## Geochemical reconstruction of late Holocene drainage and mixing in Kluane Lake, Yukon Territory

Janice Brahney (Corresponding Author), Department of Earth Science, Simon Fraser

University, Burnaby, B.C., (604) 291-3062, jbrahney@sfu.ca

John J. Clague, Department of Earth Science, Simon Fraser University, Burnaby, B.C.

Brian Menounos, Department of Geography, University of Northern British Columbia

Thomas. W.D. Edwards, Department of Earth Science, University of Waterloo

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#### Abstract

The level of Kluane Lake in southwest Yukon Territory, Canada, has fluctuated tens of metres during the late Holocene. Contributions of sediment from different watersheds in the basin over the past 5000 years were inferred from the elemental geochemistry of Kluane Lake sediment cores. Elements associated with organic material and oxyhydroxides were used to reconstruct redox fluctuations in the hypolimnion of the lake. The data reveal complex relationships between climate and river discharge during the late Holocene. A period of influx of Duke River sediment coincides with a relatively warm climate around 1300 yr BP. Discharge of Slims River into Kluane Lake occurred when Kaskawulsh Glacier advanced to the present drainage divide separating flow to the Pacific Ocean via Kaskawulsh and Alsek rivers from flow to Bering Sea via tributaries of Yukon River. During periods when neither Duke nor Slims river discharged into Kluane Lake, the level of the lake was low and stable thermal stratification developed, with anoxic and eventually euxinic conditions in the hypolimnion.

## Introduction

Kluane Lake is the largest lake in Yukon Territory, Canada, with an area of 409 km<sup>2</sup> (Figure 1; Natural Resources Canada 2003). Geological evidence indicates that the size and level of Kluane Lake have fluctuated markedly throughout the Holocene. Drowned trees and submerged beaches indicate lake levels up to 30 m below present (Bostock 1969, Rampton and Shearer 1978a, Clague et al. 2006), and raised shorelines and beach deposits occur up to 12 m above present lake level (Bostock 1969, Clague 1981, Clague et al. 2006).

The most recent rise in the level of Kluane Lake to its +12 m highstand occurred in the 17<sup>th</sup> century, during the Little Ice Age advance of Kaskawulsh Glacier. Dendrochronological evidence has constrained the time of this rise to a 50-year period beginning in AD 1650 and ending between AD 1680 and 1700 (Clague et al. 2006). Bostock (1969) hypothesized that, prior to the Little Ice Age, Kluane Lake drained southward through the Kaskawulsh River valley (Figure 1). The advance of Kaskawulsh Glacier blocked the southerly outlet and forced glacial meltwater and seasonal snowmelt from the glacier and its catchment directly to Kluane Lake via Slims River (watershed area = 1259 km<sup>2</sup>). Kluane Lake rose 12 m above its present level and overtopped the Duke River fan at the north end of the basin, establishing the current drainage route. The outflow incised the fan, lowering the lake to its present level.

The influence of Kaskawulsh Glacier on Kluane Lake prior to the 17<sup>th</sup> century is presently unknown. The glacier advanced several times before the Little Ice Age (Borns and Goldthwait 1966; Denton and Stuiver 1966; Denton and Karlén 1977) and may have contributed meltwater to the lake at those times.

Duke River, with a watershed area of 631 km<sup>2</sup>, presently bypasses Kluane Lake, joining Kluane River 4 km north of the lake outlet (Figure 1). Aerial photographs and satellite images, however, reveal abandoned Duke River channels extending to the northwest shore of Kluane Lake near Burwash Landing, but it is not known when the channels were last active.

Geochemistry of lake sediments is a complex function of processes occurring within the lake and the surrounding catchment. The concentrations of dissolved solutes and sediment originating from the watershed are modified during transport, deposition, and early diagenesis. The elemental chemistry of lake sediments can record details about weathering, runoff, lake productivity, pH, and redox conditions (Engstrom and Wright 1984; Boyle 2001). Climate change and associated watershed drainage processes can influence the types of sediments that enter the lake, as well as limnological characteristics such as lake temperature, depth, and redox conditions. Sediment geochemistry can also be used to fingerprint sediment sources in the watershed (e.g., Mosser 1991; Collins et al. 1997, 1998).

We use sediment geochemistry to provide insights into the relationship between historical climate change and drainage in the Kluane Lake basin. In addition, we infer changes in mixing depths, coincident with fluctuations in lake level.

#### Study area

Kluane Lake is located within Shakwak Trench in southwest Yukon Territory (Figure 1). The Kluane Ranges to the west are the easternmost range of the St. Elias Mountains. The Ruby Range to the east is part of the Yukon Plateau. The St. Elias Mountains support the largest ice fields and glaciers in North America, including Kaskawulsh and Donjek glaciers, which terminate, respectively, 20 km south and 40 km west of Kluane Lake.

The Denali fault extends in a northwesterly direction along the west side of Kluane Lake. It separates sedimentary and volcanic rocks of the Alexander terrane in the Kluane Ranges on the west side of the lake from high-grade metamorphic rocks of the Yukon-Tanana terrane in the Ruby Ranges to the east (Campbell and Dodds 1982). Thick glaciofluvial and glaciolacustrine sediments dating to the last glaciation (Kluane Glaciation) and one or more earlier glaciations underlie Kluane Lake and border it to the east.

During summer, when Slims River discharge is greatest due to melt of snow and ice, Kluane Lake typically rises 1-2 m above its winter level. Slims River is the dominant source of sediment to the southern portion of the lake. A sediment plume of glacial rock flour transported by Slims River covers much of the southern part of the lake during icefree periods. The rock flour is derived from both bedrock and sediments in the Slims River watershed. Turbid water from Slims River is denser than the lake water and sinks as it flows outward (Bryan 1972). Bottom waters are thus well aerated and the lake mixes throughout the summer. Silt in overflow and interflow plumes derived from Slims River rains out onto the floor of the southern half of the lake during summer and fall. Deposition of silt is restricted to water depths greater than 5 m, and most of it occurs at depths greater than 10 m. Currents are too vigorous at shallower depths for silt to accumulate there. Large amounts of silt and sand are carried from the slope of the Slims delta into deeper parts of the southern half of the lake by turbidity currents. Other sediment sources include Gladstone Creek at the northeast corner of the lake, Silver Creek at the south end of the lake, and several ephemeral streams that flow across large fans into the lake along its west side.

## Methods

#### Core collection and analysis

Thirteen percussion cores, 11 from Kluane Lake, one from Cultus Bay, and one from Grayling Lake, were collected in July 2004; four of the 13 cores were used in this study

(Figure 2). Six or more suspended sediment and floodplain samples were collected from Slims River, Silver Creek, Bock's Creek, and Duke River. In addition, a representative sample of glaciolacustrine silt was collected from the east side of the lake. All cores were split and analyzed at a high resolution for bulk physical properties. Three Kluane Lake cores (08, 10, and 36) and the Cultus Bay core (26) were selected for geochemical analysis. Plug samples, 1 cm in diameter, were taken from each core at intervals of 6-10 cm, depending on the stratigraphy.

A test was conducted to determine if bulk elemental geochemistry is affected by elements in the organic or oxyhydroxide fraction of the sediments. Samples from core 10 were treated first with tetra-sodium pyrophosphate (Na<sub>4</sub>P<sub>2</sub>O<sub>7</sub>) to remove metals associated with organic matter. Pyrophosphate does not attack sulfides and it does not dissolve amorphous iron oxyhydroxides (Ross and Wang 1993). Sodiumcitrate/dithionite ( $(Na_3C_6H_5O_7)$ ·( $Na_2S_2O_4$ )) was used as a reducing agent to remove Fe and Mn oxyhydroxides. Magnetite and silicates are not dissolved by this treatment (Ross and Wang 1993). The residual sediments were washed with distilled water and aspirated until neutral pH was achieved before the next digestion and final analysis. Major and trace elements in the extracts, residual sediment, and bulk untreated sediment were analyzed using an inductively coupled plasma-mass spectrometer (ICP-MS) and an inductively coupled plasma-atomic emission spectrometer (ICP-AES) at the Ontario Geological Survey Geosciences Laboratory. Trends of some elements in the bulk sediment samples and the treated residual sediment differ, thus the procedure outlined above was used to analyze the remaining lake and stream samples.

Plant macrofossils were radiocarbon dated by the AMS method at Beta Analytic and IsoTrace laboratories. All dates are reported as calibrated ages. Additional dating control is provided by the White River tephra, which is about 1150 years old (Clague et al. 1995). The tephra, <sup>14</sup>C ages, and the tree-ring ages for the recent rise of Kluane Lake, rising above present level at 1650 AD, were used to estimate sedimentation rates and dates of major events recorded in the cores.

#### Data analysis

Principal component analysis, cluster analysis, discriminant analysis, Euclidean distance metrics, and sediment unmixing models were used to cluster sediment intervals and ascribe them to a particular sediment source. Principal component analysis was performed using singular value decomposition on the covariance matrix. The analysis was done on the residual stream sediment data to determine if individual streams had unique principal factors, and on core data to determine whether elemental associations in principal components are related to particular stream sources. The non-hierarchical K-means cluster method was used on the residual data to group sediment samples. Cluster centres were iteratively defined and calculated using the Euclidean distance metric. To assign each sediment interval to a dominant sediment source, we used both multi-group discriminant analysis and unweighted Euclidean distances, E:

$$\mathbf{E} = \sqrt{\sum (S_i - C_i)^2} \tag{1}$$

where S<sub>i</sub> is the stream source concentration of element i, and C<sub>i</sub> is the core interval concentration of element i. Finally, constrained least squares analysis was performed to

determine the relative contribution of each stream to specific core intervals. Calculations are similar to those of Bryan et al. (1969). Composite element concentrations were used to minimize the error of proportional stream contributions to each core sediment interval. Constraints are such that each stream source proportion must be:

 $0 \leq S_i \leq \!\! 1$ 

$$\sum_{i=1-4}^{n=i} S_i = 1$$

## **Results**

#### Core descriptions

Core 36 was collected near the southeast end of Kluane Lake at a depth of 36 m (Figure 2). The core is 240 cm long and comprises three units (Figure 3). Unit 1 (240-97 cm) consists of light grey silt (Munsell colours 5Y6/1 and 5Y5/1) with black laminae up to 1 mm thick. Black laminae are most common between 120 and 97 cm depth. Light grey laminae from 120 to 97 cm are coarser grained than those lower in the core. A piece of wood at 219 cm depth yielded a radiocarbon age of  $3910 \pm 80^{14}$ C yr BP (4570-4090 cal yr BP; Table 2.1). Unit 1 interfingers with unit 2 (96-65 cm), which consists of massive brown silt (7.5YR4/1 - 7.5YR4/2) with several fine sand laminae. The White River tephra occurs at 88 cm depth. Unit 2 is abruptly overlain by unit 3, which consists mainly of light olive-grey clayey silt (5Y6/1-2). Darker laminations (5Y4/1) from 1 to 5 mm thick occur throughout the unit. Sedimentation rates in units 1 and 2 are 0.04 and 0.03 cm yr<sup>-1</sup>, respectively, and increase to 0.2 cm yr<sup>-1</sup> in unit 3.

Core 08 was collected at a depth of 25 m near the east shore of Kluane Lake between Christmas and Cultus bays (Figure 3). It is 63 cm long and comprises four units. The base of the core and the material found in the core catcher are coarse sand. Unit 2 (63-34 cm) consists of laminated fine to medium grey sand (10Y2.5/1) with a silty section between 61 and 54 cm depth. White River tephra occurs at 39 cm, and additional chronological control is provided by a radiocarbon age of  $1660 \pm 40$  <sup>14</sup>C yr BP (1690-1500 cal yr BP) at 50.5 cm. The contact between units 2 and 3 is gradational and interlayered. Unit 3 (33-17 cm) is laminated brown silt and very fine sand (2.5Y5/2 and 2.5Y3/3) with distinct bright orange layers (10YR5/6). Unit 3 is sharply overlain by unit 4 (16-0 cm), which is composed of laminated light olive-grey clayey silt (5Y5/2-3) with pale yellow beds (5Y5/2) 1-1.5 cm thick. Sedimentation rates increase from 0.02 cm yr<sup>-1</sup> in units 2 and 3 to 0.05 cm yr<sup>-1</sup> in unit 4.

Core 10 was collected in 33 m of water on the east side of the lake, north of Cultus Bay. The core is 108 cm long and consists of four units (Figure 3). Unit 1 (108-69 cm) and unit 2 (68-36 cm) are upward-fining, light grey, laminated silts (5Y4/1). The contact between the two units is gradational. Unit 3 (35-19 cm) is brown silt with several orange laminae (2.5YR4/6). Unit 3 and 4 are separated by a 2-cm-thick layer of White River tephra. Unit 4 (18-0 cm) comprises light olive-grey clayey silt (5Y4/2) with diffuse orange laminae.

Core 26 was collected in Cultus Bay in 14 m of water. Cultus Bay is separated from Kluane Lake by a spit that is breached at its south end. The core is 180 cm long and consists of five units (Figure 3). Unit 1, from the base of the core to 156 cm depth, is coarse sand with forest litter at 177 cm. The forest litter includes conifer needles, detrital wood fragments, and rootlets. This unit is sharply overlain by unit 2 (155-140 cm), which consists of fine to medium sand and scattered plant detritus. Unit 3 (139-53 cm) marks the transition to lacustrine sedimentation and is composed of organic grey silt (5Y2.5/1-5Y3/1) with fine sand laminae. Several black streaks and laminations, roots in growth position, and plant detritus occur near the base of this unit. A piece of wood at 139 cm and another at 177 cm returned identical radiocarbon ages of  $1180 \pm 40^{-14}$ C yr BP (1180-980 cal yr BP). Unit 4 (52-42 cm) consists of rhythmically laminated light grey silt (5Y2.5/1) with 22-25 couplets. Unit 5 (41-0 cm) is weakly laminated light grey silt (5Y2.5/1 – 5Y4/1) with scattered black spots and streaks. A sharply bounded bed with low organic content occurs at 37-39 cm depth.

#### Elemental abundances

#### Residual

In core 36, major changes in element abundances occur at 153, 97, and 65 cm (selected analytical data are shown in Figure 4). Phosphorus, Ca, and Na increase from 65 cm to the surface; high values of these elements were also obtained in samples at 159 and 203 cm. Calcium and Sr co-vary in units 1 and 2 ( $r^2 = 0.82$ ), likely reflecting the presence of calcite.

Concentrations of Al, Ba, Cs, Ga, K, Li, Rb, Sn, Th, Tl, U, and W increase from 97 cm to the top of the core. Silver, Hf, Nb, Sb, Ti, Y, Zr, and the lanthanide elements have relatively low concentrations in unit 3, and higher, non-varying concentrations through the remainder of the core. Cobalt, Cr, Fe, Mg, Mn, Ni, Sc, V, and Zn co-vary, with generally low values in unit 3 and in single samples at 159 cm and 203 cm. Molybdenum, S, and organic carbon co-vary in unit 1 and increase from 130 to 97 cm. Cores 10 and 08 show similar trends to core 36. Elements associated with clay minerals (Al, Ba, K, and Rb) increase upward in these cores, whereas Mn, Ti, and Y decrease upward. Calcium and Sr co-vary from the base of the cores to units 3 and 4 in cores 08 and 10, respectively. Sulfur increases from 80 to 60 cm in core 10, and phosphorus concentrations are higher in units 4 and 3 in core 10 and 08, respectively.

Concentrations of Ca, Na, Sr, and Y increase from 50 to 40 cm in core 26 and are also relatively high at 70 and 25 cm (Figure 5). Aluminum, Fe, Mg, Mn, and Sc peak at 70 and 37 cm, and P is high through much of the core. Concentrations of S are high near the base of the core and, along with Mo, have single-sample peaks at 130 cm.

#### Citrate/dithionite extracts

Concentrations of many oxyhydroxide-associated elements in core 36 are lowest from 120 to 97 cm (Figure 6). Above 97 cm, Fe and Mn increase, respectively, by 29% to 530  $\pm$  8 ppm and by 23% to 10  $\pm$  0.08 ppm. Aluminum, As, Ba, Co, Ni, Ti, U, V, Zn, and Zr also increase above 97 cm. Oxyhydroxide-associated concentrations of Au, Cu, Rb, Sn, and Sr increase from 18 cm to the top of core 10, and other oxyhydroxide-associated elements increase upward above 60 cm in this core. Oxyhyroxide-associated elements increase from 35 cm to the top of core 08. Calcium, P, Sc, Sr, Ti, U, and V peak at 70 cm and 36 cm in core 26. Barium, Co, Cr, Fe, Rb, Y, Zn, and Zr increase upward in this core.

#### Pyrophosphate extracts

Organic-bound metals and %C decrease from 97 cm to the top of core 36 (Figures 4 and 6). Cadmium, Co, Cr, Fe, Mn, Ni, S, Ti, U, V, and Zn co-vary with %C. Pyrophosphateextractable S was detected only from the base of the core to 97 cm. Calcium, Mg, and Sc decrease upward in core 36. Uranium and V increase from 120 to 97 cm; their peaks are, respectively, slightly below and above the peaks in Mo and S in the residual fraction. Sodium and Mn are high from the base of the core to 65 cm and As and Ba concentrations are highest from 97 to 65 cm. Iron, Ni, and Zn peak above 65 cm, and K, Rb, Sr, and U increase from 65 cm to the top of the core. Most metals in core 10, as in core 36, co-vary with %C. Cobalt, Cu, Mn, Ni, Pb, and Zn peak at 50 cm, whereas As, Ba, Ca, Cr, Mg. Sr, Ti, V, Y, and Zr peak at 60 cm. Copper, Au, Cd, Cr, Ni, S, Sn, Ti, U, V, Y, Zn, and Zr, generally decrease upward in core 08; K, Rb, and Sr increase upward in this core. Cadmium, Co, Mg, Mn, Ni, and U peak at 18 cm in core 08, and As, Ba, Ca, Cd, and Mn are generally high from 18 to 10 cm. Copper, Mn, and Zn peak at 50-40 cm in core 26; no trends are evident in other organic-bound metals in this core (Figure 5).

#### Data analysis

Principal component analysis successfully grouped samples from the same stream, indicating that the sediment transported by each stream is different in composition and that source reconstructions using sediment geochemistry are possible. The first three principal components, which distinguish the four streams, explain 87% of the variance (Figure 7).

Principal components analysis of geochemical data from core 36 produced three significant factors that explain 83% of the variation. Factor 1 is interpreted to reflect a Slims River sediment source, with positive loadings on elements that are abundant in the Slims watershed, specifically Ca, Na, and P, and negative loadings on elements with low concentrations in the watershed. Factor 2 has positive loadings on Ca and Sr, suggesting that a carbonate mineral occurs in the sediment. This factor is important above 45 cm and below 97 cm in core 36. Two event laminae at 65 and 74 cm also load strongly on factor 2. Factor 3 has positive loadings on elements with high concentrations in Duke River sediments.

The first three factors explain 93% and 95% of the variance in the geochemical data from cores 08 and 10, respectively. It is difficult to ascribe any of the three factors to a particular sediment source, although two factors for each core appear to be weakly associated with Slims River and Bock's Creek sediments.

Results of the cluster analysis, Euclidean distances, discriminant analysis, and sediment unmixing models are generally consistent for core 36. Discriminant analysis indicates two major periods of Duke River influence in core 36, one from the base of the core to 139 cm depth and another from 65 to 47 cm (Figure 8). A Bock's Creek influence is evident from 136 to 97 cm and a Silver Creek influence from 89 to 65 cm. Euclidean distances give similar results. Constrained least squares analysis also indicates two periods of Duke River influence, one from the base of the core to 133 cm and another from 89 to 53 cm (Figure 9). The influence of Bock's Creek is reduced when Duke River's influence is high, but is evident from 133 to 97 cm and 73 to 36. A Silver Creek signature is evident until Slims River begins to dominate the sediment supply at 47 cm. Constrained least squares analysis and Euclidean distances reveal Slims-type intervals at 203 and 159 cm, but discriminant analysis assigns only the sample at 203 cm to Slims River. Cluster analysis indicates that these intervals are unique but does not associate them with the same sediment source as unit 3. The inclusion of the glaciolacustrine silt sample in the data set does not significantly alter the results of the constrained least squares analysis. Constrained least squares analysis was also performed on the data after

removing Mo and S, which are incorporated in authigenic sulfides, but no differences were evident in the results. The results from core 10 are similar to those from core 36.

Discriminant analysis indicates that core 08 is dominated by Duke River sediment. Only the surface sample and the sample at 56-55 cm have a Silver Creek source. Likewise, constrained least squares analysis suggests that most samples from core 08 have a Duke River source. A Silver Creek influence is evident at the base of the core, but decreases upward. A Bock's Creek sediment signature is present throughout the core, increasing in the upper two samples.

### Discussion

#### Sediment sources

Residual element abundances in core 36 change at 153, 97, and 65 cm depth (Figure 4). Based on average sedimentation rates, these changes date to about 2750, 1300, and 300 cal yr BP. The most recent change is marked by a gradual increase in Ca, Na, and P from 65 cm to the top of the core. These elements are characteristic of the Slims River source, which is consistent with the inception of meltwater inputs from Kaskawulsh Glacier about 300 cal yr BP. Elevated Na and P suggest an influx of unweathered rock flour. Phosphorus, as a limiting nutrient, is transformed within a few thousand years into bioavailable forms through soil development (Filippelli et al. 2006). Similarly, Na is easily leached during weathering, but is not readily sedimented through authigenic reactions, adsorption, or biological uptake (Engstrom and Wright 1984). The upward increase in Ca, Na, and P reflects the rapid progradation of the Slims River delta from the present Kaskawulsh-Slims drainage divide to its present location. The advance of the delta in historic time was rapid, averaging about 42 m yr<sup>-1</sup> from 1899 to 1970 (Rampton and Shearer 1978b).

Constrained least squares analysis and Euclidean distances assign the samples at 203 and 159 cm to a Slims River source, largely based on their elevated Ca, Na, and P concentrations (Figures 8 and 9). Discriminant analysis also identifies the sample at 203 cm as Slims-type sediment. These results suggest that Kaskawulsh Glacier meltwater flowed into Kluane Lake at least twice before 300 cal yr BP. Based on average sedimentation rates, the earlier meltwater inputs date to about 4000 and 2800 cal yr BP. Glaciers in the Purcell, Coast, Rocky, and Selkirk mountains advanced around 4000 cal yr BP (Gardner and Jones 1985; Osborn and Karlstrom 1989; Osborn et al., 2007; Wood and Smith 2004), and Kaskawulsh Glacier and other glaciers in the St. Elias Mountains advanced about 2800 cal yr BP (Borns and Goldthwait 1966; Denton and Stuiver 1966; Denton and Karlén 1977).

The gradient of Kaskawulsh River directly downstream of Kaskawulsh Glacier is steeper than that of Slims River. Slims River is thus vulnerable to being pirated by Kaskawulsh River. Only the presence of glacier ice and outwash in the divide area prevents this piracy (Figure 1). Any pre-Little Ice Age advance of Kaskawulsh Glacier that brought the toe of the glacier close to the present divide would probably route meltwater away from Kaskawulsh River and into Kluane Lake. The weakness of the Slims sediment signal at 203 and 159 cm is likely due to the distance (25 km) of core site 36 from the point of meltwater inflow to Kluane Lake. Peaks in these elements occur in sediment intervals with lower organic carbon contents than the rest of the core, suggesting rapid sedimentation, which is consistent with a new meltwater source. Constrained least squares analysis indicates two major periods of Duke River influence on Kluane Lake sediments. When Kluane Lake was 10 m or more lower than today, Duke River may have flowed into the lake south of Brooks Arm, strengthening the Duke River signal in cores 36 and 08. A possible stream channel extends southeast along the axis of Brooks Arm and may have carried Duke River during one or more periods of low lake level (Robert Gilbert, personal communication, 2004). In addition, imbrication in the Duke River fan sediments indicates that flow was to the southeast some time between 2000 and 600 cal yr BP (Clague et al. 2006).

At times, Duke River either flowed north away from Kluane Lake, as it does today, and did not affect the lake, or it discharged into an isolated basin near Talbot Arm. Dating and geochemistry of core 36 indicate that the river flowed into Kluane Lake from about 4850 to 2400 cal yr BP, when its influence began to decline. Little or no Duke River sediment entered the lake between about 2100 and 1300 cal yr BP. A Duke River influence again becomes evident about 1300 cal yr BP and continues to about 300 cal yr BP (Figure 10).

The beginning of the most recent period of Duke River discharge into Kluane Lake coincides with a major change in the climate of southern Yukon. Anderson et al. (2005) documented shifts in the North Pacific Index (NPI), a measure of sea-level pressure over the North Pacific, at about 2800 and 1300 cal yr BP. North Pacific pressure anomalies control climate in southwest Yukon. Analysis of modern historical climate data from Burwash Landing reveals a negative correlation of the NPI ( $r^2 = 0.65$ ) with annual temperature and a positive correlation ( $r^2 = 0.59$ ) with winter precipitation. Warming around 1300 yr BP, inferred from independent paleo-climate data (Anderson et al. 2005), may have thawed permafrost in soils in the watershed. Mildly reducing conditions in the catchment may have resulted from melt of stagnant ice. In a mildly reducing environment, manganese has a greater solubility than iron; an increase in Mn relative to Fe is observed from 97-65 cm (1300 to 300 yr BP) in all geochemical fractions.

Constrained least squares analysis indicates an upward decrease in sediment of Silver Creek provenance above 97 cm (1300 yr BP). This trend is consistent with Rampton and Shearer's (1978b) interpretation of the stratigraphy of Kluane Lake sediments off the Slims River delta. Their subbottom acoustic survey revealed two sharply bounded sediment units: an upper unit derived from Slims River and a lower unit deposited by Silver Creek and other local streams.

#### Anoxia in Kluane Lake

Constrained least squares analysis of geochemical data from core 36 suggests that Duke River either did not discharge into Kluane Lake from about 2100 to 1300 cal yr BP, or that its flow was reduced. Because Slims River did not exist at this time, the level of Kluane Lake was considerably lower than today and the basin may have been closed. Sediments deposited at this time contain black laminae, consistent with deposition under reducing conditions. Reducing conditions could develop from permanent or nearpermanent stratification (meromixis) in the lake. Meromictic and anoxic conditions are not commonly associated with low stands of oligotrophic lakes, but semi-permanent stratification could develop in Kluane Lake under some conditions. Groundwater entering the lake from the south and west is rich in dissolved solutes (150-3000 mg L<sup>-1</sup>; Harris 1990). Density stratification could develop in the low-level lake in the absence of mixing and influx of fresh Slims and Duke River waters. A concentration of total dissolved solids of roughly 340 mg L<sup>-1</sup> is enough to cause a greater density difference in water masses than that created by temperature differences. Pienitz et al. (2000) reported anoxic conditions in a shallow Yukon lake due to high Mg<sup>2+</sup> and SO<sub>4</sub><sup>2-</sup> concentrations, similar to concentrations of these ions in groundwater flowing into Kluane Lake (10 mg/L Mg<sup>2+</sup> and 50 mg/L SO<sub>4</sub><sup>2-</sup>).

Under low oxygen conditions, redox-sensitive elements can complex with organic acids or precipitate as insoluble oxyhydroxides or insoluble metal sulfides. Elements involved in reactions catalyzed by free H<sub>2</sub>S or deposited as organic complexes include Cr, U, and V (Calvert and Pedersen 1993, Algeo and Maynard 2004). Elements that can form insoluble sulfides in reducing conditions include Cd, Co, Mo, Ni, Pb, and Zn (Calvert and Pedersen 1993, Algeo and Maynard 2004). The presence or absence of these elements in sediments can provide information on paleo-redox conditions in Kluane Lake.

#### Molybdenum

Molybdenum and S concentrations co-vary ( $r^2 = 0.91$ ) from 153 to 97 cm in core 36, suggesting uptake of Mo in pyrite (Figure 4). The associated increase in organic carbon is not likely due to increased productivity, but rather to preservation of organic matter in a low-oxygen environment. Molybdenum can be concentrated in anoxic bottom waters through redox cycling in the water column. MoO4<sup>2-</sup> is easily scavenged by manganese oxyhydroxides (Berrang and Grill 1974; Adelson et al. 2001) in oxygen environments

releases  $MoO4^{2-}$  into solution. Sedimentation of Mo seems to require the formation of an intermediary species, thio-oxymolybdate ( $MoO_xS_{4-x}^2$ ) (Helz et al. 1996). Molybdenum can then be sedimented by scavenging on iron sulfides or by forming bonds with sulfurized organic matter (Helz et al. 1996; Adelson et al. 2001; Tribovillard et al. 2004). Free H<sub>2</sub>S/HS<sup>-</sup> is necessary for thiomolybdate to form, thus euxinic conditions are required in the water column, rather than simply reducing conditions in the sediments. Although conversion to thiomolybdates is catalyzed by mineral surfaces (Helz et al. 1996), Crusius et al. (1996) noted that Mo does not seem to accumulate in modern marine environments that are suboxic or anoxic, but rather only in marine environments that are euxinic. Thus, molybdenum fixation probably occurs more rapidly at the sediment-water interface than at depth in the sediment. It is unlikely that Kluane Lake is productive enough to produce strong reducing conditions within sediments after burial.

#### Copper

Copper can be precipitated in anoxic or euxinic environments as an independent sulfide phase, in solid solution with iron sulfides, or as an organic complex (Morse and Luther 1999). Copper co-varies with sulfur from 130 to 97 cm (2200 to 1300 yr BP) in core 36, perhaps due to authigenic precipitation. The increase in Cu from 97 to 65 cm (1300 to 300 yr BP) in the same core probably records a change in sediment source.

#### Vanadium and uranium

Pyrophosphate-extractable V and U increase from 130 to 97 cm (2200 to 1300 yr BP) in core 36 (Figure 6). Pyrophosphate-extractable U also increases from 65 cm (300 yr BP) to the top of the core. The latter increase may be the result of a change in sediment source because U also increases in the residual fraction. Citrate/dithionite-extractable V and U

decrease from 130 to 97 cm. Vanadium and U are known to concentrate in organic sediments under mildly reducing and euxinic conditions (Emerson and Huested 1991; Klinkhammer and Palmer 1991; Algeo and Maynard 2004). Vanadium, like Mo, can be concentrated in anoxic bottom waters (Wehrli and Stumm 1989).

Vanadium occurs as the vanadyl ion  $(VO^{2+})$  under mildly reducing conditions. It can be precipitated as insoluble oxides or hydroxides under strongly reducing conditions (Wanty and Goldhaber 1992). The reduced form of U is the uranyl ion  $UO^{2+}$ . No enrichment of either element as authigenic phases is evident in Kluane Lake sediments. Residual V is highly correlated with Zn throughout core 36 ( $r^2 = 0.80$ ). Vanadium correlates with Ti in the Slims interval (65-0 cm;  $r^2 = 0.97$ ) and from 159 cm to the base of the core ( $r^2 = 0.88$ ). It has a lower correlation with Ti from 153 to 97 cm ( $r^2 = 0.62$ ), but is highly correlated with Fe over this section of the core ( $r^2 = 0.92$ ). Residual U correlates with K and Al over the length of the core ( $r^2 = 0.86$  and 0.84, respectively), and U correlates with Fe in unit 1 ( $r^2 = 0.83$ ). These strong correlations suggest that, through much of the core, residual V and U are associated with non-authigenic fractions and thus are likely controlled by sediment provenance. The absence of enrichment in the authigenic phase may be due to competitive complexation of the dissolved species with organic matter. Vanadyl and uranyl ions commonly form organic ligands (Templeton and Chasteen 1980; Lewan and Maynard 1982; Emerson and Huested 1991; Klinkhammer and Palmer 1991; Algeo and Maynard 2004), and complexation of U and V with organic material under suboxic and anoxic conditions may leave the dissolved species unavailable for precipitation in sediments under reducing conditions. The requirement

that conditions be only mildly reducing may explain peaks in V and U prior to peaks in Mo and S.

#### Chromium, cobalt, nickel, and zinc

Chromium, Co, Ni, and Zn co-vary in all Kluane Lake cores. In core 36, Co, Cr, and Ni in the residual phase correlate strongly with Mg in unit 3 ( $r^2 = 0.92$ , 0.86, and 0.9, respectively), and in units 1 and 2 ( $r^2 = 0.91$ , 0.87, and 0.91, respectively). From the base of the core to 97 cm, Zn is strongly correlated with Al ( $r^2 = 0.62$ ), Fe ( $r^2 = 0.91$ ), and V ( $r^2 = 0.96$ ). These strong correlations suggest that the elements in the residual phase reflect sediment provenance.

Cobalt, Ni, and Zn can be precipitated as independent sulfide phases in anoxic environments, but the process is kinetically slow for Co and Ni (Morse and Luther 1999). Cobalt and Ni can be incorporated into pyrite, but structural and thermodynamic properties may restrict Zn and prevent Cr from co-precipitating altogether (Huerta-Diaz and Morse 1992; Morse and Luther 1999). Huerta-Diaz and Morse (1992) noted that Mo is more rapidly incorporated into pyrite than Co and Ni. Concentrations of Cu and Mo in Kluane Lake waters are greater than concentrations of Co, and Ni (J. Bunbury and K. Gajewski, unpublished data), which may account for the elevated values of authigenic Mo and Cu in Kluane Lake sediments.

Cobalt, Cr, Ni, and Zn correlate with %C in the pyrophosphate-extractable fraction in core 36 ( $r^2 = 0.68$ , 0.52, 0.76, and 0.55, respectively) and in the citrate/dithionite-extractable fractions below 97 cm. The association of Ni with the organic fraction suggests deposition under reducing conditions in the hypolimnion. In the reduced state, dissolved Ni is preferentially incorporated into organic tetrapyrrole complexes (Lewan and Maynard 1982). Tetrapyrrole complexes degrade faster than other types of organic matter, thus preservation requires deposition in a low-oxygen environment. An association of Cr and Ni with the organic fraction is consistent with the observation of Algeo and Maynard (2004) that these elements are associated with organic matter in non-sulfide anoxic and euxinic waters. The presence of Co and Zn in the organic fraction suggests that both elements can complex with organic matter under reducing conditions.

Zinc can complex with humic and fulvic acids in anoxic environments (Achterberg et al. 1997). The brief increase in Zn pyrophosphate at 65 cm in core 36 may record in-wash of terrestrial organic matter as Kluane Lake rose during the Little Ice Age. Calcium, Fe, and Ni also increase at this depth, possibly for the same reason.

#### Iron and manganese

Iron and Mn co-vary in core 36. The residual phases probably represent both detrital and authigenic minerals; both elements can form minerals through diagenetic precipitation. Iron and Mn oxyhydroxides are soluble in their reduced states and are insoluble under oxic conditions (Engstrom and Wright 1984). Thus concentrations of both elements should be low in the sediments during periods of anoxia.

The presence of Fe and Mn oxyhydroxides from 120 to 97 cm in core 36 does not necessarily argue against euxinic conditions (Figure 6). The normal sequence of reduction reactions is  $O_2$ ,  $NO_3^-$ ,  $MnO_x$ ,  $Fe(OH)_3$ , and  $SO_4^{2-}$ . The next electron acceptor must be almost used up before the reaction moves on to the next stage (Schlesinger 1997). This sequence, however, may not always occur in natural environments due to spatial heterogeneity and variable concentrations of available electron receptors. Kelly et

al. (1982) observed some sequential reduction in seasonally stratified lakes, where  $O_2$  and  $NO_3^-$  reduction ceased before the lakes overturned but all other reactions proceeded simultaneously. Sulfate concentrations in Kluane Lake and in the lakes that surround it are up to four orders of magnitude greater than dissolved Fe and Mn concentrations, and nitrate is only present in trace concentrations.

Complete dissolution of oxyhydroxides on the floor of Kluane Lake may require lengthy exposure to reducing conditions. Iron and Mn oxyhydroxides could still be deposited at such times because Kluane Lake is relatively shallow. Dissolution would surely occur, but oxyhydroxides would have a comparatively short transit through anoxic or euxinic bottom waters. The presence of Fe and Mn oxyhydroxides may also be an artifact of sampling. Sediments were sampled with a 1-cm diameter plug, and most samples contained both dark and light laminae.

Iron and Mn can be strongly complexed by organic matter (Engstrom and Wright 1984). Iron and Mn pyrophosphates increase with organic carbon from 153 to 97 cm in core 36. Reduction of oxyhydroxides may have liberated Fe and Mn, which were subsequently complexed with organic matter.

Core 10 is only slightly shallower than core 36 (33 m vs. 36 m), thus its sediments probably would have experienced bottom water anoxia too. Core 10 sulfur concentrations are elevated from 80 to 50 cm (3600 to 200 yr BP), and redox-sensitive elements (Co, Cu, Ni, Pb, and Zn) peak at 50 cm. Citrate extractions of As, Ba, Co, Fe, Mn, Ni, U, V, Y, Zn, and Zr are relatively low in this section of the core.

Core 08 is shallower than cores 36 and 10 and does not show the same associations with redox-sensitive elements. Pyrophosphate extractable metals decrease upward from near the base of the core. Secondary peaks of Co, Ni, and U occur at 17 cm. The trends may reflect higher concentrations of organic matter near the base of the core, associated with shallower water. The peak at 17 cm (300 yr BP) marks the base of the upper unit in core 08 and possibly records in-wash of terrestrial organic matter during the Kluane Lake high stand.

A return to mixing in the basin

A return to oxygenated conditions in Kluane Lake about 1300 cal yr BP is suggested by a rapid increase in Fe and Mn oxyhydroxides and associated trace elements at 97 cm (1300 yr BP) in core 36 (Figure 6). Citrate/dithionite-extractable Fe and Mn increase, respectively, 29% (to  $530 \pm 8$  ppm) and 23 % ( $10 \pm 0.08$  ppm) at this level. Arsenic, Ba, Co, Ni, Ti, U, V, and Zr increase from 29% to 260%, probably because they were scavenged by oxyhydroxides. Similarly, Fe and Mn oxyhydroxide concentrations in the citrate/dithionite extracts of cores 10 and 08 increase above, respectively, 35 cm and 25 cm (1300 and 600 yr BP).

Constrained least squares analysis suggests that Duke River began to flow into Kluane Lake about 1300 yr BP. Increased input of fresh cold water may have initiated mixing in the lake and re-oxygenated its bottom waters.

Aluminum, Mg, P, and Rb concentrations in citrate extracts increase at 65 cm in core 36. They also increase in the residual fraction, implying a change in source material rather than a change in oxidation state. Phosphorus sedimentation and retention in sediments are influenced by Fe and Mn oxyhydroxides (Engstrom and Wright 1984). Phosphorus does not correlate with either element throughout core 36, suggesting that its sedimentation may not be controlled by Fe or Mn for much of Kluane Lake's history. Phosphorus concentrations in Kluane Lake sediments appear to be related to a change in supply rather than redox state. Citrate-extractable P increases above 108 cm and again above 65 cm (1600 and 300 yr BP). Co-variation of citrate-extractable P and citrateextractable Sr and Ca suggests that carbonates may control their concentrations.

## Summary

Changes in the flow of Duke and Slims rivers affected the level of Kluane Lake and its redox state during the late Holocene. Duke River at times flowed into the lake and at other times bypassed it to the north. Sediment geochemical data indicate that Duke River flowed into Kluane Lake before 2100 cal yr BP and between about 1300 and 200 cal yr BP. Slims River has flowed into Kluane Lake for the past several centuries and, over this period, has deposited a thick wedge of sediments in the southern part of the basin (Clague et al. 2006). Older Slims River sediment, dating to about 4000 and 2800 cal yr BP, is present in one core in the southern part of the lake. Fluctuations in discharge and location of both rivers appear to be associated with climate fluctuations. Climate warmed around 1300 cal yr BP, and the glacier advanced at about 4000, 2800, and 300 cal yr BP.

Kluane Lake was low and stratified when Duke River and all meltwater from Kaskawulsh Glacier bypassed the lake. Meromixis led to anoxic and eventually euxinic conditions in the hypolimnion, possibly causing precipitation of Mo and Cu sulfides. The lack of enrichment of many other redox-sensitive elements can be explained by their low availability in the water column or by competitive complexation with humic and fulvic acids.

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<sup>14</sup> C age <sup>1</sup>	Laboratory no	Core no. and	Material	Calendar age <sup>2</sup>
$(^{14}C \text{ yr BP})$		sample depth		(cal yr BP)
$1660\pm40$	Beta-200708	08, 50.5 cm	wood	1500-1630, 1660-1690
$1180\pm40$	Beta-200709	26, 177 cm	wood	980-1180
$1310\pm40$	Beta-200710	31, 103 cm	wood	1170-1300
$1180\pm40$	Beta-213014	26, 139 cm	wood	980-1180
$3910\pm80$	TO-12468	36, 219 cm	spruce needle and twig	4090-4570

Table 1 Radiocarbon ages from Kluane Lake cores

<sup>1</sup> Radiocarbon laboratory: Beta-Beta Analytic Inc.; TO-IsoTrace Radiocarbon Laboratory (University of Toronto).

 $^{2}$  Determined from the calibration data set IntCal98 (Stiuver et al. 1998); calibrated age ranges are reported as  $\pm 2\sigma$ .

## **Figure captions**

Fig. 1. Kluane Lake, southwest Yukon Territory. Modified from Clague et al. (2006) with permission from Elsevier.

Fig. 2. Kluane Lake bathymetry and core locations. Cores described in this paper are designated by black dots. Modified from Clague et al. (2006) with permission from Elsevier.

Fig. 3. Lithostratigraphy of Kluane Lake and Cultus Bay cores.

Fig. 4. Representative concentrations of elements in the residual sediment fraction of core 36.

Fig. 5. Representative concentrations of elements in the residual, cithrate/dithionite, and pyrophosphate fractions of core 26.

Fig. 6. Representative concentrations of elements in the citrate/dithionite and pyrophosphate extracts from core 36.

Fig. 7. Principal component bi-plots showing separation of sediment sources.

Fig. 8. Sediment sources for samples from cores 08, 10, and 36 based on discriminant analysis and Euclidian distances.

Fig. 9. Constrained least squares results for core 36. Note the two major periods of Duke River influence. The Slims River source dominates the sediment from 42 cm to the surface.

Fig. 10 Reconstructions of Kluane Lake at three times, showing inferred flow directions of Duke River. The photographs show representative sediment deposited at each time.









Core 26







## **Principal Components Analysis**

Suspended stream and floodplain sediment



## Core 36

vr BP	Durith	District	Euclidean
J. 21	Deptn	Discriminant	Distance
	3-2	Slims	Slims
	9-10	Slims	Slims
	17-18	Slims	Slims
	24-25	Slims	Slims
	30-31	Slims	Slims
	36-37	Slims	Duke
	42-43	Slims	Duke
	47-48	Duke	Duke
	53-54	Duke	Duke
	56-57	Duke	Silver
200	64	Duke	Bocks
3007	67-68	Silver	Silver
	73-74	Silver	Silver
	74	Duke	Bocks
	80-81	Silver	Silver
	89-90	Silver	Bocks
1300	96-97	Duke	Bocks
	103-104	Bocks	Bocks
	105.5-106.5	Bocks	Bocks
	110-111	Duke	Silver
	113-114	Silver	Silver
	118-119	Bocks	Bocks
	121-122	Bocks	Bocks
	127-128	Bocks	Bocks
	133-134	Bocks	Duke
	136-137	Bocks	Duke
	139-140	Duke	Duke
	146-147	Duke	Duke
	153-154	Duke	Duke
2750-	159-160	Duke	Slims
	165-166	Duke	Duke
	173-174	Duke	Duke
	178-179	Duke	Duke
	184-185	Duke	Duke
	190-191	Duke	Duke
	196-197	Silver	Duke
	201-202	Duke	Duke
	203-204	Slims	Slims
	210-211	Duke	Duke
	216-217	Duke	Duke
4850-	232-233	Duke	Slims

Core 26

Depth	Discriminant	yr BP
0-1	Silver	
8-9	Slims	
14-15	Duke	
21.5-22.5	Duke	
26-27	Duke	
37-38	Duke	200
40-41	Slims	200
50-51	Bocks	200
62-63	Duke	500
70.5-71.5	Duke	
78-79	Duke	
102-103	Bocks	
116-117	Bocks	
131-132	Silver	
144-145	Slims	-1200

# Core 10

Depth	Discriminant	
2-3	Slims	
9-10	Slims	
16-17	Silver	-300
21-22	Bocks	
50-51	Duke	
60-61	Duke	
80-81	Duke	
103-104	Slims	<b>−</b> 5000



Depth	Discriminant	
1-2	Silver	
10-11	Duke	L300
18-19	Duke	500
25-26	Duke	
36-37	Duke	
55-56	Slims	-2400



