions concerning both the recent and long term history of changes in this basin. In this study, we report on recent (<1100 yr) siliceous microfossil assemblage succession in two short (~30-cm) sediment cores from Lake Baikal. Specific objectives of this project were to analyze the extent of anthropogenic modification in recent history of two basins in Lake Baikal and to determine if any climatic signals were evident in recent sedimentation.

Materials and methods

Box cores were recovered from two sites in Lake Baikal during summer 1991 as part of the Baikal Drilling Project (Lake Baikal Paleoclimate Project Members, 1992). Core 324 is from Baikal's north basin (55° 15' N, 109° 30' E) and Core 316 is from between the central and southern basins (52° 28' N, 106° 05' E) west of the Selenga River delta (Fig. 1). Subsampled cores from each box were sectioned by extrusion in ten 0.5-cm intervals from 0 to 5.0 cm, and select 1.0-cm intervals were taken at 6–7, 8–9, 14–15, 20–21, 26–27, 28–29 (Core 324 only), and 32–33 cm (Core 316 only). Core sections were refrigerated for further processing.

Weight percent sedimentary biogenic silica was analyzed spectrophotometrically with the molybdate blue method (Pilskaln & Paduan, 1992). Extraction methods using Na2CO3 were adapted from Leinen (1977), Eggimen et al. (1980) and Mortlock & Froelich (1989).

For siliceous microfossil preparation, a split from each section was freeze-dried to reduce microfossil breakage inherent with air or heat drying. A weighed dry subsample, between 11 and 13 mg, was boiled for 30 min in 30% H2O2 at 110 °C. After boiling, 25 ml of concentrated HNO3 was added to the peroxide-sediment solution. This resulted in a vigorous exothermic reaction within 5 min, after which the solution was further heated at 120 °C for an hour. Byproducts of oxidation were removed with six rinses of distilled water, allowing a minimum of six hours of settling between rinses. The entire sample was then poured into Battarbee drying chambers (Battarbee, 1973), allowed to evaporate to dryness, and the four replicate coverslips mounted in Hyrax™.

A minimum of five hundred microfossils were enumerated on a single transect on each of two replicate coverslips using brightfield oil immersion optics (N.A. > 1.32) capable of 1200 × magnification. Diatoms were identified to the lowest taxonomic designation and fragments of diatoms mathematically reconstructed to whole valve units. Chrysophyte cyst morphotypes were given a numerical identifier when first encountered (e.g., Figs 28–33). Sponge spicules were rare but also counted. Raw counts were converted to absolute abundance and reported as diatom valves or microfossils per g dry sediment. Counts were also converted to percent abundance and reported relative to total microfossil abundance.

To analyze the relationships among microfossil assemblages within the two cores, cluster analyses were performed on the most abundant taxa (Stoermer et al., 1985a), defined as those found in core sections with five occurrences at > 1% relative abundance (Tables 1 & 2). Discussion of the sedimentary distribution of taxa will be limited to these species. Cluster analyses were run with SYSTAT (Wilkinson, 1989) and were based on Euclidean distance measure and average distance clustering (Carney, 1982).

The limited subsample size available for our analyses precluded any dating estimates to be made directly on material from box cores 316 and 324. In lieu of direct measurements, we estimated dates for each core using linear sedimentation rates calculated by Edgington et al. (1991). Edgington et al. (1991) collected short cores from throughout the Baikal basin in 1988, analyzed 210Pb and 137Cs inventories, and calculated linear and mass sedimentation rates in each of Baikal's three basins and near the Selenga River delta. Two of the Edgington et al. (1991) stations (3 and 4) were near our Core 316 site and their two cores had mean sedimentation rates (including both with and without

<table>
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<tr>
<th>Table 1. Diatom and chrysophyte taxa used in cluster analysis, Core 324. Taxa chosen were present in five or more sample intervals at &gt; 1% relative abundance</th>
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<td><em>Aulacoseira baikalensis</em> (C. Meyer) Simonsen</td>
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<td><em>Aulacoseira baikalensis</em> fo. <em>square-punctae</em></td>
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<td><em>Aulacoseira 'spore'</em> (Stoermer et al., 1995)</td>
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<td><em>Cyclotella baikalensis</em> Skv.</td>
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<td><em>Cyclotella ornata</em> (Skv.) Flower</td>
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<td><em>Cyclotella minuta</em> (Skv.) Antipova</td>
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<td><em>Stephanodiscus inconspicicus</em> Mak. et Pomazkina</td>
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<td>benthic <em>Fragilaria</em> spp.</td>
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<td>chrysophyte cysts 1, 3, 18, 22, 31, 35</td>
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mixing rates) of 1.0 mm yr\(^{-1}\). Using this rate, our Core 316 records an estimated sedimentation history from the 1660's to 1991 A.D. Station 8 from Edgington et al. (1991) was located in Baikal’s north basin and their core had a linear sedimentation rate of 0.26 mm yr\(^{-1}\). With this sedimentation rate used to estimate dates, our Core 324 preserves a sediment record from the late 800's to 1991 A.D.

**Results**

Biogenic silica analysis indicated that the depositional environments of the two sampling sites differ (Fig. 2). On a weight percent basis, Core 316 had less silica content at all depths and it remained relatively constant downcore, ranging only from 19 to 31 weight percent. In contrast, Core 324 had over 80 weight percent in the surface layer, dropped to 43% by 5.0 cm and remained near that level downcore, except at 26–27 cm where biogenic silica fell to only 23%.

The distribution of total microfossils in both cores (Fig. 3) was very similar to their respective biogenic silica profiles. Core 324 had greater microfossil abundance at all depths compared to core 316, except at 26–27 and 28–29 cm. Of the three types of siliceous microfossil remains, diatoms were the most abundant, while chrysophyte cysts made up a significant second portion of both cores. The third category, sponge spicules, were enumerated and found in both cores, but never in great abundance. However, they were included in total microfossil abundance. Because of their size and robust construction, sponge spicules may represent significant portions of biogenic silica in cores (Conley & Schelske, 1993); however, we believe that
Table 2. Diatom and chrysophyte taxa used in cluster analysis, Core 316. Taxa chosen were present in five or more sample intervals at >1% relative abundance

_Aulacoseira baicalensis_ (C. Meyer) Simonsen
_Aulacoseira baicalensis_ fo. _square-punctae_
_Aulacoseira baicalensis_ fo. _compacta_
_Aulacoseira ‘spore’_ (Stoermer et al., 1995)
_Cyclotella baicalensis_ Skv.
_Cyclotella ornata_ (Skv.) Flower
_Cyclotella minuta_ (Skv.) Antipova
_Stephanodiscus binderanus_ (Kütz.) Krieg.
_Stephanodiscus inconspicuus_ Mak. et Pomazkina
_Synedra acus_ Kütz.
_Achnanthes_ spp.
_Cocconeis_ spp.
benthic _Fragilaria_ spp.
chrysophyte cysts 1, 3, 22, 35

Fig. 2. Weight % biogenic silica content (g biogenic silica per 100 g dry sediment) of two Lake Baikal cores (Core 324-filled circles, Core 316-open squares). Vertical scale is depth (cm) from sediment surface.

because of their low abundance in Baikal sediments, their contribution was minimal.

The sedimentary diatom assemblage contained remains of both planktonic and benthic/periphytic diatom communities. In both cores, planktonic diatom remains represented the greatest portion of the assemblage. In the northern basin core (Fig. 4), planktonic diatoms made up between 69 and 79% of the microfossil assemblage with slightly greater contribution in the upper sediments. Benthic diatom remains made up between 1 and 5.6% with somewhat greater benthic proportions below 4.0 cm. Core 316 had a greater relative proportion of benthic diatoms compared to Core 324, with between 6.7 and 19.6% (Fig. 5). A peak in benthic abundance occurred at 6–7 cm with declining abundance downcore. Planktonic diatoms contributed from 77.7 to 85% of the microfossil assemblage in Core 316. Their distribution nearly mirrored the benthic diatom assemblage with greatest relative abundance in the 0.5 to 5.0 cm section with a second peak at 20–21 cm.

Chrysophyte cysts are very common in Lake Baikal sediments and form a morphologically diverse assemblage. Over 100 morphotypes have been identified to date. In Core 324, the nearsurface sediments contained 10 to 15% cysts, while below 5.0 cm, cysts made up nearly 20% of the microfossil assemblage (Fig. 4). Core 316 had slightly lower chrysophyte abundance with 6 to 10% abundance from 0.5 to 5.0 cm and between 7.5 and 21% abundance downcore (Fig. 5).

Core 324-North Basin

To simplify further presentation of results, the distribution of specific taxa and the results of cluster analyses from the two cores will be presented separately.

Cluster analysis on Core 324 resolved four microfossil assemblage zones (Fig. 6). Zone 1 represents samples from the surface to 2.5 cm and covers estimat-
ed dates from 1890’s to 1991 A.D. Zone 3 (c. 1640’s–1830’s A.D.) had the highest order of differentiation in our cluster analysis, and represents samples from 4.5 to 8–9 cm. The last cluster incorporates two zones within Core 324. Zone 2 identifies three samples from 3.0 to 4.0 cm and represents sediments deposited between the 1830’s and 1890’s A.D. Also included in this cluster is Zone 4, the four deepest samples from this core. Dates of Zone 4 are estimated between the 870’s and 1450’s A.D. and include sediments from 14–15 to 28–29 cm. These four zones correspond to distinct changes in the absolute and relative abundance of specific taxa that have occurred in the recent history of northern Lake Baikal.

Aulacoseira baicalensis (C. Meyer) Simonsen is the most conspicuous member of the spring plankton community and is an endemic species to Lake Baikal. Aulacoseira baicalensis begins its vernal development especially under clear ice during February–March, reaches its population maximum in April, and continues to dominate the plankton until May–June (D. H. Jewson, pers. comm.). It has a narrow temperature optima of 4–6 °C and forms physiological resting cells in darkness and during sedimentation (Bondarenko et al., 1993). Its population numbers vary dramatically from year to year producing what are known as ‘Aulacoseira’ years every 3–4 years (Kozhov, 1963). Valve morphology of A. baicalensis is

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**Fig. 4.** Absolute abundance (top panels, no. × 10^6 microfossils per g dry sediment) and % relative abundance (lower panels, % relative to total microfossil assemblage) of benthic and planktonic diatoms and total chrysophyte cysts in Core 324.

**Fig. 5.** Absolute abundance (top panels, no. × 10^6 microfossils per g dry sediment) and % relative abundance (lower panels, % relative to total microfossil assemblage) of benthic and planktonic diatoms and total chrysophyte cysts in Core 316.
extremely variable (Likhoshway et al., 1993), possibly similar to the silica-related variation seen in *A. islandica* (O. Müll.) Simonsen and *A. granulata* (Ehrenb.) Simonsen (Stoermer et al., 1981). Skvortzow (1937) recognized three formae of this taxon based on degree of silicification and areolar structure; the nominate, fo. *compacata*, and fo. *oblonga-punctata*. We also separated three different forms of *A. baicalensis* in our analysis. The most coarsely silicified morphology, fo *compacata* (Fig. 15), and those valves slightly less silicified, the nominate (Fig. 16), correspond to Skvortzow’s (1937) similar formae. The most lightly silicified valve morphology had areolae that appear square in the light microscope. We have termed this form ‘square-punctae’ (Fig. 17) in our analyses. It corresponds to Skvortzow’s (1937) ‘immature’ cell wall.

All three forms of *A. baicalensis* were well preserved in Core 324 (Fig. 7). This taxon was the most abundant siliceous microfossil except in sediment Zone 3, where *Cylotella minuta* was more abundant. Of the three morphotypes, the coarsely silicified form, *A. baicalensis* fo. *compacata*, was most abundant at all levels. It had two abundance peaks in Zones 1 and 4, at 1.0 and 20–21 cm, respectively. Zone 3 represented the period of lowest relative abundance of *A. baicalensis* in Core 324. The more lightly silicified morphs, the nominate and fo. *square-punctae*, were present at all levels, but had their greatest development in near surface Zone 1, together making up >10% of the assemblage.

A second *Aulacoseira* species is also found as a less common member of Baikal’s plankton and sedimentary assemblage. We feel that it is presently of uncertain taxonomic affinity, and in this study will refer to it as *Aulacoseira* ‘spore’ (Figs 18–19) following Stoermer et al. (1995). This taxon has been reported as *Melosira baicalensis* ‘sporangial frustule’ (Skvortzow, 1937), *A. islandica ssp. helvetica* (O. Müll.) Simonsen (Kozhov, 1963; Kozhova et al., 1982), and *A. islandica* (Genkal & Popovskaya, 1991; Bradbury et al., 1994). In sedimentary assemblages, it is found nearly exclusively in the resting spore morphology (Fig. 18). Vegetative valves are much more lightly silicified than spores and are subject to breakage and/or dissolution during sedimentation (Fig. 19, arrow). This taxon has similar ecological characteristics as *A. baicalensis*, i.e. is found most abundantly from March to May, and may bloom under the ice (Kozhova et al., 1982; D. H. Jewson, pers. comm.). Its growth optimum is 4–6°C, and it forms true resting spores in the absence of light (Bondarenko et al., 1993). In surficial sediment collections, *Aulacoseira* ‘spore’ was more common in southern Baikal (Stoermer et al., 1995). Sedimentary distribution of *Aulacoseira* ‘spore’ in Core 324 was represented by two peaks in abundance (Fig. 7), the first in the near surface sediment Zone 1 (1.0 cm) and maximum abundance in the deepest sediment Zone 4 (28–29 cm).

Three endemic species of *Cylotella* are commonly found in the plankton and sediments of Lake Baikal. All three were originally described as varieties of *C. baicalensis* (Skvortzow, 1937), but we have separated these three taxa in full agreement with the recent detailed ultrastructural analysis by Flower (1993). The *Cylotella* taxa are characteristic of mid-summer to late-fall pelagic plankton collections in Lake Baikal, but can be found year round. They usually have their strongest development in *Aulacoseira*-poor years. The modern planktonic *Cylotella* assemblage is dominated by *C. minuta* (Skv.) Antipova (Fig. 24). *Cylotella ornata* (Fig. 23) and *C. baicalensis* (Fig. 22) have become less common in recent years (Flower, 1993).

All three *Cylotella* taxa had their abundance peaks in Zone 3. *Cylotella baicalensis*, the largest taxon, increased in abundance upcore to a peak at 6–7 cm, then generally declined towards the surface sediments except for a small peak at 3.0 cm (Fig. 8). *Cylotella ornata* had greater abundance than *C. baicalensis* at
all depths but had a similar distribution, increasing upcore to a peak at 6–7 cm in Zone 3, then decreasing towards the surface (Fig. 8). *Cyclotella minuta* was the dominant *Cyclotella* at all depths in Core 324 (Fig. 8). Its lowest absolute abundance was in the bottom two samples (26–27 and 28–29 cm) after which it increased to a peak in relative and absolute abundance in Zone 3. *Cyclotella minuta* then decreased upcore to a minimum in relative abundance in the top four samples.

*Stephanodiscus inconspicuus* Mak. et Pomazkina (Figs 25–27) is a small diatom originally reported as *Stephanodiscus* sp. 2 (Khursevich, 1989; Bradbury et al., 1994), and it has only recently been fully described (Makarova & Pomazkina, 1992). Stoermer et al. (1995) found *S. inconspicuus* widely distributed throughout surficial sediments in the Baikal basin. Aspects of its ecology are unknown, but it appears to be another Lake Baikal endemic. Bradbury et al. (1994) suggested that this taxon may be distributed in shallow waters, and related to nutrient input and warm water areas. In Core 324, *S. inconspicuus* was found in all levels and had a peak in absolute and relative abundance in Zone 3 at 4.5 cm (Fig. 7). Above and below this depth there was a general decrease in its abundance, with an absolute and relative minimum at 1.5 cm.

In our cluster analyses, all species of benthic *Fragilaria* (sensu lato) were grouped. *Fragilaria* identifications must often be made in girdle view in sediment slide preparations which unfortunately do not provide absolute taxonomic certainty at the specific level. However, we feel that the similar ecological role
that small *Fragilaria* play justifies grouping these taxa. In Lake Baikal sediments, the small *Fragilaria* spp. were most commonly *F. pinnata* Ehrenb., *F. construens* Ehrenb. and *F. construens* var. *minuta* Temp. and M. Perag. The benthic *Fragilaria* ranged in distribution from nearly zero to 3% relative abundance (Fig. 8). *Fragilaria* increased in abundance upcore from Zone 4 to their peak abundance in Zone 3 at 6–7 cm. Their abundance then generally declined towards the surface sediments except for a small peak at 2.0 cm.

Chrysophyte cysts comprised up to 20% of the microfossil assemblage at some depths in Core 324. Six different chrysophyte cyst morphotypes were present in sufficient quantity to be included in our cluster analysis. Cyst #1 (Fig. 28) had generally consistent absolute distribution in the core but had higher relative abundance in downcore Zones 3 and 4 (Fig. 8). Cyst #3 (Fig. 29) also had its greatest relative abundance in Zones 3 and 4, and fluctuated at lower absolute abundances upcore (Fig. 9). Cyst #18 (Fig. 30) had relatively stable absolute and relative abundances from 0.5 to 5.0 cm, then increased downcore to a peak in Zone 3 at 8–9 cm (Fig. 9). Below this depth, cyst #18 contributed between 4 and 5% relative abundance except for an absolute and relative minimum at 26–27 cm (Fig. 9). Cyst #22 (Fig. 31) had a nearly identical distribution profile as cyst #18 although was found in slightly less quantity throughout the core (Fig. 9). Cyst #31 (Fig. 32) had its maximum abundance within Zones 2 and 3, before declining to a minimum at 26–27 cm downcore. Lastly, cyst #35 (Fig. 33) exhibited a unimodal sedimentary distribution centered around a peak in Zone 3 at 8–9 cm, and declined in absolute and relative abundance both up and downcore (Fig. 9).
Core 316-Central/Southern Basin

Cluster analysis performed on the Core 316 assemblages revealed four distinct zones of microfossil succession (Fig. 10). One cluster, Zone 3, was represented by a single depth, 20–21 cm. This depth approximately corresponds to sediments laid down during the late 1700’s. Zone 1 clustered sediments levels from 0.5 to 4.0 cm which were estimated to date between the early 1950’s and 1991 A.D. Zone 2 (c. 1840’s–1940’s A.D.) resolved sediment depths from 5.0 to 14–15 cm. The last cluster, Zone 4, groups the two lowest depths Core 316, 26–27 and 32–33 cm. These deepest sediments have been estimated to date between the 1660’s and 1730’s A.D. In presenting the distribution of specific taxa from Core 316, we will refer to their abundance within these four cluster analysis zones (Fig. 10).

All three morphologies of *Aulacoseira baicalensis* were present in each depth of Core 316, with *A. baicalensis* fo. *compacta* (Fig. 15) the dominant morphotype at all levels (Fig. 11). The three morphotypes had their greatest abundance in Zone 1 sediments. Below Zone 1, the absolute and relative abundances of all morphotypes declined sharply. *Aulacoseira baicalensis* fo. *square-punctae* (Fig. 17) and the nominate variety (Fig. 16) were present only in very low abundance below 4.5 cm. *Aulacoseira baicalensis* fo. *compacta* (Fig. 15) continued to be the dominant morphotype downcore, contributing slightly less than 5% relative abundance in Zones 2–4.

*Aulacoseira* ‘spore’ (Figs 18–19) was found at its greatest abundance in the Zone 1 sediments from 0.5
Fig. 10. Cluster analysis based on most abundant diatom taxa and chrysophyte cyst morphotypes (Table 2) in Core 316. Horizontal scale is relative Euclidean distance. Vertical scales represent sample depths, Zones of inferred similarity, and dates associated with Zone boundaries.

to 4.0 cm. Below this depth it had relatively constant low abundance before a slight increase in Zone 4 of Core 316 (Fig. 11).

*Cyclotella minuta* (Fig. 24) was the most common *Cyclotella* taxon in Core 316. It was the dominant taxon downcore before sharply decreasing to its lowest relative abundance in Zone 1 (Fig. 12). It had a strong absolute abundance peak at 4.5 and 5.0 cm in Zone 2 and a downcore absolute minimum at 26–27 cm in Zone 4, although it still represented 30% of the microfossil assemblage at that depth. *Cyclotella ornatata* (Fig. 23) had a very similar pattern of distribution as *C. minuta*, indicating autecological similarity. It was generally at lowest relative abundance in Zone 1 sediments (Fig. 12). Two peaks of absolute and relative abundance occurred in Zone 2 at 4.5 and 14–15 cm. In downcore Zones 3 and 4, *C. ornatata* represented between 6.5 and 9.0% relative abundance. The largest Baikal endemic *Cyclotella, C. baicalensis* (Fig. 22), was the least abundant member of this genus at all depths (Fig. 12). It too was found at low abundance in Zone 1 sediments, but increased downcore to generally contribute between 1 to 3% relative abundance.

Both *Stephanodiscus binderanus* (Kütz.) Krieg, and *S. binderanus var. baicalensis* Popovsk. et Genkal have been reported from Lake Baikal (Fig. 20). However, because their identification in sediments must often be made in girdle view (Fig. 20), the two varieties are sometimes indistinguishable, and we have grouped them under the nominate variety. This diatom has been reported from Lake Baikal since the 1930’s (Skvortzow, 1937). It is a vernal taxon in shallow areas, particularly some of the larger bays (Kozhov, 1963) but also has strong development in some years in pelagic waters (Popovskaya, 1991). In the Laurentian Great Lakes, *S. binderanus* is common in more eutrophic inshore waters, and is found in spring offshore collections in the lower lakes (Stoermer & Yang, 1969).

The distribution of *S. binderanus* in Core 316 was characterized by relative and absolute abundance peaks in Zones 1 and 3 (Fig. 12). Zone 3 was represented by a single sample that had greater than 20% relative abundance of *S. binderanus*. A very sharp increase in abundance between samples 4.5 and 4.0 cm delimits the separation between Zones 2 and 1.

*Stephanodiscus inconspicuus* (Figs 25–27) was found in all sediment depths of Core 316, but had its greatest absolute abundance in near surface sediment Zone 1 (Fig. 11). Moving downcore there was a second abundance peak in Zone 2 at 14–15 cm before abundances dropped to constant levels through Zones 3 and 4.

*Synedra acus* (Fig. 21) has become an increasingly important member of the Baikal plankton. It is found year-round, but most prominently from April–June. It has noticeably increased in abundance since the 1950’s (Popovskaya, 1993), and has been shown to have a wide breadth of growth optima (Bondarenko *et al.*, 1993). It has also apparently been a conspicuous member of the plankton during the history of Baikal. Bradbury *et al.* (1994) reported on its periodic occurrence since the late Pleistocene in Lake Baikal, suggesting its association with more nutrient input and warmer climates. Sedimentary remains of *S. acus* were usually fragmented.

*Synedra acus* had two abundance peaks in Core 316 (Fig. 12). The first occurred in near surface Zone 1 with maximum abundance at 1.5 cm. The second peak was in Zone 4 and represented the absolute and relative abundance maximum for *S. acus*.

Because of their similar ecological role in the diverse periphyton of Lake Baikal (Skvortzow, 1937; Foged, 1993), *Achnanthes* and *Cocconeis* taxa were each grouped at the generic level for statistical analyses. *Achnanthes* species most commonly encountered in Core 316 sediments were *A. oestrupi* (A. Cl.) Hust., *A. lanceolata* (Bréb.) Grun., *A. calcar* Cl., *A. striata* Skv. and C. Meyer, and *A. minutissima* Kütz. The *Cocconeis* assemblage was dominated by *C. placen-
At the top of Zone 2, abundances fell to <2% compared to Core 324. The four that were included, however, had their greatest absolute abundance in Zone 1, with secondary peaks at 8–9 cm in Zone 2 and in Zone 4. The small benthic Fragilaria taxa were also grouped in Core 316, similar to Core 324. The Fragilaria assemblage contributed >2% relative abundance at all core depths except at 14–15 cm. Cocconeis spp. were also distributed at nearly constant relative abundance downcore (Fig. 13). In general, however, they had their greatest absolute abundance in Zone 1, with secondary peaks at 8–9 cm in Zone 2 and in Zone 4.

Fewer chrysophyte cyst morphotypes met our inclusion criteria for cluster analysis of Core 316 compared to Core 324. The four that were included, however, had also been selected for the Core 324 cluster analysis. Cysts #1 (Fig. 28) and #3 (Fig. 29) had similar distributions in Core 316 (Fig. 13). Both had highest abundances in Zone 4 and then dropped to approximately 2% relative abundance through Zones 2 and 3. At the top of Zone 2, abundances fell to <2% where they remained to the surface. Cyst #22 (Fig. 31) had quite variable distribution in the core. It was found in greatest absolute and relative abundance in the top-and bottom-most depths (Fig. 14). Cyst #35 (Fig. 33) had two peaks in abundance in Zones 2 and 3 at 8–9 and 20–21 cm respectively (Fig. 14). Lower abundances of cyst #35 were generally found in Zones 1 and 4 except for a local minimum at 14–15 cm.

Discussion

Lake Baikal has no modern analog. Its size, depth, and latitudinal position create unique limnological conditions and, because of the lake’s great age and relative geographic isolation, the Baikal flora and fauna have developed an extremely high degree of endemism. Endemism is reflected at all levels of the food web, and is especially evident in the diatom community. The plankton community is represented by very few endemic taxa adapted to Baikal’s unique pelagic environment (Skvortzow, 1937; Bradbury et al., 1994). This phenomenon of nondiverse plankton is not observed in other large Pleistocene-age lakes (Stoermer & Yang, 1969). The benthic and periphytic diatom communities have also developed endemically but contrary to planktonic assemblages are extremely diverse and contain ‘swarms’ of related species (Skvortzow & Meyer, 1928; Skvortzow, 1937; Stoermer et al., 1986; Kociolek & Stoermer, 1988; Foged, 1993). This confounds any effort to create a modern analytical calibration set (Fritz et al., 1991) for resolving downcore changes in Lake Baikal’s past. Our downcore interpretations of siliceous microfossil remains must instead be based on autecological characteristics of Baikalian algae, an analytical approach that has proven successful in paleolimnological interpretations of other large lake systems (Stoermer et al., 1993).

The sedimentary environment in Lake Baikal is characterized by extremely high biogenic silica levels and large depositional zones (Williams et al., 1993). In contrast, large Pleistocene-age lakes often have discrete depositional basins and an order of magnitude less sedimentary biogenic silica content than Lake Baikal (Stoermer et al., 1985b). Characteristics of Baikal that create these sedimentary conditions are its great depth, steep sided basin morphometry, rather high levels of dissolved silica, cold waters, and incomplete vernal and autumnal mixing (Weiss et al., 1991; Granina et al., 1993; Williams et al., 1993). Biogenic silica profiles of Cores 324 and 316 also reflected the difference in depositional environments between the northern and southern basins, with much greater silica content in northern basin sediments (Fig. 2). Several studies (Granina et al., 1993; Williams et al., 1993; Stoermer et al., 1995) have documented identical trends and suggested that biogenic silica sedimentation patterns in Lake Baikal reflect both the slightly higher water column silica values in the northern basin and, in southern Baikal sediments, dilution by terrigenous clastic inputs, especially from the Selenga River (Fig. 1). This dilution factor masks sedimentary signals of greater water column productivity in the south.

The northern basin of Lake Baikal is removed from most direct anthropogenic impacts affecting the central and southern basins. It has lower productivity and less terrigenous input than the southern basin. Modern phytoplankton assemblages are dominated by the endemic Aulacoseira and Cyclotella with very few of the more ubiquitous diatoms encountered in southern Baikal (Stoermer et al., 1995). The four successional zones resolved with cluster analysis (Fig. 6) of Core 324 were separable based on their assemblage characteristics, and we suggest that these changes reflect climatic patterns over the last ~1100 years. The most recent sediments in the northern basin (Zone 1, c. 1890’s–1991 A.D.) were characterized by an increased abundance
Fig. 11. Absolute (top panels, no. x 10^6 microfossils per g dry sediment) and relative (lower panels, %) abundance of some major diatom taxa in Core 316. Left vertical scale is depth (cm) from sediment surface, right scale indicates microfossil succession zones from cluster analysis (Fig. 10).

of *Aulacoseira baicalensis* and *Aulacoseira* 'spore' (Fig. 7). Zones 1 and 2 also identify a period of increasing biogenic silica accumulation in northern Lake Baikal (Fig. 2). Zone 1 sediments encompass the recent climatic warming that had its maximum c. 1940 A.D. (Khotinskiiy, 1984). The abundance of vernal *Aulacoseira* taxa and increased biogenic silica accumulation suggests warmer climates associated with longer, more productive vernal growth and milder winter conditions. However, caution should be exercised when correlating these recent assemblage changes and productivity increases solely to climatic variables. In other ultra-oligotrophic north temperate lakes, recent assemblage shifts appear to also reflect unique changes associated with modern day atmospheric loadings (Stoermer et al., 1985c; Stoermer et al., 1990).

In contrast, Zone 3 sediments (c. 1640's–1830's A.D.) were dominated by the endemic *Cyclotella* with reduced abundance of *Aulacoseira*. Estimated dates from Zone 3 correspond to a climatic period, known as the Little Ice Age (late 1400's to 1850 A.D.), which was characterized by cooler climatic conditions in Siberia (Khotinskiiy, 1984). The shift in dominance in the planktonic community from *Aulacoseira* to *Cyclotella* provides an indicator of this climate change. During cooler conditions, heavier ice and possibly snow cover would decrease light penetration below the ice and shorten the spring turnover period, thereby limiting late winter and vernal growth of the endemic *Aulacoseira* taxa (Lund, 1966; D. H. Jewson, pers. comm.). Concomitantly, greater abundance of the dominant summer and autumn planktonic *Cyclotella* would be expect-
ed. The microfossil assemblage between Zones 1 and 3 (Zone 2, c. 1830's–1890's A.D.) and in Zone 4 (c. 870's–1450's A.D.) are similar to one another (Fig. 6) suggesting they represent transitional intervals between cooler and warmer climates as Baikal's plankton shifted between *Cyclotella* and *Aulacoseira*-dominated assemblages. Bradbury & Dieterich-Rurup (1993) interpreted a similar shift in planktonic dominance from *Fragilaria crotonensis* Kitton to *Stephanodiscus minutus* Grun. in Elk Lake, Minnesota, as a response to cooler conditions during the Little Ice Age.

Other taxa showing similar downcore patterns between Zones 1 and 3 of Core 324 as the *Cyclotella* spp. are chrysophyte cyst #35 (Fig. 9), *Stephanodiscus inconspicuus* (Fig. 7), and the benthic *Fragilaria* spp. (Fig. 8). While we know little about the ecology of the Baikal chrysophytes, increased deposition of cysts has been correlated to persistence of higher productivity into summer (Bradbury & Dieterich-Rurup, 1993), a condition that would have also promoted strong growth of the endemic *Cyclotella*. Relatively high abundance of *Fragilaria* and *S. inconspicuus* during the Little Ice Age period would also be expected. Along with these two taxa, the grouped benthic taxa also had their maxima in Zone 3 sediments (Fig. 4) suggesting enhanced microfossil contribution from shallow water productivity relative to offshore production during this cooler climate period. Thicker ice cover and lengthier ice breakup periods during cooler climates may have produced extended periods with ice-free bays and shallows similar to the moat hypothesis of Smol (1988).

In contrast to the north basin, the southern end of Lake Baikal has been the site of some regional urbanization and industrialization. Impacts on the lake from atmospheric and direct loadings are cen-
Fig. 13. Absolute (top panels) and relative (lower panels) abundance of some major benthic diatom taxa and chrysophyte cyst morphotypes in Core 316. Scale units as in Fig. 11.

Fig. 14. Absolute (panels 1&3) and relative (panels 2&4) abundance of two major chrysophyte cyst morphotypes in Core 316. Scale units as in Fig. 11.
Plate 1. Dominant Lake Baikal diatoms found in sediment assemblage. Scale bar = 10 μm. All figures at same magnification (Scale bar in Fig. 21). Fig. 15. Aulacoseira baikalensis fo. compacta. Fig. 15a. High focus. Fig. 15b. Mid-focus showing extreme thickness of valve mantle. Fig. 16. Aulacoseira baikalensis nominate form. Fig. 16a. High focus. Fig. 16b. Mid-focus showing thinner mantle region. Fig. 17. Aulacoseira baikalensis fo. square-punctae. Figs 18–19. Aulacoseira ‘spore’, note collapsed remains of vegetative frustule in Fig. 19 (arrow). Fig. 20. Stephanodiscus binderanus. Fig. 21. Synedra acus.
Plate 2. Dominant Lake Baikal diatoms and chrysophyte cysts found in sediment assemblage. Scale bars = 10 μm for Figs 22-24, scale bars = 5 μm for Figs 25-33. Fig. 22, Cyclotella baicalensis. Fig. 23, Cyclotella ornata. Fig. 24, Cyclotella minutula. Figs 25-27, Stephanodiscus inconspicuus. Fig. 28, Baikal chrysophyte cyst #1. Fig. 29, Baikal chrysophyte cyst #3. Fig. 30, Baikal chrysophyte cyst #18. Fig. 31, Baikal chrysophyte cyst #22. Fig. 32, Baikal chrysophyte cyst #31. Fig. 33, Baikal chrysophyte cyst #35.

tered around industrial towns and sites on the southern shore and loading from upstream cities (Ulan Ude and Selengisk) on the Selenga River (Fig. 1; Galazii, 1991; Belt, 1992). Confounding sedimentary interpretation of anthropogenic effects on the lake are climatic changes that must have also affected southern Baikal.
Core 316 preserved a much shorter history than Core 324, thus pre-settlement conditions are poorly resolved. Higher sedimentation rates found between Baikal's southern and central basin (Edgington et al., 1991) allowed recovery of a record extending from c. 1660's–1991 A.D. Our cluster analysis resolved four zones of microfossil succession in Core 316 (Fig. 10). Zone 1 (c. early 1950's–1991 A.D.) corresponds to post-World War II industrialization, a period when planktonic communities became dominated by endemic *Aulacoseira* (Fig. 11). The increase in *Aulacoseira* abundance is probably indicative of both the recent climatic warming (Khotinskiy, 1984) and increased productivity from nutrient loadings. An increased abundance in Zone 1 of non-endemic diatoms, such as *Stephanodiscus binderanus* (Fig. 12), *Synedra acus* (Fig. 12), and *Cyclostephanos dubius* (Fricke) Round (not figured), all considered more eutrophic taxa, also lends credence to a hypothesis that anthropogenic impacts from development and increased nutrient loadings have occurred in this region of Lake Baikal. These changes also appear to have impacted certain chrysophyte populations in southern Baikal. Cysts #1, #3, and #35 experienced reduced abundance in the transition from Zone 2 to Zone 1 (Figs 13, 14).

In sediments predating Zone 1 in Core 316 (Zones 2–4, c. 1660's–1950's A.D.), endemic *Cyclotella* taxa dominated the planktonic community in southern Lake Baikal (Fig. 12). Zone 2 sediments (c. 1840's–1950's A.D.) were also characterized by decreased *Aulacoseira* abundance (Fig. 11). Zone 2 may represent restabilization following initial European occupation of the region. Similar 'relaxation' of eutrophication following initial destruction of forests is commonly noted in cores from the Laurentian Great Lakes (Stoerner et al., 1993).

Zone 3 is represented by a single depth (20–21 cm) and is uniquely characterized by an abundance peak of *Stephanodiscus binderanus* that had begun in Zone 4. This depth has been dated c. 1780's A.D., and may indicate that anthropogenically-induced changes occurred in Lake Baikal during early settlement and development in the southern basin. This date corresponds roughly to a time period when Irkutsk and Ulan Ude were seeing significant Russian expansion and development. Changes in the lake may have been driven by nutrient loadings from land clearance, slash and burn agriculture, and urban loading along the Selenga River (Riasanovksy, 1984). Apparently the diatom community responded with increased production of smaller taxa (*S. binderanus*) since the 20–21 cm depth was a local maximum of absolute microfossil abundance but did not show increased biogenic silica deposition (Figs 2, 3). The most recent sedimentary assemblages in Core 316 suggest that a similar response has occurred in southern Baikal since the early 1950's (0.5–4.0 cm in Fig. 3), supporting Popovskaya's (1991) observed increase of small-celled species in the plankton since 1958. The appearance of *S. binderanus* was still overshadowed by dominance of the endemic *Cyclotella* taxa during Zone 3, which suggests that nutrient enrichment was probably most prevalent in nearshore areas of southern Baikal. In the Laurentian Great Lakes *S. binderanus* may form dense nearshore blooms with little invasion of offshore waters (Lorefice & Munawar, 1974).

The two lowest depths in Core 316 were resolved as Zone 4 (c. 1660's–1730's A.D.). This zone had abundant *Cyclotella* but was also characterized by increases in *Synedra acus* and *Aulacoseira* 'spore' (Figs 11, 12). The few samples available make interpretation difficult but, following previous arguments, these samples were likely deposited under conditions that allowed maximum growth of summer-blooming taxa.

The climatic signals recorded in the planktonic assemblages in Core 324 should also provide evidence of Late Holocene climatic shifts in future down-core interpretations. Historical vegetation patterns in Siberia suggest that there have been many major and minor climatic optima since the last glaciation (Khotinskiy, 1984) which should be reflected in northern basin sediments. The southern sediment record provides strong evidence that Lake Baikal is no longer in its pristine state. The appearance of non-endemic diatoms in Baikal points towards changes that have occurred quite dramatically since World War II, which are probably associated with industrialization and urbanization around southern Baikal and along the Selenga River. Changes in the primary producer communities should act as a harbinger of the potential for future changes in all levels of the food chain. In other formerly oligotrophic large lake systems that experienced anthropogenic impacts, changes in primary production levels and assemblage characteristics identified with paleo-limnological analysis were the first signs of environmental impact and future problems (Stoerner et al., 1985b).
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References

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Note added in proof

Stephanodiscus inconspicuus Makarova et Pomazkina has recently been transferred to the new genus Craτeriportula (Flower & Håkansson, 1994), as C. inconspicua (Mak. et Pom.) Flower et Håkansson.