Dynamic diatom response to changing climate 0–1.2 Ma at Lake El’gygytgyn, Far East Russian Arctic

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Abstract. The Lake El’gygytgyn sediment record contains an abundant diatom flora through most intervals of the lake’s history, providing a means to create and test hypotheses concerning the lake’s response to changing climates. The 0–1.2 Ma core interval is characterized by shifts in the dominant planktonic genera and events of exceptional concentration and diversity. Warm interglacial intervals have enhanced concentration and diversity of the plankton. This response is most extreme during exceptional events corresponding to marine isotope stages (MIS) 11 and 31. Diatom concentration and diversity also increase during some cold intervals (e.g., MIS 2), suggesting conditions of lake circulation and nutrient cycling promoting diatom production during these events. Short intervals of low plankton concentration accompanied by shifts in the dominant genus of the lake suggest conditions during certain cold events generate a severe impact on plankton production. The absence of these events during extended intervals of low summer insolation variability suggests a muted cold-event response of the lake system linked to regional climate.

1 Introduction

The sediment record from Lake El’gygytgyn, Far East Russian Arctic, contains a unique archive of terrestrial paleoclimate spanning the last 3.6 million years (Melles et al., 2012). The majority of recovered sediment contains significant but variable biogenic silica (Meyer-Jacob et al., 2013) mostly in the form of diatom valves. Understanding the factors influencing the production and preservation of diatoms in the lake is one key to understanding many of the other proxy records used to unravel the response of the ancient lake system to changing climates. Diatom species assemblages also offer a powerful recorder of changes in the lake potentially linked to paleoclimate through ice conditions, circulation, nutrient delivery, and water chemistry (e.g., Smol, 1988; Douglas and Smol, 1999).

Early research on the modern diatom flora in El’gygytgyn has focused on descriptions of extant taxa found in lake-bottom samples. These first studies describe many unique species, consistent with a long-lived lake system (Sechkina, 1956; Jouse and Sechkina, 1960; Kharitonov, 1980, 1993; Genkal and Kharitonov, 1996, 2005, 2006). Prospects of the recovery of the complete sediment record have inspired a renewed interest in the modern diatom flora (Cremer and Wagner, 2003; Cremer et al., 2005; Cremer and van de Vijver, 2006; Stachura-Suchoples et al., 2008). However, because of its remote location, on-site studies have been only short duration, and the diatom response to interannual changes in the lake system is poorly constrained.

Previous diatom studies of sediment cores from Lake El’gygytgyn show a generally abundant, diverse, and variable flora. The core PG1351 record included 150 samples spanning the last 250 ka (Cherepanova et al., 2007, 2010). Although these initial investigations demonstrated the potential of these diatom records, the interpretations from these early core studies were complicated by the presence of turbidites and by taxonomic uncertainty, especially in the morphologically variable dominant planktonic diatom included in the Cyclotella ocellata complex (sensu Cremer and Wagner, 2003).

This study utilizes samples from a more recent core (Lz1024; recovered in 2003 extending to 340 ka) and initial
samples from the nearby deep drilling site (5011-1) to construct a continuous low-resolution composite record spanning the last 1.2 Ma. Four intervals are analyzed in finer temporal resolution, corresponding to marine isotope stages (MIS) 1, 5e, 11, and 31, to contribute to the research focus on exceptional interglacial events (Melles et al., 2012; Lozhkin and Anderson, 2013; D’Anjou et al., 2013; Vogel et al., 2012) and to help understand conditions during the local Last Glacial Maximum (Holland et al., 2013). This study provides initial interpretations of these diatom records along with guidance and prospects for future investigation.

1.1 Setting

Lake El’gygytgyn (67°30′ N, 172°5′ E; Fig. 1) is situated in an impact crater dated 3.58 ± 0.04 Ma (Layer, 2000). The modern lake (12 km diameter and 175 m deep) is oligotrophic (Nolan and Brigham-Grette, 2007). In recent observed years, the lake is ice free during the period of approximately July–October and completely mixes in summer. Weak sub-ice convection is also suggested by gas bubble patterns in the lake ice (Nolan and Brigham-Grette, 2007). At the present lake level, a shallow shelf (less than 10 m deep) occupies approximately 11% of the lake basin area (Nolan and Brigham-Grette, 2007). However, morphological evidence suggests higher and lower lake levels in the past, substantially impacting the area of shallow-water lake bottom (Fedorov et al., 2008; Juschus et al., 2011). Additional details of the modern setting and limnology are described elsewhere (Nolan and Brigham-Grette, 2007; Fedorov et al., 2012; Nolan, 2012; Nolan et al., 2012; Wennrich et al., 2013).

2 Methods

Diatom samples were prepared from a measured mass of freeze-dried sediment, treated with boiling 30% H$_2$O$_2$. Systematic diatom counts were conducted at 1000 × magnification with a Leica DMLB light microscope with differential interference contrast. At least 500 valves were counted in all but a few sparse samples. Quantitative diatom concentrations were calculated using measured transect areas on slides prepared from a measured sediment mass and added calibrated microspheres settled through a water column onto a microscope cover slip (modified from Battarbee et al., 2001).

Counted diatoms were identified to species level, wherever possible, using a wide variety of taxonomic references. Taxonomy of the dominant diatoms was confirmed by scanning electron microscopy of selected samples. Names assigned to the taxa are generally those used by previous El’gygytgyn investigators (e.g., Sechkina, 1956; Cremer and Wagner, 2003; Cherepanova et al., 2007; Stachura-Suchoples et al., 2008). Presentation and discussion of planktonic diatoms in this study focus on genus-level variations. In most cases, each genus is dominated by a single species as noted in the text. Cyclotella contains multiple morphotypes which, with further taxonomic work, will warrant further subdivision. Here they are combined in the Cyclotella ocellata complex (Cremer and Wagner, 2003) because of the difficulty to distinguish these taxa in routine counts. Aulacoseira subarctica (O. Müller) Haworth is the dominant member of its genus. Minor occurrences of other Aulacoseira species also occur and are excluded from the Aulacoseira plots. Genus-level diversity in the plankton was estimated utilizing the Shannon index (e.g., van Dam, 1982). Further observations of periphyton assemblage, valve size and preservation were noted and quantified in certain cases.

The age model for the Lz1024 core is provided by methods described elsewhere (e.g., Juschus et al., 2007). Below 340 ka, the chronology is based on the detailed age model derived primarily from paleomagnetic reversal and orbital tuning (Melles et al., 2012) and correlation to the base of the Lz1024 record. The initial analysis interval presented here provides an approximately 4–5 kyr average resolution for the
0–1.2 Ma record and 1 kyr resolution for the intervals analyzed in more detail.

Diatom accumulation rates (valves cm$^{-2}$ kyr$^{-1}$) are calculated from concentrations in individual samples (valves g$^{-1}$) using age model sediment accumulation rates and dry bulk density calculated from gamma ray attenuation porosity evaluator (GRAPE) scans (Gebhardt et al., 2013) and are presented for some core intervals. The main results are presented as simple concentrations to describe independently the occurrence of diatoms in the sediment and to avoid introducing uncertainties related to the age model.

3 Results

3.1 The composite record 0–1.2 Ma

The 0–1.2 Ma record presented here preserves a complex cyclicity in the lake’s diatoms, reflected in the concentration, preservation, valve size, and species assemblage (Fig. 2). Because of the great differences in size, silica content, and valve concentration in dominant planktonic diatoms and the sporadic occurrence of some other planktonic taxa, absolute concentrations, rather than percentage within an assemblage, are presented for comparison. Numerical valve counts of total diatoms show a broadly similar trend to the biogenic silica record (Frank et al., 2012; Meyer-Jacob et al., 2013). Subtle differences in these trends may be attributed to minor variation in valve size and preservation observed in these intervals and by contributions from larger, less numerically abundant taxa.

The most notable down-core changes are shifts in the dominant genera. *Pliocaenicus* seczkinae Stachura-Suchoples, Genkal et Khursevich, previously observed to emerge since 15 ka, has a long complex history in El’gygytgyn, exhibiting substantial morphological variability. *Cyclotella* likewise becomes a minor component of the plankton during certain intervals. During other times, both diatoms co-exist, as in the Holocene.

One clear aspect of the 0–1.2 Ma diatom record is the occurrence of intervals of exceptional diatom concentration that correspond to inferred warm intervals (yellow in Fig. 2). Both plankton and periphyton concentrations increase, and the valves exhibit better preservation. Other planktonic taxa including *Stephanodiscus* cf. minutulus (Kützing) Cleve and Möller also occur during some of these events. *A. subarctica* occurs in abundance during some of these events prior to 550 ka, and is the numerically most abundant taxa during particular intervals from 800 to 1200 ka. The most extreme of these exceptional events correspond to MIS 11 and MIS 31. Here both total plankton and periphyton reach concentration peaks unmatched in the studied interval, and a high diversity of planktonic diatoms occurs. Many of the high diatom concentration events are characterized by temporary size increases in the morphologically plastic planktonic genera (especially *Cyclotella* and *Pliocaenicus*). For example, during MIS 11 the initial cell diameter of *Cyclotella* is 25–30 µm, compared to 15–20 µm during inferred colder intervals immediately preceding and following this event. During MIS 31 the initial cell diameter of *Pliocaenicus* increases to 35–50 µm from less than 30 µm before this event.

A few intervals of increased plankton and periphyton abundance correspond to apparently cold intervals (purple in Fig. 2). These are distinguished by their distinctive periphyton assemblage, particularly the absence of species found in inferred warm intervals, especially *Achnanthidium kriegeri* (Krasske) Hamilton, Antoniades et Siver and *Cocconeis placenta* Ehrenberg (see Sect. 3.2). None of these intervals is characterized by significant concentration of *Pliocaenicus*.

Short (< 20 cm) intervals of low plankton abundance also occur in the diatom record. Seven of the 234 samples systematically analyzed for this study contain less than $10^7$ valves g$^{-1}$ of total planktonic diatoms (green in Fig. 2). Some of these samples have zero observed planktonic diatoms or an order-of-magnitude-lower concentration than any other samples. Corrections for age model sedimentation rate and density confirm exceptionally low diatom accumulation rates. Many of these zones have persisting moderate concentrations of well-preserved periphyton. These intervals, in many cases, correspond to shifts in the dominant plankton in the lake. For example, at 924 ka *Pliocaenicus* replaces *Cyclotella*. One such zone occurs at 225 ka. Although *Cyclotella* persists, this interval corresponds to an abrupt decline in a morphotype of the *Cyclotella ocellata* complex (Cherepanova et al., 2010), which dominates the plankton 225–550 ka. This long-term dominance and abrupt decline may justify the establishment of this morphotype as a separate distinct species.

3.2 MIS 1 and 2 (0–34 ka)

The genus *Cyclotella* dominates the sediment record of the last 34 ka (Fig. 3). Minor and sporadic planktonic diatoms vary dramatically across this portion of the record. The 20–25 ka interval exhibits the greatest diversity in plankton with peaks in *Fragilaria cf. navana* Lange-Bertalot, *S. cf. minutulus*, and *Asterionella formosa* Hassal often associated with higher nutrient concentrations (e.g., Lotter et al., 1998; Bennion et al., 2004). This interval corresponds to the highest Si/Ti ratios and biogenic silica concentrations in the last 34 ka, some intervals exceeding 30 %. Average diatom accumulation rates of 20–25 ka approximately equal the average from 0–5 ka (1.8 × 10$^7$ valves cm$^{-2}$ kyr$^{-1}$). The comparatively large, heavily silicified *Pliocaenicus* occurs in abundance only since 15 ka, although a few individual valves are observed on slides from earlier samples.

The periphyton species assemblage also changes with comparable timing. Small Achnanthaceae are numerically the most abundant valves in most samples. *A. kriegeri* and *Achnanthidium minutissimum* (Kützing) Czarnecki occur in greatest abundance since 15 ka. In contrast, *Planolithidium*...
Fig. 2. Concentration (valves g$^{-1}$) of planktonic species, total periphyton, total diatoms, and plankton diversity (Shannon index) from 0 to 1.2 Ma. Exceptional events are indicated by colored intervals. Diatom concentrations, except total diatoms, are plotted on a log scale. For comparison, the marine isotope stack (Lisiecki and Raymo, 2005) and 65° N June insolation (Laskar et al., 2004) are also plotted.
Fig. 3. Concentration (valves g\(^{-1}\)) of planktonic species, total periphyton, total diatoms, and plankton diversity (Shannon index) from 0–34 ka. For comparison, other El'gygytgyn climate proxies including Si/Ti measured by XRF core scanning and MTWM (mean temperature warmest month) derived from pollen data (best modern analog approach) (Melles et al., 2012). Other pollen data (Lozhkin et al., 2007) suggest continuation of a cold assemblage through the 20–34 ka interval (not shown).

3.5 MIS 31 (1057–1113 ka)

The interval corresponding to MIS 31 peak warmth contains a diatom abundance second only to MIS 11 in the 0–1.2 Ma record (Fig. 6). Aulacoseira is the most consistent and numerically abundant member of the plankton assemblage. The robust Pliocaenicus, although numerically less abundant, is also a significant and consistent part of the assemblage. Similar to MIS 11, additional genera occur in the more recent portion; in this case Stephanodiscus and Asterionella occur 1065–1076 ka. Above this interval there is a low plankton abundance zone (< 10\(^7\) valves g\(^{-1}\)) corresponding to the extirpation of Pliocaenicus and return of Cyclotella to dominance in the lake. This low-abundance zone is missed in the 0–1.2 Ma record because of the lower sample resolution, although it is suggested by the observed shift in dominant genus.

3.4 MIS 11 (360–430 ka)

The interval corresponding to MIS 11 exhibits the greatest peak in diatom concentration driven mostly by Cyclotella (Fig. 5). During the first part of this interval (411–416 ka) peak abundance corresponds to numerically abundant but small-diameter Cyclotella and Stephanodiscus. From 411 to 400 ka, Cyclotella occurs in lower abundance but increased size. Diversity also increases significantly in this interval, especially 401–403 ka, where Fragilaria and Asterionella occur 1065–1076 ka. Above this interval there is a low plankton abundance zone (< 10\(^7\) valves g\(^{-1}\)) corresponding to the extirpation of Pliocaenicus and return of Cyclotella to dominance in the lake. This low-abundance zone is missed in the 0–1.2 Ma record because of the lower sample resolution, although it is suggested by the observed shift in dominant genus.

3.3 MIS 5e (111–135 ka)

The genus Cyclotella also dominates the 111–135 ka core interval (Fig. 4). Sparse Pliocaenicus and planktonic Fragilaria occur in a few samples. The inferred peak warmth of MIS 5e corresponds to an approximate doubling of the diatom concentration driven mostly by Cyclotella. Calculated genus-level plankton diversity does not significantly increase in this interval. This index may underestimate the true plankton diversity in this interval because the dominant Cyclotella contains at least two morphotypes with significantly different diameter. The peak in diatoms seen at 135 ka corresponds to the preceding cold productive mode (Fig. 2).

The genus Cyclotella also dominates the 111–135 ka core interval (Fig. 4). Sparse Pliocaenicus and planktonic Fragilaria occur in a few samples. The inferred peak warmth of MIS 5e corresponds to an approximate doubling of the diatom concentration driven mostly by Cyclotella. Calculated genus-level plankton diversity does not significantly increase in this interval. This index may underestimate the true plankton diversity in this interval because the dominant Cyclotella contains at least two morphotypes with significantly different diameter. The peak in diatoms seen at 135 ka corresponds to the preceding cold productive mode (Fig. 2).

**oestrupii** (Cleve-Euler) Round et Bukhtiyarova peaks from 19 to 24 ka. Diatoms with valves greater than 75 µm length vary more irregularly. However, *Frustulia rhomboides* var. amphipleuroides (Grunow) DeToni is almost exclusively observed 15–25 ka, and *Eunotia pseudopectinalis* Hustedt is more characteristic of the interval since 15 ka.
4 Discussion

4.1 Peak warmth intervals

Observing the Holocene and other identified interglacial intervals from the composite core record, a clear pattern in diatom response is apparent. Concentration increases in both periphyton and plankton. Genus-level diversity increases in the plankton. Valves are generally well preserved with only minor dissolution noted. The occurrence of certain periphytic taxa also characterizes each of these intervals, especially A. kriegeri and C. placentula. These taxa also occur in the deeper record, even as the dominant plankton varies. In general, the diatom response is consistent with a relatively favorable environment for diatom growth that might be anticipated from enhanced nutrient delivery, prolonged open-water growing season, and expanded diversity in habitats in the lake and its catchment.

The emergence of Pliocaenicus since 15 ka may be a response to warming and enhance open-water conditions. Little is known about the ecological tolerance of this taxon. Flower et al. (1998) speculate the similar Pliocaenicus costatus var. sibiricus (Skabitch.) Round et Håkansson responded favorably to 20th-century warming in Lake Bolschoi, Siberia. The appearance of Pliocaenicus in El’gygytgyn at 15 ka may also partially reflect the elimination of a competitive exclusion from larger members of the Cyclotella ocellata complex. These forms disappear from the record during an inferred cold, diatom-poor interval at ca. 70 ka. Thus, favorable conditions for this large, heavily silicified Pliocaenicus may first recur after 15 ka, allowing expansion into this vacated niche in the phytoplankton. The linkage between Pliocaenicus abundance and climate is further strengthened by the correspondence with its emergence 13–15 ka, peak at 9–10 ka, and subsequent decline through the Holocene with the pollen-derived summer temperature (Fig. 3) (Melles et al., 2012).

Although not yet quantified in detail, the observed size increase in Pliocaenicus or Cyclotella, particularly in MIS 11 and 31, may also be a response to conditions during these times. Diatom size control may vary with conditions of a specific lake (e.g., Finkel el al., 2009). In a nutrient-limited lake system, higher nutrients (including silica) may favor larger plankton. Further, larger plankton could be an adaptation to greater mixing/turbulence due to more open water, allowing larger diatoms to remain in the photic zone without sinking. On longer timescales, individual species may adapt to competition introduced during intervals favorable for phytoplankton production. The El’gygytgyn sediment record may provide an exceptional opportunity to explore these phenomena on glacial-interglacial timescales.

Observed interglacials within the 0–1.2 Ma record are highly variable in the magnitude of their diatom response. MIS 11 and 31 are exceptional events both in diatoms and other proxies (Melles et al., 2012). Both intervals have peaks in species usually associated with enhanced nutrient status. The occurrence of these assemblages corresponds to sediments Facies C, sensu Melles et al. (2012), characterized by reddish-brown, finely laminated silt, explained by high productivity in the growing season and anoxic bottom waters during winter. The unique diatom response may result from enhanced nutrient delivery from the catchment or a threshold in lake-water oxygenation and circulation, altering the nature and timing of nutrient cycling in the lake (e.g., Rippey et al., 1997; Brüchman and Negendank, 2004). This exceptional diatom production and accumulation plays an important role in the formation of this sediment facies, ranging between 20 and 50 % biogenic silica during these peaks.
Fig. 5. Concentration (valves g\(^{-1}\)) of planktonic species, total periphyton, total diatoms, and plankton diversity (Shannon index) from 360 to 430 ka. Other El’gygytgyn climate proxies from Melles et al. (2012) are included for comparison (see Fig. 3).

Correlation between the plankton assemblages and other proxy records during MIS 11 and 31 (Figs. 5 and 6) affirms their climate sensitivity. Initial warming at 427 ka (MIS 11) is mainly expressed as an increase in *Cyclotella*. Following modest declines in Si/Ti and pollen-derived summer temperature, *Stephanodiscus* occurs at levels comparable to *Cyclotella* 400–417 ka. *Asterionella* also occurs sporadically in this interval. This event corresponds to the peak of MIS 11, inferred from pollen and biomarker temperature reconstructions (Melles et al., 2012; D’Anjou et al., 2013). These species abruptly decline as reconstructed summer temperatures first approach near-modern values at ca. 400 ka. During MIS 31, initial warming ca. 1085–1090 ka corresponds to the emergence of *Aulacoseira* as the dominant planktonic diatom. *Stephanodiscus* and *Asterionella* similarly rise during the peak in pollen-indicated summer temperature (1076 ka) and abruptly decline at near-modern conditions (1065 ka). This lag further supports the hypothesis of a threshold in climate-driven nutrient conditions in the lake during extreme interglacials.

The distribution of *A. subarctica*, a widespread diatom preferring intermediate nutrient conditions (Gibson et al., 2003), illustrates some of the complexity of comparing particular interglacials. This diatom, occurring almost exclusively in
identified interglacial intervals, is a minor component since 550 ka but dominates in some intervals between 830 and 1150 ka. This species is also abundant in Lake Baikal during approximately this same time, peaking ca. 850, 1150, and 1190 ka (Grachev et al., 1998), suggesting a broader biogeographic influence on its occurrence and distribution.

4.2 Cold productive intervals

The occurrence of high biogenic silica (20–35 %) and diatom production during MIS 2 may help to decipher the environment of the lake during this interval and to understand the dynamics of the lake system and its response to changing climate. The substantially different overall assemblage, including blooms of delicate planktonic taxa, generally favoring higher nutrient status and not observed in substantial quantities in the Holocene, suggests that this trend is more than a preservation effect or lack of dilution from other inputs. More likely, the environment in the lake was similarly favorable for diatom production compared to the recent lake. Yet other proxies, such as pollen, sediment facies, and diatom isotopes suggest at least 4 °C colder summer air temperatures compared to modern conditions (Melles et al., 2012; Chapligin et al., 2012), and in other cold stages this diatom response is absent.

Periodic circulation of a mostly ice-covered lake is one hypothesis to generate such production and diversity. Periodic expansion in plankton diversity may reflect adaptation of the dominant species to a more limited range of conditions or periodic changes in nutrient resource distribution (e.g., Kilham et al., 1996). The general dominance of Cyclotella or Pliocaenicus may represent a competitive exclusion of other taxa during the typical range of conditions (e.g., Stoermer and Edlund, 1999). The particular occurrence of these nutrient-favoring phytoplankton species during MIS 2 may indicate conditions providing seasonal to extra-seasonal pulses of nutrients to the plankton, allowing for a temporary increase in the population of these widespread taxa. Circulation-induced seasonal nutrient input timed to persistent transparent ice cover may also promote the expansion of these otherwise sporadically occurring taxa. Such a scenario may also be consistent with the lack of significant Pliocaenicus, perhaps requiring more extensive open-water conditions.

Other observations from the MIS 2 sediment record provide additional perspective on the lake’s response to this event. Melles et al. (2012) suggest that the sediment facies indicates more extensive ice cover leading to oxygen-depleted
bottom waters, but sufficient melting of the lake’s margins to allow some fluvial input. However, this sediment Facies A does not always correspond to the diatom response observed in MIS 2. Melles et al. (2007) attribute high total organic carbon in MIS 2 to some combination of increased productivity and preservation due to anoxia. Investigations of biomarkers and organic matter δ13C indicate limited connection to atmospheric CO2 and limited oxygen depletion in the water column (Holland et al., 2013) and, although not indicating enhanced aquatic production or nutrients, imply that enhanced production would need to be sustained by internal biogeochemical cycles.

The periphyton assemblage also changes during this same interval, suggesting environmental changes across all diatom habitats. Their abundance suggests at least periodic melting sufficient in the lake margins to promote the diversity and abundance of the lake’s periphyton community. The periphyton may respond directly to climate-induced shallow-water habitat changes, or indirectly to suggested nutrient conditions and lowered lake level during MIS 2 (Juschus et al., 2011). However, the uncertain tolerance ranges of the dominant El’gygytgyn periphyton make a specific interpretation of these changes difficult.

Additional short productive intervals (sensu MIS 2) lacking the characteristic interglacial periphyton suggest the potential recurrence of conditions favorable to diatoms during some cold events. Similar to MIS 2, several of these intervals correspond to low insolation and heavy marine oxygen isotopes in the tuned chronology. Thus, interpretations of biogenic silica and diatom concentration in the deeper record may be complicated by additional such events, requiring confirmation with other proxy records.

4.3 Sparse plankton events

Events in which plankton dramatically decreases in abundance and subsequently re-emerges with a different character may help to constrain the nature of these intervals of the lake’s history. These observations imply that phytoplankton in the lake is stressed such that competitive exclusion is eliminated for some time. However, the recurrence of species either unique to El’gygytgyn or only narrowly distributed over the long history of the lake suggests that minimal populations are sustained in the lake, and that they survive these events. Given the correlation of these events to relatively cold intervals, these extreme declines in the lake’s plankton are most easily explained by a prolonged light limitation from extended summer ice and snow cover on the lake. However, the sparse plankton event at 1060 ka corresponds to a zone of near-modern pollen-derived summer temperature and mean annual precipitation (Melles et al., 2012). Thus, any climate-related mechanism to drive these events is either consistent with such conditions or is short lived and unsolvable in the available pollen record. The role of changing lake level in these events cannot be excluded based on the diatom observations. The persistence of well-preserved near-typical periphyton concentration across some of these intervals suggests some shallow-water habitats are maintained during these times.

The low-plankton events are not evenly distributed through the lake record. None has yet been observed from 225 to 550 ka in the low-resolution sampling completed, and none is anticipated based on the apparent stability of the dominant Cyclotella. The interval 680–920 ka also lacks low-plankton zones with a consistent dominance of Pliocenoicus. These intervals both correspond to a time of lower-amplitude summer insolation variability (e.g., Laskar et al., 2004), suggesting that these events may be triggered by conditions unique to summer insolation lows. This observation is similar to the interval of sustained high diatom concentration in the Lake Baikal record 370–580 ka (Prokopenko et al., 2002). The lake system and the regional climate are sensitive to intervals of extreme insolation lows, perhaps inducing extended periods of light limitation in planktonic habitats. Careful observations of diatom assemblages provide clues to the nature of exceptional cold events only subtly expressed in other proxies.

5 Conclusions and future work

From 0 to 1.2 Ma, sediments recovered from Lake El’gygytgyn record dramatic variations in diatom species assemblage and abundance. During warm interglacial intervals, both plankton and periphyton abundance increases and plankton diversity expands. Extreme warm events (MIS 11 and MIS 31) have the highest diatom concentrations observed, and abrupt plankton assemblage changes within these intervals suggest a threshold in climate-driven nutrient conditions in the lake. Exceptional cold events where plankton abundance and diversity exceed some interglacials suggest the possibility of enhanced sub-ice production promoted by periodic lake circulation and transparent ice. During some cold events, short intervals occur with extremely low plankton abundance and accumulation rates, sometimes corresponding to a shift in the dominant planktongenus. These events are absent from intervals of lower magnitude summer insolation variation, especially 225–550 ka, suggesting a similar response seen in the Lake Baikal record.

Additional diatom studies from this interval and the complete sediment record extending to 3.6 Ma provide numerous opportunities to further test the interpretation of preliminary results presented here. Systematic observations of plankton size and preservation, and periphyton assemblages across these events, may help to distinguish open water from sub-ice production. The patterns of diatom response may also change during earlier intervals of anticipated extreme and continuous warmth. The El’gygytgyn diatom record further provides a unique setting to observe diatom evolution linked to changing climate in this relatively isolated environment.
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