

Complex responses of phototrophic communities to climate warming during the Holocene of northeastern Ontario, Canada

Elmslie, B. G., Gushulak, C. A. C., Boreux, M. P., Lamoureux, S. F., Leavitt, P. R., & Cumming, B. F. (2019). Complex responses of phototrophic communities to climate warming during the Holocene of northeastern Ontario, Canada. *Holocene*. Advance online publication. https://doi.org/10.1177/0959683619883014

Published in: Holocene

Document Version: Peer reviewed version

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1 2 3 4 5 6	Complex responses of phototrophic communities to climate warming during the Holocene of northeastern Ontario, Canada
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9 10	Brett G. Elmslie ¹ , Cale A.C. Gushulak ¹ , Maxime P. Boreux ^{1,2} , Scott F. Lamoureux ² , Peter R. Leavitt ^{3,4} , Brian F. Cumming ¹
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13 14 15 16 17	 Paleoecological Environmental Assessment and Research Laboratory (PEARL), Department of Biology, Queen's University, Kingston, Canada Department of Geography and Planning, Queen's University, Kingston, Canada Institute of Environmental Change and Society, University of Regina, Regina, Canada Institute of Global Food Security, Queen's University Belfast, Belfast, United Kingdom
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19	
20	Corresponding author: C.A.C. Gushulak
21	Email address: 16cacg@queensu.ca
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31 Abstract

32 Although the Holocene climate of North America has been generally well-studied, high regional 33 variability can obscure understanding of mechanisms underlying large-scale and long-term 34 variability. To address such a shortcoming, historical changes in Holocene climate in 35 northeastern Ontario were quantified using analysis of sedimentary pollen, diatoms, and 36 pigments in a small boreal lake. Modern Analog Technique (MAT) based on pollen was used to 37 reconstruct average temperature over the Holocene record of Charland Lake and showed average 38 temperature was ~2 °C warmer than present conditions ~7800–4500 cal yr BP, a time period 39 which we define as the Holocene Thermal Maximum (HTM). Further investigation of pollen 40 data suggests a two-phase HTM: warm and dry conditions followed by warm and wet conditions 41 based on the occurrence of *P. strobus* and Cupressaceae (*Thuja*) pollen. Concomitant changes in 42 landscape features affected the sedimentological and phycological nature of the lake, as diatom 43 assemblages reflect delayed stabilization of the landscape following the draining of Glacial Lake 44 Ojibway ~8200 cal yr BP. In contrast, during a Post-HTM period, which was warmer than today 45 but cooler than the HTM, diatom and pigment data appear to be more influenced by indirect 46 effects of climate, such as lake-water thermal stratification and changes in plankton seasonality. 47 This study suggests that while climate greatly influenced regional forest ecology, landscape, 48 hydrological, and indirect climate effects were more influential on the phytoplankton 49 communities during the Holocene in northeast Ontario. These findings may provide insight into how freshwater environments will respond to future anthropogenic warming in the region. 50 51 52 Keywords: *Thuja*, *pollen*, thermal stratification, landscape dynamics, diatoms, sedimentary 53 pigments

- 54
- 55 Introduction
- 56

57 The Holocene Thermal Maximum (HTM) was a period of enhanced warmth that 58 occurred during the early and middle Holocene (Viau et al., 2006; Renssen et al., 2009, 2012). 59 This warming has been well recorded and well-studied in the mid-to-high latitudes of the 60 Northern Hemisphere that experienced major climatic alterations following the last glacial 61 maximum (LGM) and deglaciation (Whitlock and Bartlein, 1997; Carlson et al., 2008). The 62 cause of the HTM has largely been attributed to an increase in solar insolation, due to orbital 63 variation, which maximized at ~10 ka BP (thousand years before present) (Ritchie et al., 1983; 64 Bond et al., 2001). Despite the overall maximum solar insolation occurring at this time, the 65 specific regions of North America may not have experienced maximum warmth at the same time 66 or to the same degree (Renssen et al., 2009, 2012). In general, regions at higher latitudes 67 experienced greater amounts of warming than those at lower latitudes (Renssen et al. 2009, 2012; 68 Briner et al., 2016) and the western portion of the continent warmed earlier than the east, likely 69 due to downwind cooling effects of the melting Laurentide Ice Sheet (LIS) (Carlson et al., 2008; 70 Viau and Gajewski, 2009). Alongside the variation in onset, duration, and magnitude of warmth 71 experienced during the HTM in North America, reconstructions of Holocene precipitation levels 72 suggest a climatic dipole in which the west of the continent was dry while the east was wet 73 during the HTM (Shuman and Marsicek, 2016). To date, however, these studies have focused on 74 the continental United States and the Arctic or subarctic regions (Viau and Gajewski, 2009; Briner et al., 2016; Shuman and Marsicek, 2016) and relatively little is known regarding the 75 76 nature of the HTM in the boreal region of east-central Canada (e.g., northern Ontario). 77 The boreal forest of northern Ontario spans ~ 1500 km, is one of the largest continuous 78 forests globally, and contains abundant fresh water in its wetlands, rivers, and innumerable small 79 lakes. Despite its size, the region is susceptible to anthropogenically-forced climate change

80 (Price et al., 2013). Therefore, a characterization of landscape responses to analogs of past 81 climates may be useful for predicting effects of atmospheric warming and changes in regional 82 hydrology (Klemm et al., 2016; Navarro et al., 2018; Teller et al., 2018). Atmosphere-ocean-83 vegetation models have suggested that the HTM in northern Ontario was $\sim 2-3$ °C warmer than 84 the modern climate (Renssen et al., 2009, 2012). Recent work from northwest Ontario (~1200 85 km to the west of our study site) reveals that the HTM occurred from ~8500–4500 cal yr BP 86 (calendar years before present) and was ~2 °C warmer than present conditions. This estimate was 87 calculated using a modern analog technique (MAT) calibration based on fossil pollen and a 88 regional set of modern pollen and climate data (Moos and Cumming, 2011). Associated dryer 89 conditions resulted in lower lake levels, increased algal production, an eastward shift of the 90 prairie-forest ecotone, and an increase in fire activity (Moos et al., 2009; Moos and Cumming, 91 2011, 2012; Karmakar et al., 2015a, 2015b). Despite this research, there has been little work 92 examining the HTM of northeastern Ontario, Canada, which is a problem this paper seeks to 93 address.

94 Only one Holocene-scale palynological study has documented Holocene climate 95 variability in northeast Ontario (Liu, 1990). This study examined three sediment cores in a north-96 south transect of eastern Ontario. One of these sites, Lake Six, is 6 km away from our study site 97 so comparisons to Liu (1990) refer to the Lake Six record. In Liu (1990), the author showed a 98 delayed, but protracted HTM period occurring from ~7-2.5 ka BP at Lake Six. This was defined 99 by increases in *Pinus strobus*, a typical taxon of the warm Great Lakes-St. Lawrence forest, and 100 in Cupressaceae. Liu (1990) interpreted the presence of Cupressaceae pollen as belonging to 101 Thuja based on its co-occurrence with Thuja macrofossils within the sediment core. These data suggest that the HTM of northeast Ontario was wet, agreeing with studies at similar longitudes to 102

103 the south (Viau et al., 2006; Shuman and Marsicek, 2016). A similar interpretation has been 104 made by Carcaillet et al. (2001) who used charcoal analysis to infer that low fire activity during 105 the this time in western Ouebec was due to enhanced water availability. Unfortunately, Liu's 106 (1990) climatic results from Lake Six are supported by only three radiocarbon dates. With these 107 few data points there is likely considerable error regarding the temporal boundaries of HTM 108 warming and it is likely that the upper boundary (~ 2.5 ka BP) is overestimated based on studies 109 from northwest Ontario (Moos and Cumming, 2011, 2012). Improved geochronology, as well as 110 replicate sites in the region, are important to better define the onset and duration of the HTM in 111 northeast Ontario, as well as its effects on lake ecosystems.

112 This paper attempts to quantify changes in climate systems in northeastern Ontario using 113 robust analysis of fossil pollen to reconstruct climatic and vegetation characteristics, as well as 114 fossil diatoms and pigments from aquatic phototrophs to quantify the effects of climate 115 variability on lake production and community composition. Diatoms are ideal for these 116 objectives, as certain species have well-documented optima for nutrients and other physical and 117 chemical limnological characteristics (water-depth, pH, salinity, stratification regime, etc.). By 118 examining changes in diatom species which are known to be nutrient-controlled, changes in 119 nutrient levels can be inferred (Douglas and Smol, 1999; Lotter et al., 1999; Rühland and Smol, 120 2005; Cumming et al., 2015). Similarly, fossil pigments (chlorophylls, carotenoids, derivatives) 121 often preserve after the loss of morphological remains of non-siliceous phytoplankton and 122 phytobenthos, and are used to estimate historical changes in the abundance of primary producers, 123 as well as their gross community composition (Leavitt et al., 1994a, 1997; Hodgson et al., 1998; 124 Hall et al., 1999). Taken together, analysis of these proxies are can be used to answer three 125 questions related to the HTM: a) when and for how long did the HTM take place in northeast

Ontario; b) what was the degree of warming that occurred during the HTM of this region and was the warmer climate wet or dry, and; c) what was the response in algal production to this warmer climate?

129

130 Methods 131 132 Study area 133 134 Charland Lake (N 48°34' 50.8", W 80°53' 46.5) (Figure 1) is located east of Timmins, 135 Ontario, Canada, immediately west of Kettle Lake Provincial Park and is located within 6 km of 136 Lake Six, site of the only other study of Holocene climate in northeast Ontario (Liu, 1990). 137 Charland Lake, like other basins in the region, formed when Glacial Lake Ojibway drained off 138 the landscape of northeastern Ontario ~8200 cal yr BP (Veillette, 1994; Carlson et al., 2008; Roy 139 et al., 2015; Margold et al., 2018). The lake is small (~20 ha), 16 m deep, and 274 m above sea 140 level. The climate of the region is humid continental (Koppen Dfb) which is characterized by wet 141 summers and long, cold winters. Average temperature is ~ 1.8 °C (Environment Canada, 142 2019). Modern boreal forest in the area is composed mainly of white (*Picea glauca*) and black 143 spruce (*Picea mariana*) in lowlands with jack pine (*Pinus banksiana*) and white birch (*Betula* 144 papyrifera) on upland sites. Charland Lake occurs in the 'Great Clay Belt' of northern Ontario. 145 The bedrock is composed of metamorphosed Precambrian granites and granodiorites which are 146 overlain by Quaternary glacial landforms and extensive glaciolacustrine clay deposits laid down 147 by Glacial Lake Ojibway (Veillette, 1994; Roy et al, 2011, 2015). [insert Figure 1.] 148 149 Figure 1. (A) Bathymetric map of Charland Lake with 2 m contours. The black star represents

150 the location of the piston core that was taken from a depth of ~16 m. (B) A reference map of

151 Ontario with the Charland Lake coring site indicated by a black star.

152

153 Sample collection

154 Bathymetric data points were collected from Charland Lake using a Garmin GPS-Map 155 Sounder 238 connected to a Toughbook computer with Bathymetric Automated Survey System 156 (B.A.S.S.) software, v 2.4 (Levec, 2001). These data points were converted to shape files and 157 used to produce the bathymetric map in ArcMap 10.5 (Esri, 2016). A 473 cm long sediment core 158 was collected from a depth of ~16 m on 19 June 2014 using a 1-m square-rod Livingstone piston 159 corer with an internal diameter of 5.1 cm (Wright, 1967; Wright et al. 1984; Glew et al., 2001). 160 Approximately 10 cm of material from the top of the core was lost during horizontal extrusion in 161 the field. To assure collection of an undisturbed sediment-water interface, and to be able to 162 accurately date the top of the piston core, a gravity core (internal diameter 7.62 cm) was taken at 163 the same site using a modified gravity corer (Glew 1989) and was sectioned into 0.5 cm 164 intervals in the field. The piston core exhibited distinct units, with organic gyttja 0–313 cm, grey 165 clays from 313–337 cm, organic-rich material from 337–417 cm, and basal clay deposition 166 below that level. In this paper, we refer to these clay layers in order of deposition, making the 167 clay deposit from 417–473 cm 'the first clay layer' and the deposit from 313–337 cm 'the second 168 clay layer'. The piston core was wrapped in 1-m sections on site and transported in a cooler to 169 the Paleoecological Environmental Assessment & Research Laboratory (PEARL) at Queen's 170 University where they were stored horizontally in a cold room at ~4° C. Half of the core was 171 then sectioned into 1 cm intervals, while the other half was archived.

172

173 Analyses

174 Chronology

175 176 The top 13 cm of the gravity core were dated by using a constant rate of supply (CRS) model applied to ²¹⁰Pb gamma activity of the sediments. This was done to assess the date of the 177 178 top of the piston core as the sediment-water interface was lost during extrusion. The ¹⁴C 179 chronology for the piston core was determined by dating a concentrated sample of pollen grains 180 from eight sediment samples with accelerator mass spectrometry (AMS). Pollen grains were 181 isolated from lake sediments by LacCore using a procedure similar to Brown et al. (1989) and 182 measured for ¹⁴C at the Lawrence Livermore National Laboratory. An age-depth relationship 183 based on these radioisotopic data was constructed using Bayesian age modeling with the BACON 184 (v. 2.2) modeling package in R (R Core Team, 2015) with the IntCal13¹⁴C calibration curve 185 (Blaauw and Christen, 2011; Reimer et al., 2013). Default settings outlined by Goring et al. 186 (2012) were used to determine the gamma distribution of the accumulation rate, which were in 187 agreement with the posterior distribution of activities (Blaauw and Christen, 2011). The prior 188 memory was set to a mean of 0.3 and shape of 25 to accommodate small shifts in accumulation 189 rates. The sensitivity of prior memory parameters was tested and changes in mean and shape did 190 not result in large changes to the model.

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Dry mass and organic matter
Dry mass and percent organic matter was determined through standard loss-on-ignition
procedures (Heiri et al., 2001) on 60 intervals throughout the sediment core.
Pollen
Pollen samples were prepared at every 8 cm (60 samples total) using a modification of
the method of Bennett and Willis (2001). Sediment samples were spiked with two exotic
Lycopodium tablets (batch no. 1031) and digested with 10% hydrochloric acid and 10%
potassium hydroxide before being sieved through a 10 μ m mesh. The remaining sediment was
further digested in 40% hydrofluoric acid and then acetolyzed, deflocculated with 10% sodium
metaphosphate, stained with Safranin, and mounted on microscope slides in silicon oil. Pollen
grains were counted with a Leica light microscope with a 40x differential-interference-contrast
objective. A minimum of 400 pollen grains were counted per sample except those in which
pollen concentration was low and multiple coverslips needed to be counted. In those cases, a
minimum of 300 grains were counted. Pollen grains were identified to the lowest possible
taxonomic resolution based on published references (Bassett et al., 1978; McAndrews et al.,
1978; Kapp et al., 2000).
Pigments
Subsamples of wet sediment were used for determination of photosynthetic pigment

215 concentrations from 59 intervals throughout the core at the University of Regina's Institute of 216 Environmental Change and Society (IECS). Sedimentary pigment analysis was undertaken 217 following procedures outlined in Leavitt and Hodgson (2001). Pigment concentrations are reported as nmoles pigment g⁻¹ gram organic matter, a unit which is linearly proportional to the 218 219 standing stock of phytoplankton in decadal-scale monitoring programs (Leavitt et al., 1994a). An 220 Agilent model 1100 high-performance liquid chromatography (HPLC) system with photodiode 221 array detector was calibrated using commercial pigment standards from DHI (Denmark). 222 Analysis included the main a- and b-phorbins (chlorophyll, pheophytin), as well as chemically-223 stable, taxonomically-diagnostic pigments representing total algal abundance (β -carotene), 224 cryptophytes (alloxanthin), total cyanobacterial (echinenone), colonial cyanobacteria 225 (myxoxanthophyll), Nostocales cyanobacteria (canthaxanthin), chlorophytes (Chl b), 226 chlorophytes + cyanobacteria (lutein-zeaxanthin), siliceous algae (fucoxanthin), mainly diatoms 227 (diatoxanthin), and anaerobic purple sulfur bacteria (okenone) following Leavitt and Hodgson, 228 (2001). Historical changes in the lacustrine preservation environment were recorded as changes 229 in the ratio of labile precursor (Chl a) to stable products (pheophytin a), both ubiquitous 230 pigments used to estimate total phototrophic abundance (Leavitt and Hodgson, 2001). 231

232 Diatoms

Diatom samples were prepared using ~0.2–0.3 g of wet sediment, taken every 8.0 cm, for a total of 59 subsamples. Samples were digested in a 1:1 molar solution of concentrated nitric and sulphuric acids before being brought to a slightly acidic pH through repeated rinses with double-deionized water. Diatom slurries were reduced to a volume of ~5 ml by aspiration following sedimentation, and spiked with a solution of microspheres of known concentration

equivalent to 4 ml of a 2.0×10^7 spheres/ml solution for organic samples and 0.2 ml of a 2.0×10^7 238 239 10⁷ for clay-rich samples. Samples were plated on coverslips in a series of four dilutions and then 240 mounted to slides using Naphrax[®]. Diatoms valves were identified and enumerated using a Leica 241 DMRB microscope under a $100 \times$ Fluotar objective (NA of objective = 1.3) and differential 242 interference contrast optics at 1000× magnification. For most diatom samples a minimum of 400 243 valves were counted, or, if the concentration of valves was exceptionally low, until five transects 244 were completed. Diatoms were identified to species or lower taxonomic units using the standard 245 references (Krammer and Lange-Bertalot, 1986, 1988, 1991a, 1991b; Cumming et al., 1995). 246 Chryosphyte scales were enumerated alongside diatoms, but were not taxonomically identified.

247

248 Numerical analyses

249 Fossil pollen, pigment, and diatom data were plotted using the computer program Tilia 250 v.2.0.2 (Grimm, 2004). Pollen taxa present at greater than 2% abundance in at least two samples 251 and diatom species with greater than 5% abundance in at least three samples were included in the 252 plots. Pollen concentration was calculated with the formula: (exotic Lycopodium spores added x 253 fossil pollen counted) / exotic Lycopodium spores counted (Bennett and Willis, 2001) and 254 standardized to dry mass. A depth-constrained cluster analysis (CONISS) (Grimm, 1987) was 255 performed on the pollen assemblage to identify major differences in the pollen assemblage. The 256 statistical significance of zones delineated by CONISS were validated with a broken-stick model 257 by rioja package in R (Juggins, 2015; R Core Team, 2015).

Modern Analog Technique (MAT) was used to quantitatively reconstruct mean annual temperature over the Holocene record of Charland Lake using the C2 software program (Juggins, 2003). Modern pollen taxa and climate data were collected from the North American pollen

261 database (Whitmore et al., 2005; Williams et al., 2006) to form a regional calibration set. A total 262 of 305 samples between 45-60 °N and 75-90 °W were selected to form the calibration set which 263 captured the transition between the boreal and mixed-wood forests to the south of Charland 264 Lake. The fossil pollen from Charland Lake are well represented in the modern pollen dataset so 265 it is likely that this group of sites should provide strong analogs for the assemblages found in the 266 Charland Lake sediment core over the Holocene. Cupressaceae pollen was removed from the 267 reconstruction model as this pollen type can represent both Juniperus and Thuja which have very 268 different climatic preferences (Yu, 1997). Annual average temperature was reconstructed using 269 the five closest analogs compared to fossil pollen interval with squared-chord measurement of 270 similarity (Overpeck et al., 1985; Viau et al., 2006; Viau and Gawjewski, 2009) and bootstrap 271 cross-validation (bootstrapped $r^2 = 0.95$, root-mean-square-error-of-prediction (RMSEP) = 0.88). 272 The 305 site calibration set was tested against itself using the same method described above to 273 discern the distribution of dissimilarities among the analogs used for the temperature 274 reconstructions. Each of the five closest analogs used for each fossil interval was within the 20th percentile of the 305 sites. The majority were within the 10th percentile. This resulted in average 275 dissimilarity values to be less than 30% which corresponds to the top 10th percentile for each 276 277 reconstructed site.

Pigments were restricted to common sedimentary compounds of known chemical stability (see above). Analysis of similarities (ANOSIM) using a Bray-Curtis similarity coefficient and 999 permutations, were used to test the null hypothesis that there was no difference in diatom and pigment composition between the pollen-inferred climate zones (Clarke and Warwick, 1994). Analyses were performed using both non-transformed and square-root transformed species data. ANOSIM tests were performed on diatom relative abundance and

284	concentration and pigment concentration data. ANOSIM tests were calculated using the PAST 3
285	software package (Hammer et al., 2001; Hammer and Harper, 2006). The sample from a depth of
286	440 cm was determined to be an outlier and was removed prior to any statistical analyses.
287	Following the ANOSIM analysis, SIMPER tests (also performed in PAST), were used to
288	calculate the contribution of each species to the average dissimilarity between the two groups.
289	Post-hoc t-tests assuming unequal variance were performed to identify the significant difference
290	between the pollen-inferred climate zones for individual sedimentary pigments using Microsoft
291	Excel. The index of chrysophyte scales to diatom frustules (scale-to-diatom index) was
292	calculated using the formula: scales/ (diatoms + scales) \times 100 (Moos et al., 2005).
293	
294	Results
293 296 207	Age model
297 298	The activity of ²¹⁰ Pb in the Charland Lake gravity core was adequate to produce an age-
299	depth model for the top 12 cm of sedimentation (Supplemental Figure 1). Background levels of
300	²¹⁰ Pb were reached below that depth. Approximately 10 cm were lost from the first section of the
301	piston core during extrusion. Based on our CRS model , this would mean the top of the piston
302	core would correspond to an age of 0 cal yr BP with $BP = 1950$.
303	Concentrated pollen samples from eight sediment intervals were dated using AMS (Table
304	1). Bayesian modeling of the depth- ¹⁴ C age relationship revealed a relatively constant rate of
304 305	1). Bayesian modeling of the depth- ¹⁴ C age relationship revealed a relatively constant rate of deposition over the period of study (Figure 2). Overall, the errors of the inferred ages for
304305306	1). Bayesian modeling of the depth- ¹⁴ C age relationship revealed a relatively constant rate of deposition over the period of study (Figure 2). Overall, the errors of the inferred ages for intervals in the sediment core were relatively low (± 30 –40 years). The calibrated ¹⁴ C dates
304305306307	1). Bayesian modeling of the depth- ¹⁴ C age relationship revealed a relatively constant rate of deposition over the period of study (Figure 2). Overall, the errors of the inferred ages for intervals in the sediment core were relatively low (\pm 30–40 years). The calibrated ¹⁴ C dates increased in age with increasing depth and follows an approximately linear trend with
 304 305 306 307 208 	1). Bayesian modeling of the depth- ¹⁴ C age relationship revealed a relatively constant rate of deposition over the period of study (Figure 2). Overall, the errors of the inferred ages for intervals in the sediment core were relatively low (\pm 30–40 years). The calibrated ¹⁴ C dates increased in age with increasing depth and follows an approximately linear trend with

- 309 *Table 1.* Summary of the ¹⁴C-dating results on pollen isolated from selected intervals from the
- 310 sediment cores from Charland Lake. All analyses were performed based on pollen isolated at the
- 311 LacCore Facility at the University of Minnesota, and dated at Lawrence Livermore National
- Laboratory. The mean age of the distribution is presented in "Cal yr BP". The top date was
- 313 generated through ²¹⁰Pb dating and a CRS model.
- 314

Core	Cumulative depth in piston core (cm)	$\delta^{13}C$	¹⁴ C Age ± SD	Cal yr BP	2-σ cal yr BP range (min-max)	CAMS #
C2S1	0	N/A	N/A	0	-6–28	N/A
C2S1	45.5	-28	1170 ± 30	1081	976–1187	171192
C2S2	143.5	-28	2505 ± 40	2623	2451-2749	171205
C2S2	178.5	-28	3035 ± 30	3260	3133–3360	171201
C2S3	204-206	-28	3480 ± 30	3762	3649–3863	176019
C2S3	262.5	-28	4400 ± 30	4925	4846–5037	171194
C2S4	305-306	-28	4560 ± 35	5390	5238-5570	176020
C2S4	401.5	-28	6235 ± 30	7159	6999–7272	171206
C2S5	437.5	-28	6995 ± 40	7815	7670–7947	171202

316

Figure 2. Age-depth model run for the Charland Lake core using BACON (Version 2.2). Top left
panel: Markov Chain Monte Carlo model iterations. Top middle panel: Prior (heavy line) and
posterior (solid) distribution of accumulation rate. Top right panel: Prior (heavy line) and
posterior (solid) distribution of model memory. Bottom panel: calibrated ¹⁴C dates and the agedepth model. The outer dotted lines indicate 95% confidence intervals. The central dotted line is
the 'best' model based on the weighted mean age.

324 Pollen

325

326 Four statistically-significant pollen zones were identified in the Charland Lake core

327	through CONISS and validated with the broken stick model (Figure 3; Supplemental Figure 2).
328	CONISS analysis demonstrated that breaks between zones occur at cumulative depths of 85, 253,
329	and 445 cm. Reconstructed average temperature showed major changes in line with these
330	boundaries and we labelled the zones as the Pre-HTM, HTM, Post-HTM, and Modern climate
331	zones (Figure 3). Standard error for the temperature reconstructions were generally low and on
332	average did not exceed 0.5 °C above or below the estimate (Figure 4). Error is larger in the Pre-
333	HTM zone, but even considering the upward limit of the error there is still a large and noticeable
334	increase in temperature when entering the HTM zone (Figure 4).
335	The Pre-HTM zone occurred from 473–445 cm (~8200–7800 cal yr BP) and showed
336	mixed dominance of Picea and Pinus; P. bankisana and P. resinosa could not be distinguished.
337	This zone also included a relatively low abundance of arboreal pollen compared to rest of the
338	core, as well as more abundant spores and pollen from Sphagnum, fern, and Cyperaceae.
339	Reconstructed average temperature was approximately -0.5–0 $^\circ$ C in this zone. The HTM zone
340	occurred from 445–253 cm (~7800–4500 cal yr BP) and showed a major increase in
341	reconstructed temperature (Figure 3, Figure 4). Average temperatures of $\sim 2-3$ °C were consistent
342	across this zone, but there appears to be two distinct floral assemblages within the warm period.
343	The first was marked by an increase in the P. banksiana/resinosa and the second with an
344	increase in Pinus strobus and Cupressaceae. Organic sedimentation also began in this zone,
345	concomitant with pollen concentration rapidly increasing before falling substantially at the onset
346	of the second clay layer. The Post-HTM period (253-85 cm, ~4500-1700 cal yr BP) showed an
347	increase in Picea, P. banksiana/resinosa, and Betula pollen, as those of P. strobus and
348	Cupressaceae declined. This is reflected in the MAT reconstruction as average temperature fell
349	by ~1 °C. Pollen concentration was variable in this zone but generally increased during the

350	earlier portion before declining after ~3300 cal yr BP. The Modern Zone (85–0 cm, ~1700–0 cal
351	yr BP) showed a general decrease in pollen concentration and further decreases of <i>P. strobus</i> .
352	Picea became predominant in this zone with P. banksiana/resinosa and Betula experiencing
353	minor decreases compared to the Post-HTM zone. Reconstructed average temperature slowly
354	declined by ~0.5 °C throughout this zone. The concentration of pollen types was also plotted and
355	can be found in the supplemental materials (Supplemental Figure 3).
356	[insert Figure 3]
357	Figure 3. Relative abundance of abundant (>2% found in 2 sections of the core) pollen taxa in
358	Charland Lake. Total pollen concentration is also shown. Darker zones in Sediment Type
359	represent organic sedimentation in the piston core. Layered zones represent clay deposits.
360	Climate Zones are derived through CONISS. Reconstructed average temperature was estimated
361	through Modern Analog Technique calibration (bootstrapped $r^2 = 0.95$, RMSEP = 0.88).
362	[insert Figure 4]
363	Figure 4. Reconstructed average temperature with error bars and percent dissimilarity of the
364	analogs used in the MAT temperature reconstruction of the Charland Lake record. Percent
365	dissimilarity refers to the mean dissimilarity of the five closest analogs used to reconstruct
366	annual temperature for each fossil interval. Zones are the CONISS-derived climate zones from
367	Figure 3.
368	
369	
370	Pigments
372	The pollen climate zones were used as a framework to assess the relationship between
373	climate change and variation in pigment assemblages (Figure 5). Overall, preservation of

374 pigments was poor in the Pre-HTM zone, with fossil concentrations below detection limit in 375 most samples. A similar absence of fossil pigments in the second clay layer suggests that the 376 near absence of sedimentary organic matter favored complete decomposition of carotenoids and 377 chlorophylls, irrespective of their inherent chemical lability, similar to patterns seen in glacially-378 fed alpine lakes (Bunting et al., 2010). In contrast, concentrations of ubiquitous pigments (β -379 carotene, Chl a, pheophytin a), and those from cryptophytes (alloxanthin), diatoms 380 (diatoxanthin), total cyanobacteria (echinenone), Nostocales cyanobacteria (canthaxanthin), 381 chlorophytes (pheophytin b) and chlorophytes+cyanobacteria (lutein-zeaxanthin) all increased 382 markedly at the base of the HTM zone, whereas more labile compounds (fuxoxanthin, Chl b) did 383 not. These patterns reflect the increase in okenone from obligate anaerobic purple sulfur bacteria, 384 an indicator of anoxia in deep-waters or sediment. Transition from the HTM to Post-HTM zone 385 was marked by significant increases in the concentration of most fossil pigments (Table 2, Figure 386 6). The Post-HTM zone was characterized by elevated concentrations of most pigments, often to 387 a historical maximum relative to other zones (e.g., β -carotene, lutein-zeaxanthin, okenone, and 388 canthaxanthin). In the Modern zone (0-85 cm; ~1700-0 cal yr BP), pigment assemblages were 389 marked by the first appearance of myxoxanthophyll from colonial cyanobacteria, increases in 390 concentrations of compounds from Nostocales cyanobacteria (canthaxanthin) and secondarily 391 total cyanobacteria (echinenone), and historical maxima of less chemically-stable pigments from 392 diatoms (fucoxanthin, diatoxanthin) and chlorophytes (Chl b). In contrast, fossil levels of 393 ubiquitous β -carotene and the mixed chlorophyte-cyanobacterial indicator, lutein-zeaxanthin, 394 declined slightly in the most recent zone. Relative to individual pigments, the ratio of labile 395 chlorophyll a to stable phaeophytin a (indicating the preservation environment) remained

396 relatively constant through the core, with some variance within and between the clay bands of

the Pre-HTM and HTM zones (Figure 5). [insert Figure 5.]

398

399 Figure 5. Concentration of photosynthetic pigments (nmoles per gram organic matter) in

400 Charland Lake sediment core over time (cal yr BP). The ratio of chlorophyll *a* to phaeophytin *a*,

401 an indicator of preservation, and organic matter (%) are also shown. Darker zones in Sediment

402 Type represent organic sedimentation in the piston core. Layered zones represent clay deposits.

403 The dotted lines indicate the pollen-derived climate zones from Figure 3.

404

Table 2. Summary table of the results from a series of two sample post hoc t-tests assuming
 unequal variance conducted to identify significant differences in pigment concentrations between
 the HTM, Post-HTM, and Modern zones for sedimentary pigments. Bold cells indicate
 significance.

409

Zones	Canthaxanthin	Echinenone	Okenone	Alloxanthin	β-carotene
НТМ,	t (22) -8.7 = p =	t (12) -2.5 = p =	t (21) = -6.3 =	t (26) -5.1 = p =	t (21) = -1.9 p
Modern	<0.01	0.03	<i>p</i> = <0.01	<0.01	= 0.07
HTM, Post-	t (44) -5.0 = <i>p</i> =	t (43) = -4.5 p =	t (42) = -4.2 =	t (31) -4.2 = p =	<i>t</i> (43) = -4.1 <i>p</i>
HTM	<0.01	<0.01	<i>p</i> = <0.01	<0.01	= <0.01
Post-HTM,	t (23) -4.2 = p =	t (11) = -0.8 <i>p</i> =	t (25) = -2.2 =	t (32) -1.4 = p =	t (24) = 1.7 p =
HTM	<0.01	0.44	<i>p</i> = 0.04	1.69	0.1
	Lutein-	Myxoxanthopyll	Fucoxanthin	Chlorophyll b	Diatoxanthin
	zeaxathin				
НТМ,	t (32) -2.5 = p =	<i>t</i> (10) -3.3 = <i>p</i> =	t (16) -19.4 = p	t (10) -8.4 = p =	t (25) -10.7 =
Modern	0.02	<0.01	= <0.01	<0.01	<i>p</i> = <0.01
HTM, Post-	t (36) -4.4 = p =	N/A	t (25) -6.3 = p	t (22) -3.1 = p =	t (41) -6.4 = p
HTM	<0.01		= <0.01	<0.01	= <0.01
Post-HTM,	t (25) 2.3 = p =	<i>t</i> (10) -3.3 = <i>p</i> =	t (31) -4.6 = p	t (26) 0.3 = p =	t (19) -6.0 = p
Modern	0.03	<0.01	= <0.01	0.77	= <0.01

⁴¹⁰

411 One-way pair-wise ANOSIM tests confirmed that there were significant differences in

412 pigment concentration among the three most recent zones. Specifically, the null hypothesis of no

413 difference between pigment concentrations among zones was rejected in comparisons of the

414 HTM and Post-HTM zones, and between the Post-HTM and Modern zones (Table 3). A

415 SIMPER test identified lutein-zeaxanthin as the predominant pigment contributing to ~40% of

416 the difference in composition between the HTM and Post-HTM zones (Table 4), and ~29% of

417 the difference in composition seen between Post-HTM and Modern zones (Table 4). [insert

- 418 Figure 6].
- 419

420 Figure 6. Boxplots of pigment concentrations (nmol/g OM) within the a priori defined pollen-

- 421 derived climate zones (HTM (n= 24), Post-HTM (n= 21), Modern (n= 11)). ANOVA tests were
- 422 run to test for significant differences between zones for A) β -carotene (F(2, 53) = 8.2, p-value <
- 423 0.01) B) lutein-zeaxanthin (F(2, 53) = 7.6, p-value < 0.01) C) alloxanthin (F(2, 53) = 11.4, p-
- 424 value < 0.01) D) echinenone (F(2, 53) = 7.3, p-value < 0.01) E) okenone (F(2, 53) = 10.1, p-
- 425 value < 0.01) F) canthaxanthin (F(2, 53) = 26.8, p-value < 0.01) G) chlorophyll b (2 outliers
- 426 removed in the Post-HTM zone) (F(2, 53) = 6.5, p-value < 0.01) H) fucoxanthin (F(2, 53) = 52.1,
- 427 p-value < 0.01) I) diatoxanthin (F(2, 53) = 47.1, p-value < 0.01) J) myxoxanthophyll (F(2, 53) =
- 428 22.9, p-value < 0.01). Asterisks indicate zones that are significantly different based on post-hoc
- 429 t-tests (Table 2). Pre-HTM zone omitted due to the small number of samples in this zone.
- 430

431 *Table 3* Summary of the one-way ANOSIM pair-wise tests (Bray-Curtis dissimilarity) on

432 Charland Lake diatom relative abundances, diatom concentrations, and HPLC data between the

433 pollen-derived climate zones. Significance levels indicated in brackets. Bold cells indicate

- 434 significance (999 permutations).
- 435

	Diatom	Diatom Relative	Diatom	Diatom	HPLC	HPLC
	Relative Abundance	Abundance (sqrt)	Concentration	Concentration (sort)	concentration	concentration (sort)
Global R	0.25 (0.001)	0.28 (0.001)	0.23 (0.001)	0.24 (0.001)	0.23 (0.001)	0.31 (0.001)
Modern, Post-HTM	0.02 (0.294)	0.08 (0.12)	0.21 (0.008)	0.27 (0.003)	0.39 (0.001)	0.58 (0.001)

Modern, HTM	0.35 (0.002)	0.37 (0.001)	0.3 (0.002)	0.27 (0.002)	0.12 (0.1)	0.21 (0.017)
Post-	0.34 (0.001)	0.33 (0.001)	0.22 (0.001)	0.24 (0.001)	0.21 (0.001)	0.27 (0.001)
нтм,						
нтм						

Table 4. Summary of the one-way SIMPER tests (Bray-Curtis dissimilarity) on Charland Lake

diatom relative abundances, diatom concentrations, and HPLC data between pollen-derived

climate zones. Percent contribution of driving taxa indicated in brackets.

	Diatom	Diatom	Diatom	Diatom	HPLC	HPLC
	Relative	Relative	Concentration	Concentration	concentration	concentration
	Abundance	Abundance		(sqrt)		(sqrt)
		(sqrt)				
Modern, Post-	S. minutulus	D. stelligera	S. minutulus	S. minutulus	Lutein (29.0%)	Myxoxanthophy
HTM	(32.8%)	(18.8%)	(46.2%)	(26.7%)		(17.4%)
Modern, HTM	S. minutulus	S. minutulus	S. minutulus	S. minutulus	Lutein (27.9%)	Diatoxanthin
	(29.7%)	(17.1%)	(46.0%)	(26.3%)		(17.4%)
Post-HTM,	S. minutulus	S. minutulus	S. minutulus	D. stelligera	Lutein (40.2%)	Lutein (22.3%)
НТМ	(22.8%)	(14.6%)	(27.7%)	(17.8%)		

Diatoms

445	Application of pollen climate zones to the diatom assemblages of Charland Lake showed
446	that diatom concentrations were low in both the Pre-HTM zone and the second clay band, but
447	increased in the organic layer between the clay bands during the HTM to $\sim 23 \times 10^8$ valves per
448	gram dry weight (Figure 7). Diatom concentrations also increased after the second clay band and
449	remained relatively stable between 284–145 cm (range ~9 to 21 x 10^8 valves per gram dry
450	weight). The scale-to-diatom index was low within both clay bands, with an increase in scaled
451	chrysophyte abundance in the intervening organic layer. The abundance of scaled chrysophytes
452	increases during the second clay band and continued to increase throughout the HTM and Post-
453	HTM zones. After a Post-HTM zone peak, scaled chrysophyte abundance declined into the
454	Modern zone and remained low to the top of the core (Figure 7). [insert Figure 7.]

456	Figure 7. Relative abundance of dominant (>5% found in 3 sections of the core) diatom taxa in
457	Charland Lake arranged by age (cal yr BP). The diatom taxa are arranged by their weighted-
458	average optima based on cumulative depth in the core. The scale-to-diatom index and total
459	diatom concentrations (valves/g dry weight x 10 ⁸) are also shown. Darker zones in Sediment
460	Type represent organic sedimentation in the piston core. Layered zones represent clay deposits.
461	The dotted lines indicate the pollen-derived climate zones from Figure 3.
462	
463	The largest shift in diatom assemblages occurred after the second clay band, before the
464	boundary of the HTM and Post-HTM zones. Prior to this point, the Pre-HTM and HTM zones
465	were predominated by benthic taxa, while the Post-HTM and Modern zones were composed
466	mainly of planktonic taxa. Only one sample from the Pre-HTM zone contained enough diatoms
467	for enumeration so this zone was removed from the analysis.
468	HTM taxa prior to the second clay band included Staurosira construens and Staurosirella
469	pinnata. Lindavia intermedia appeared as an important taxon immediately before and after this
470	clay band. After the resumption of organic sedimentation, Stephanodiscus minutulus was the
471	predominant taxon for the remainder of the zone. The importance of S. minutulus increased in
472	the Post-HTM -zone and remained common until \sim 220 cm when relative abundance of
473	Discostella stelligera increased. This pattern was reversed at ~105 cm. Diatom assemblages
474	within the Modern zone were composed largely of S. minutulus and Stephanodiscus parvus;
475	however, D. stelligera abundance increased again at \sim 70 cm until it became a subdominant
476	species at the top of the core (Figure 7).

477	The difference in diatom assemblages between the four pollen-derived climate zones
478	were assessed using ANOSIM tests. The null hypothesis of no difference between diatom
479	assemblages in the HTM, Post-HTM, and Modern zones was rejected in two of the pair-wise
480	tests comparing the HTM zone to the Post-HTM zone and the HTM zone to the Modern zone
481	(Table 3). SIMPER tests identified S. minutulus as the predominant species contributing to $\sim 23\%$
482	of the difference in species composition between the HTM and Post-HTM zones (Table 4).
483	Similarly, there was a significant shift in species composition between the HTM and Modern
484	zones with S. minutulus driving species change and contributing to $\sim 30\%$ of the difference in
485	species composition (Table 4).

487 **Discussion**

488 The HTM in northeast Ontario took place from ~7800–4500 cal yr BP based on large 489 changes in pollen data that suggest elevated temperatures. Average temperature reconstructions 490 from the pollen assemblages in Charland Lake showed that the climate was ~1.5–2.0 °C warmer 491 than the modern day. In this context, the 'modern day' refers to the top of our piston core which 492 corresponds to ~1950. All discussion referring to 'the modern day' or 'present day' relate to this 493 time frame. Algal response to climate change over the Holocene of this region was variable and 494 somewhat unexpected as the warm HTM zone did not experience the greatest amount of algal 495 abundance. Instead, the cooler Post-HTM and Modern zones showed increased algal production 496 which may be related to various indirect climate and landscape effects. These effects, along with 497 more thorough interpretations of pollen and algal data, are discussed in the forthcoming sections. 498

499 Pollen as a climate proxy in northeastern Ontario

500 *Pre-HTM zone* (~8200–7800 *cal yr BP*)

501 The climate during this interval was cooler than present as is shown in the temperature 502 reconstruction (Figure 3). This inference of cooler temperatures arises because of the 503 predominance of *Picea* pollen, combined with a relatively low amount of *Pinus sp.* pollen. 504 Thermophilous taxa, such as deciduous hardwood trees, were rare in the Pre-HTM zone, further 505 contributing to the inference of cool conditions. In addition, the landscape was likely poorly 506 developed with low vegetation cover and immature, highly inorganic soils, as inferred by the 507 relatively high abundances of ferns and other spore producing plants. It is likely that the boreal 508 forest had not completely colonized the watershed of Charland Lake so soon after the draining of 509 Glacial Lake Ojibway ~8200 cal yr BP (Liu, 1990; Prentice et al., 1991).

510

511 HTM zone (~7800–4500 cal yr BP)

Reconstructed temperature shows a sharp increase by ~2 °C at the onset of this zone ~7800 cal yr BP. This climate shift is most driven in the decline of *Picea* and the increase in *P. banksiana/resinosa*, followed by increases in *P. strobus* and Cupressaceae at ~6000 cal yr BP. This is similar to patterns recorded in the mid-Holocene of Lake Six as described by Liu (1990). This two-phase HTM likely represents an increase in precipitation levels after ~6000 cal yr BP as inferred by increase of Cupressaceae.

518 Interpretation of the Cupressaceae peak is difficult, as this pollen morphotype cannot be 519 easily distinguished between *Juniperus* and *Thuja* based on morphological characteristics alone 520 (McAndrews, 1973; Yu, 1997). As *Juniperus* and *Thuja* are indistinguishable as pollen types, 521 and the two genera have differing climatic optima, we have elected to remove Cupressaceae 522 from our temperature reconstructions. It is for this reason as well, that we did not attempt to 523 reconstruct annual precipitation over the Holocene based on the pollen in the Charland Lake 524 core. Juniperus is known to prefer dry habitats, while Thuja prefers wetter conditions (Johnston, 525 1990; Yu et al., 1996; Yu 1997). Nonetheless, we infer that Cuppressaceae were composed 526 mainly of *Thuja* at Charland Lake due to the presence of *Thuja* stomata observed on pollen 527 slides. This interpretation agrees with that of Liu (1990) who inferred that *Thuja* represented 528 most Cupressaceae pollen at Lake Six based on the occurrence of fossil Thuja seeds. Carcaillet et 529 al. (2001) also inferred the presence of *Thuja* rather than *Juniperus* because charcoal analysis 530 shows that forest fire frequency did not increase in the area during the HTM which would be 531 consistent with a wetter, Thuja-rich environment. 532 *Pinus strobus*, which increased in relative abundance alongside Cupressaceae likely

533 expanded northward from the mixed Great Lakes-St. Lawrence forest as average temperature 534 increased (Terasmae and Anderson, 1970; Richard, 1980; Bartlein et al., 1984, Liu, 1990; Hall et 535 al., 1994). Concomitant changes in P. strobus and Cupressaceae during the HTM also argues for 536 the presence of Thuja, as competition for the drier upland areas would not have allowed the 537 coexistence of Juniperus and P. strobus (Liu, 1990). Thuja would have had minimal habitat 538 competition with P. strobus, allowing the species to coexist (Fowells, 1965; Liu, 1990). It is 539 therefore likely that *Thuja* proliferated in the widespread wetlands of the Clay Belt lowlands, 540 while P. strobus occupied dry upland sites (Liu, 1990).

We infer that the HTM manifested in two phases in the Charland Lake region. The first phase, from ~7500–6000 cal yr BP, saw the increase of *Pinus banksiana/resinosa* and increased average temperature ~2 °C compared to the previous zone. The second phase occurred from ~6000–4500 cal yr BP and experienced increases in *P. strobus*, a large increase in *Thuja*-inferred Cupressaceae, and a further increase in temperature by ~0.5–1.0 °C to a maximum of ~3 °C. This

546	reconstruction makes the maximum HTM temperature \sim 1.5–2.0 °C warmer than current
547	conditions, a value which is in line with temperature estimates provided by climatic modelling
548	for this region (Renssen et al., 2009, 2012). We also postulate that the second phase of the HTM
549	was wetter than present in this region, as Thuja would have inhabited widespread wetlands
550	which developed during this time. This interpretation is consistent with Prentice et al. (1991)
551	whose precipitation reconstructions of the Holocene of eastern North America shows an increase
552	in annual precipitation in northeast Ontario after ~6000 cal yr BP.
553	
554	The Post-HTM zone (~4500–1700 cal yr BP)
555	The Post-HTM zone was a transitional time between the warmer HTM and cooler
556	modern climate regimes. The reconstructed temperature shows a decline in temperature from the
557	maximum Holocene value of ~3.0 °C at ~5000 cal yr BP to ~1.5 °C ~4500 cal yr BP. Average
558	temperature remained around 1.5 °C for the majority of this zone. Floristically, this zone saw
559	clear decreases in P. strobus and Cupressaceae and increases in Picea and Pinus
560	banksiana/resinosa. In addition, the observed increase in Betula may suggest increased fire
561	activity and drier conditions which in turn would not allow for the continued high abundance of
562	Thuja after ~3700 cal yr BP (Carcaillet et al., 2001) which is shown in the concentration diagram
563	(Supplemental Figure 3).
564	

565 Modern zone (~1700–0 cal yr BP)

566 During the Modern interval, *Picea* increased, while *P. strobus* declined further,
567 contributing to the inferred decrease in average temperature to ~1.0 °C. *Betula* and other
568 common and uncommon arboreal tree types remained largely unchanged throughout this period.

569 It is quite likely that this climate zone experienced an increase in precipitation as represented by 570 the increased abundance in *Picea* and decrease in *P. banksiana/resinosa* (Liu, 1990; Prentice et 571 al., 1991).

572

573 Lake and algal responses to Holocene climate change

574 Definition of climate zones based on historical changes in terrestrial vegetation allows us 575 to evaluate both how regional climate variation may have affected the linkage between land and 576 water, and the responses of past changes in the production and community composition of 577 aquatic primary producers. Here we evaluate changes in community composition of phototrophs 578 (using fossil pigments and diatom assemblages) better understand changes in climate on aquatic 579 environments.

580

581 *Pre-HTM zone (~8200–7800 cal yr BP)*

582 Northeastern Ontario was inundated by Glacial Lake Ojibway during the early Holocene. 583 This glacial lake deposited thick clay layers which form the Great Clay Belt in the Cochrane 584 District of northern Ontario before draining into the Tyrrell Sea ~8200 cal yr BP (Veillette, 1994; 585 Roy et al., 2011, 2015) resulting in the formation of Charland Lake. Algal production (both 586 pigments and diatoms) was too low to be detected or analyzed statistically in the Pre-HTM zone. 587 This may be due to the deposition of the first clay layer during this time. Clastic or mineral-rich 588 sedimentation is common in the early ontogeny of post-glacial boreal lakes (Liu, 1990; Teller et 589 al., 2018) and is known to interrupt fossil preservation and signals of algal abundance when 590 sedimentary organic matter is extremely low (Leavitt and Hodgson, 2001; Bunting et al., 2010).

592 HTM zone (~7800–4500 cal yr BP)

593 The climate rapidly warmed at the onset of this zone as shown in the temperature 594 reconstruction based on pollen (Figure 3), but pollen data suggests a two-phase warming event; 595 first warm and dry conditions followed by warm and wet conditions. Charland Lake continued to 596 experience major clastic input at the beginning of this zone, although microfossils from algae 597 and cyanobacteria begin to appear at this time. High abundance of S. construens and S. pinnata 598 during the early HTM suggests low-light or turbid conditions (Haworth, 1976; Bradshaw et al., 599 2000; Fritz et al., 2004) consistent with high clastic influx. These benthic diatoms are known to 600 be tolerant of low-light environments (Punning and Puusepp, 2007; Kingsbury et al., 2012) 601 which, along with the low concentrations of both diatoms and pigments, suggests an unproductive and light-limited environment. Influx of clastic material in Charland Lake may 602 603 reflect high rates of terrestrial erosion from the undeveloped landscape. The young forest 604 surrounding Charland Lake was likely open during the early HTM, favoring high rates of 605 weathering and transport of clay particles to the lake (Dearing, 1983; Dearing and Foster, 1986; 606 Almquist-Jacobson, 1992).

607Deposition of organic matter begins in the HTM, although algal fossils remain rare until608~7000 cal yr BP (Figure 5). The appearance of fossil pigments from many but not all groups was609likely a response to HTM warming, soil development, and the influx of organic matter which610would have created anoxic sediments which provided a better environment for pigment611preservation (Leavitt, 1993; Leavitt and Hodgson, 2001; Leavitt et al., 2003). Consistent with612this pattern, the ratio of chlorophyll *a* to phaeophytin *a* suggests a high degree of pigment613preservation (Leavitt and Hodgson, 2001) following the period of high clastic influx. Lower light

614 penetration during the final period of fine inorganic matter influx may have reduced photo-615 oxidation of pigments and favoured elevated fossil concentrations (Furlong and Carpenter, 1988; 616 Hurley and Armstrong, 1990, 1991; Leavitt, 1993). Finally, inferred low water-levels (based on 617 the high abundances of benthic diatoms) may have favoured development of benthic mats of 618 chlorophytes and cyanobacteria (as canthaxanthin). Typically, labile pigments (such as Chl a) are 619 preserved better if they are produced in benthic mats (Leavitt et al., 1994b; Leavitt et al., 2003). 620 At ~6000 cal yr BP, Thuja-inferred Cupressaceae increases rapidly, signifying the onset 621 of wet conditions (Figure 3). Several major phycological and sedimentological changes 622 coincided with this wetter climate. First, elevated organic matter sedimentation persisted until 623 \sim 6100 cal yr BP and was characterized by increased pigment concentrations, elevated relative 624 abundance of planktonic diatoms, and an increase in chrysophyte scales indicative of deeper lake 625 conditions (Zeeb and Smol, 2001; Figure 5, 7). Contemporaneous with increased temperatures 626 and lake-levels, landscape stabilization was occurring, as indicated by a hiatus in clastic 627 sedimentation which suggests forest closure and reduced erosion (Liu, 1990). The transition to 628 gyttja-based sediments also suggests the presence of soluble nutrients leaching from fresh, 629 organic soils (Liu, 1990).

Organic sedimentation was interrupted by the deposition of the second clay layer ~6000– 5500 cal yr BP, at which time concentrations of phototrophic fossils declines and diatom assemblages reverted to benthic taxa. Paradoxically, high Cupressaceae pollen abundance suggests a wet climate and high lake-levels. We propose that water levels increased sufficiently to let Charland Lake join with surrounding aquatic ecosystems, including nearby Fredrick House Lake, a site which is only ~2 m below the study basin. As Fredrick House Lake is turbid, conjoined waters may have introduced clastic material from the Fredrick House Lake catchment

637 into Charland Lake. Aquatic conditions in this second clay band were similar to those seen in the
 638 early HTM period, where diatom assemblages were composed mainly of benthic species and
 639 pigment and frustule concentrations were low.

640 After ~5500 cal yr BP, organic sedimentation resumed with no noticeable change in 641 pollen assemblages. In contrast, diatom assemblages rapidly shifted to planktonic taxa, similar to 642 that observed within the organic layer between clay bands, but with elevated abundance of the 643 eutrophic taxon Stephanodiscus minutulus. Stephanodiscus taxa are common in productive 644 waters (Cumming et al., 2015) and compete for silica better than other planktonic species 645 (Mechling and Kilham, 1982). Past research has associated higher water levels with lower Si:P 646 ratios and an associated dominance of *Stephanodiscus* (Kilham and Kilham, 1990). The 647 eutrophication may reflect increased internal loading of nutrients from anoxic sediments, as 648 indicated by the presence of okenone, a pigment from purple sulfur bacteria. These prokaryotic 649 taxa are obligate anaerobes and are present only when light penetrates to anoxic environments 650 (see Leavitt et al., 1989). Regardless of the mechanism, the increase in nutrients must have been 651 relatively minor, as neither frustule concentration nor that of most pigments suggests a strong 652 increase in primary production during this period.

653

654 *Post-HTM zone* (~4500–1700 cal yr BP)

Reconstructed average temperature shows that this zone experienced a cooler climate than the HTM but was still warmer than modern conditions. Algal groups responded to these climate changes; both diatom frustules and pigment concentrations increased moderately, possibly signaling an overall increase in primary production. The eutrophic *S. minutulus* continued to dominate the diatom assemblage until ~3900 cal yr BP when it was replaced by the

660 more oligotrophic Distostella stelligera (Figure 7). Presently, we are uncertain whether observed 661 changes in fossil patterns reflect direct effects of climate, or indirect variation in lake structure 662 and function, such as a change in thermal stratification. For example, increased abundance of D. 663 stelligera has been shown to correspond with a longer period of stratification (Dean et al., 1994; 664 Bradbury et al., 2002; Wiltse et al., 2016) and a shallow mixing depth (Saros et al., 2012). This 665 interpretation is supported by changes in the chrysophyte scale-to-diatom index, as chrysophytes 666 have a competitive advantage over diatoms when the water column is strongly stratified (Eimers 667 et al., 2009). An increase in the abundance of chrysophytes has also been related to less 668 eutrophic conditions, which agrees with an increase in D. stelligera (Reynolds, 1988; Brabury et 669 al., 2002).

670 Despite a trend towards oligotrophication as suggested by diatoms, interpretation of fossil 671 pigment data suggests an increase in algal production during this period. Several mechanisms 672 could underlie this pattern. First, increased thermal stratification can lead to the formation of 673 metalimnetic blooms which tend to be over-represented in lake sediments (Leavitt et al., 1989; 674 Leavitt and Hodgson, 2001). Alternately, changes in the seasonality of lake stratification could 675 favor production of phototrophic groups other than diatoms. For example, both lutein-676 zeaxanthin, a stable indicator of green algae and cyanobacteria, and alloxanthin, a stable 677 indicator of cryptophytes, had significantly higher concentrations (Figure 6) during this period compared to the HTM zone. Lower concentrations of other pigments from cyanobacteria (e.g., 678 679 echinenone, canthaxanthin, myxoxanthophyll) suggest limited contributions of those taxa to the 680 lutein-zeaxanthin signal (Brock et al., 2006). Both cryptophytes and chlorophytes (together 681 'flagellates') tend to replace negatively buoyant diatoms during periods of thermal stratification, 682 due to their motility (Hickman, 1974; Reynolds, 1984). As such, higher concentrations of these

algal groups suggest a longer period of thermal stratification during the Post-HTM period and,
perhaps, reduced diatom recruitment, which favoured growth of other phytoplankton groups,
such as cryptophytes and chlorophytes.

686

687 Modern zone (~1700–0 cal yr BP)

688 At Charland Lake, the Modern period of northeastern Ontario was characterized by a cold 689 and wet environment based on the decrease of P. strobus and the dominance of Picea. However, 690 algal proxies from Charland Lake suggest the onset of more eutrophic conditions as 691 concentrations of diatoms, and pigments characteristic of diatoms (fucoxanthin, diatoxanthin), 692 were highest during this period. Statistical analyses of individual pigments revealed that 693 canthaxanthin and okenone increased significantly (Figure 6), indicating an abundance of 694 colonial cyanobacteria and purple sulfur bacteria, respectively (Leavitt and Hodgson, 2001). 695 Given that the Chl *a*:pheophytin ratio declined slightly prior to the Modern period (Figure 5), but 696 okenone increased, we infer that these changes may reflect in part a minor change in the 697 preservation environment at Charland Lake. However, because this interval is also marked by the 698 first occurrence of chemically-robust myxoxanthophyll, a ubiquitous compound in colonial 699 cyanobacteria, we infer that the Modern zone has also been more productive than previous 700 periods.

The movement towards colder conditions during this period may have also promoted more lake mixing and less-stable thermal stratification, allowing some algal species, such as *S. minutulus*, to proliferate (Reynolds and Reynolds, 1985; Kilham et al., 1986; Makulla and Sommer, 1993; Bradbury et al., 2002). Growth requirements of *Stephanodiscus* species may have been largely satisfied by the regeneration of phosphorus from nutrient-dense water to the

photic zone during spring circulation, resulting in large blooms of *S. minutulus* and *S. parvus*(Kilham and Kilham, 1978; Bradbury et al., 2002). The inference of extended periods or
vigorous lake-mixing is also supported by the decrease in the chrysophyte scale-to-diatom index,
further suggesting less stratification (Reynolds, 1984; Rott, 1984).

710 Interpretation of the influence of climate on canthaxanthin (Nostocales cyanobacteria) 711 and okenone (purple sulfur bacteria) is not straightforward, as both of these algal groups would 712 be expected to decrease under cooler and wet conditions (Lami et al., 2009). Okenone is found in 713 anaerobic purple sulfur bacteria which prefer the oxic-anoxic interface in lakes (Massé et al., 714 2002), hence higher concentrations of okenone can be related to seasonal anoxia in lakes 715 (Maheux et al., 2016). On the other hand, low concentrations of purple sulfur bacteria have been 716 related to cool conditions, likely coupled with increased mixing of the water column (Schmidt et 717 al., 2002). Overall, concentrations of okenone are much lower than those seen in meromictic or 718 strongly stratified ecosystems (Leavitt et al., 1989) and suggests the presence of only seasonal 719 (summer) anoxia, perhaps restricted to deep littoral sediments. Furthermore, canthaxanthin, an 720 indicator of Nostocales cyanobacteria, is often associated with high nutrients in the water column 721 (Kleppel et al., 1988; Lami et al., 2009), but can also be a marker of N₂ fixation (Hayes et al., 722 2018). One possibility is that the Modern period experienced in an influx of dissolved organic 723 matter (DOM) which would have helped proliferate the heterotrophic growth of the 724 cyanobacteraial groups (Stevenson et al., 2016). Regardless it seems likely that factors other than 725 climate must be considered to understand how limnologic conditions became ideal for these 726 groups in the Modern period.

727

728 Conclusions

729 Pollen data from Charland Lake suggests that the HTM in northeast Ontario took place 730 from ~7800-4500 cal yr BP. The onset of warmth is ~1000 years later than recorded in 731 northwest Ontario (Moos et al., 2009; Moos and Cumming, 2011, 2012; Karmakar et al., 2015a, 732 2015b). Differences between our findings and Liu's (1990) may be due to the more rigorous 733 carbon dating model used in our study as our pollen analysis agrees with Liu's (1990) quite well. 734 Analysis of fossil pollen showed that the HTM in northeastern Ontario was warm and dry from 735 ~7500–6000 cal yr BP and then become wet from ~6000–4500 cal yr BP as indicated by the high 736 abundance of *Thuja*-inferred Cupressaceae pollen. Algal abundance was variable over the 737 Holocene, as inferred from both fossil pigment and diatom concentrations, apparently responding 738 mainly to variation in the linkage between land and water.

739 This work represents one of the first investigations in northeast Ontario to assess the 740 relationship between climate and in-lake production during the Holocene Epoch. Overall, the 741 interaction between climate and limnological conditions were complex within Charland Lake, 742 providing a striking comparison with lakes in northwest Ontario at millennial time scales. In 743 general, we demonstrate that limnological conditions were more optimal for algae over the last 744 \sim 4500 years than in the preceding millennia. This pattern is in stark contrast to those seen in 745 lakes from northwest Ontario, where paleolimnological studies have linked the warmer HTM to 746 enhanced lake-water production (Moos and Cumming, 2009; Karmakar et al., 2015b). In 747 Charland Lake, such changes were evident, but small in comparison, and did not indicate the 748 warmer temperatures of the HTM directly influenced lake-water production. These findings 749 suggest indirect climate effects, such as changes in thermal stratification and mixing, may be 750 more influential on algal production in some lakes in northeast Ontario. If this were broadly true, 751 it suggests that lakes in northeast Ontario may respond differently to future climate change

compared to lakes elsewhere in northern Ontario, and that the response and susceptibility oflakes to climate change can vary across an ecozone.

754

755 Acknowledgements

756 The authors thank the LacCore team at the University of Minnesota and Tom Brown at 757 the Lawrence Livermore National Laboratory for sample preparation and carbon dating. Thanks 758 also to Matthew Peros for his helpful advice on pollen sample preparation and to Kathleen Laird 759 for her assistance with diatom processing and taxonomy. We thank Deirdre Bateson at U. Regina 760 for analysis of fossil pigments. Special thanks to Graham Mushet, Cécilia Barouillet, and Gladys 761 Kong for help in the field. Additional thanks are given to two anonymous reviewers whose 762 comments greatly improved this manuscript. Funding for this project was provided by an 763 NSERC Discovery Grant to BFC, SFL and PRL, an NSERC PGS-D scholarship to CACG, and 764 funding from Canada Research Chair and Canada Foundation for Innovations programs.

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1083 1084	Supplemental figure 1. A) Activity of ²¹⁰ Pb, ²¹⁴ Pb, ²¹⁴ Bi, and ¹³⁷ Cs in the Charland Lake gravity
1085	core. B) Constant Rate of Supply (CRS) model inferred ages for the Charland Lake gravity core.
1086	
1087	Supplemental figure 2. Broken stick model based on a constrained cluster analysis (CONISS) of
1088	abundant pollen taxa (>1% in two or more intervals) from Charland Lake.
1089	

- 1090 Supplemental figure 3. Total concentration of pollen grains (x10⁴) from Charland Lake. Darker
- 1091 zones in Sediment Type represent organic sedimentation in the piston core. Layered zones
- 1092 represent clay deposits. Climate Zones are derived through CONISS from Figure 3.