



Volcanic ash layers in Lake El'gygytyn: eight new regionally significant chronostratigraphic markers for western Beringia

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Abstract. Ash layers from explosive volcanic eruptions (i.e., tephra) represent isochronous surfaces independent from the environment in which they are deposited and the distance from their source. In comparison to eastern Beringia (non-glaciated Yukon and Alaska), few Plio-Pleistocene distal tephra are known from western Beringia (non-glaciated arctic and subarctic eastern Russia), hindering the dating and correlation of sediments beyond the limit of radiocarbon and luminescence methods. The identification of eight visible tephra layers (T0–T7) in sediment cores extracted from Lake El'gygytyn, in the Far East Russian Arctic, indicates the feasibility of developing a tephrostratigraphic framework for this region. These tephra range in age from ca 45 ky to 2.2 My old, and each is described and characterized by its major-, minor-, trace-element and Pb isotope composition. These data show that subduction-zone-related volcanism from the Kurile–Kamchatka–Aleutian Arc and Alaska Peninsula is the most likely source, with Pb isotope data indicating a Kamchatkan volcanic source for tephra layers T0–T5 and T7, while a source in the Aleutian Arc is possible for tephra T6. The location of Lake El'gygytyn relative to potential source volcanoes (> 1000 km) suggests these tephra are regionally distributed over a large area. These deposits provide a unique opportunity to correlate the high-resolution paleoenvironmental records of Lake El'gygytyn to other terrestrial paleoenvironmental archives from western

Beringia and marine records from the western North Pacific and Bering Sea, and to move towards the development of a robust integrated framework between the continuous paleoclimatic records of Lake El'gygytyn and other terrestrial and marine records in NE Eurasia.

1 Introduction

1.1 Lake drilling program and resultant cores

Lake El'gygytyn is located in the Far East Russian Arctic (67°30' N, 172°5' E), 100 km north of the Arctic Circle (Fig. 1). The 12 km wide and 175 m deep lake formed within a meteorite impact crater created 3.58 ± 0.04 million years ago (Gurov et al., 1978, 2007; Layer, 2000), and was cored under a jointly funded project led by the International Continental Scientific Drilling Program (ICDP) in winter 2008/2009. Two pilot cores (Lz1024, PG1351) were collected from Lake El'gygytyn during initial pre-site surveys (Juschus et al., 2009), followed by the main drilling program that collected three cores from the center of the lake (ICDP site 5011: holes 5011-1A, 5011-1B, 5011-1C; 67°29.98' N, 172°06.23' E), and one permafrost core at the lake margin (ICDP site 5011-3; 76°29.1' N, 171°56.7' E) (Melles et al., 2011, 2012). Approximately 318 m of lacustrine sediment

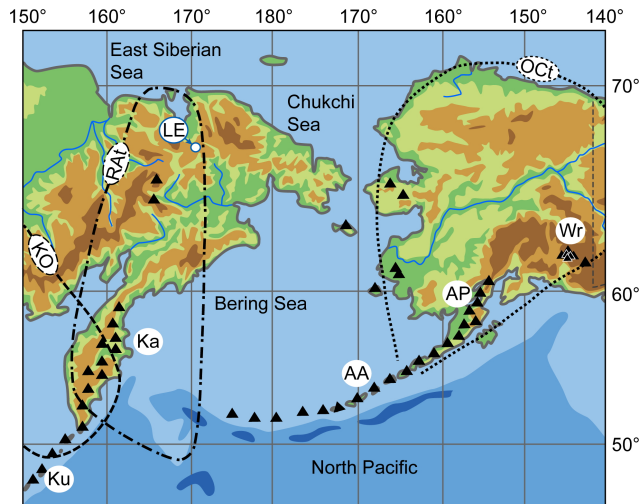


Figure 1. Location of Lake El'gygytyn (LE) in relation to volcanic areas in the Kuriles (Ku), Kamchatka (Ka), Aleutian Arc (AA) and Alaska Peninsula (AP). Reconstructed distribution patterns of Old Crow tephra (OCt; ~124 ka; Preece et al., 2011b), Rauchua tephra (RA; ~177 ka; Ponomareva et al., 2013b) and KO tephra (KO; ~7 ka; Derkachev et al., 2004) illustrate the potential for widespread tephra distribution in this region. The Wrangell volcanic field (Wr) and other smaller cinder cones and maars, indicated by additional triangular markers, are too far afield and/or do not produce the type of tephra deposits that are found in Lake El'gygytyn. During glacial times, the Bering shelf between Russia and Alaska was exposed, and only mountainous areas in this region experienced limited glaciation, creating the glacial refugium known as Beringia. As a result, Beringia contains terrestrial paleoenvironmental records that rival marine deposits in their richness and length.

were cored without hiatuses, spanning the Mid Pliocene to present (Melles et al., 2012). Tephra layers were identified and collected from all five lake cores, and were initially correlated by stratigraphy and assigned numerically based names, T0–T7.

1.2 Background

Tephra resulting from explosive volcanic eruptions can be distributed over vast areas. Because an eruption occurs over days to weeks, each tephra layer represents an isochronous horizon in the different types of depositional environments, in which it is preserved. Each tephra potentially has a unique geochemical fingerprint, allowing the correlation of the archives they occur in, and some can even be independently dated by physical dating methods (e.g., Shane et al., 1996; Westgate et al., 2001; Giaccio et al., 2012). The value of tephra layers has become increasingly recognized in climate and environmental studies over the last few decades. Tephra have been recovered from soil records, loess deposits, lake sediments, peat bogs, archaeological sites, marine and ice cores. As cryptotephra layers – a volcanic ash layer not visible to the naked eye – they have been traced

as far as 7000 km from their source (e.g., Dugmore, 1989; Grönvold et al., 1995; Pilcher et al., 1996; van den Bogaard and Schmincke, 2002; Wastegård, 2002; Davies et al., 2003, 2012; Dörfler et al., 2012; Jensen et al., 2012; Pyne-O'Donnell et al., 2012; Lane et al., 2013).

The tephra beds present in Lake El'gygytyn are of particular interest because of its location within western Beringia (Brigham-Grette et al., 2007). Western Beringia is the Eurasian half of Beringia, a large region extending from eastern Siberia to far northwestern Canada that was never glaciated (e.g., Hultén, 1937; Hopkins, 1967; Schirmermeister et al., 2013). The lack of glaciation and presence of permafrost has resulted in exceptionally well-preserved paleoenvironmental archives that go well beyond the last glacial-interglacial cycle (e.g., Westgate et al., 1990; Sher, 1997; Froese et al., 2009; Minyuk and Ivanov, 2012). In eastern Beringia (Yukon and Alaska), abundant volcanic ash from the Aleutian Arc–Alaska Peninsula and Wrangell volcanic field have been instrumental in providing a means to date and correlate these archives back approximately 2.5 My (e.g., Preece et al., 1999, 2011a, b; Jensen et al., 2008, 2011, 2013; Reyes et al., 2010). However, because few tephra have been recognized in western Beringian sediments, and the idea that prevailing wind directions tend to keep tephra “out of reach” of this region, little work has been done to examine the exposures for their presence. This has had a significant impact on attempting to identify sediments that are beyond the limits of radiocarbon and luminescence dating (e.g., Sher, 1991).

Lake El'gygytyn sediment cores contain eight visible tephra layers that are up to several centimeters thick. The closest known volcanic field to Lake El'gygytyn is more than 300 km away, and volcanoes with known explosive eruptions that have been active during the Late Neogene are ≥ 1000 kilometers away in the Kurile–Kamchatka–Aleutian Arc and Alaska Peninsula (Fig. 1). This implies that this lake holds a record of distant very large magnitude eruptions, which must be distributed widely over western Beringia. In fact, initial analyses of the “T1 tephra” from the pilot core Lz1024 indicate that it correlates to the Rauchua tephra that is thought to be Kamchatka-sourced and present at coastal bluffs on the East Siberian Sea as well as in the Bering Sea (SO201-2-81KL) and western North Pacific (SO201-2-40KL) marine cores (Ponomareva et al., 2013b). The correlation of T1, a tephra only 1 cm thick in the Lake El'gygytyn cores, to the Rauchua tephra, suggests that each individual layer (several > 1 cm thick) has the potential to play an important role in correlating and dating the disparate paleoenvironmental records of this region.

With this study our goal is to provide a detailed geochemical fingerprint for each individual visible tephra layer in the Lake El'gygytyn cores. These data support the stratigraphic correlations that have been made between the five lake cores, and enable us to identify the most likely source regions. The geochemical fingerprints – comprising major, minor- and trace-element geochemistry and lead (Pb) isotopes –

will allow for the future identification of these tephra beds in other depositional settings, where they have been preserved, providing important new chronostratigraphic markers for western Beringia. These tephra will also facilitate independent correlation of the Lake El'gygytyn climate records, particularly to the marine records from the Bering Sea and western North Pacific thus providing a means to compare age models, and allow inter-regional comparisons with sufficiently high temporal precision to discuss climatic synchronicity.

2 Summary of analytical methods

All visible tephra layers identified in the cores were sampled and investigated petrologically prior to analysis by electron probe microanalysis (EPMA) and laser ablation inductively coupled plasma–mass spectrometry (LA-ICP-MS) for compositional fingerprinting, whereby the major-, minor-, trace-element and Pb isotopic composition of single volcanic glass shards was determined (all data provided in the Supplements). Major and minor-element geochemistry was determined using wavelength dispersive EPMA at the University of Alberta, Edmonton (e.g., Jensen et al., 2008, 2011), and at GEOMAR, Kiel (e.g., Ponomareva et al., 2013b). Both laboratories contributed to the recent INTAV (International Quaternary Association's focus group on tephrochronology) comparison of tephrochronology laboratories, which confirmed the high quality of the major-element data produced by both instruments (Kuehn et al., 2011). Additionally, both laboratories run the Lipari obsidian standard as a secondary glass standard to track the quality of calibration and performance through each run (Kuehn et al., 2011). Therefore, both instruments produce data sets that are identical within statistical deviations, and the secondary standard data allow direct comparison of the data sets and help correct for potential issues arising from interlaboratory calibration differences.

Trace-element analyses were performed by LA-ICP-MS at Aberystwyth University, UK, on the glass component of individual shards using ablation craters with diameters in the range of 10–20 μm . Most analyses were carried out on the same sample mounts used for the majority of the EPMA. Detailed methods are described in Pearce et al. (2011); NIST 612 was used for calibration (Pearce et al., 1997), and ^{29}Si as the internal standard, using concentrations determined by EPMA. After rejection of analyses for the incorporation of phenocrysts and/or microlites (e.g., using elevated Ca or Sr for the presence of feldspar) into the ablated material (see analytical protocols established in Pearce et al., 2002, 2004a, 2007), a total of 226 individual shards were analyzed from the eight tephra units in Lake El'gygytyn. Analysis of the MPI-DING ATHO-G reference rhyolitic glass (GeoReM; Jochum et al., 2006) shows that accuracy is good.

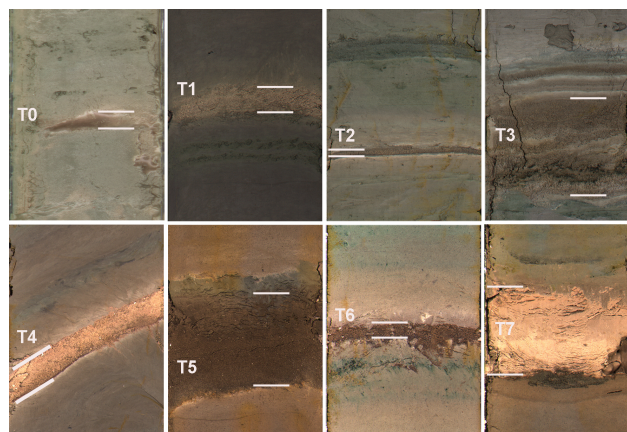


Figure 2. Photos of core sections with detected tephra layers from Lake El'gytyn: T0 (core Lz1024-5 I), T1 (core 5011-1B-1H2), T2 (core 5011-1B-7H2), T3 (core 5011-1B-10H2), T4 (core 5011-1A-19H1), T5 (core 5011-1A-19H2), T6 (core 5011-1A-25H1), T7 (core 5011-1A-35E1). Length of each segment is 12 cm.

Pb isotope analyses were also performed by LA-ICP-MS at Aberystwyth University on a total of 190 individual glass shards from all tephra deposits. Methods used were those described by Westgate et al. (2011a). Analyses were performed using 20 or 30 μm ablation craters, and calibrated against the Pb isotope ratio of the NIST 612 glass, with analytical accuracy monitored by analysis of US Geological Survey (USGS) glass reference materials. At such relatively small crater diameters, precision and accuracy are reduced compared to analysis of larger samples (e.g., by solution ICP-MS or large spot sizes using LA-ICP-MS), but this provides reconnaissance data suitable for preliminary interpretations (Westgate et al., 2011a). Full details of all analytical methods and results are given in Appendix A and the Supplement.

3 The El'gygytyn tephra record

Eight tephra layers were recognized in the main cores (T0–T7), with two present in the pilot cores (T0–T1) (Table 1). The layers are several millimeters up to 6.9 cm thick. In most layers, the top few millimeters of the layers appeared reworked. The ash horizons are composed of 99 % volcanic material as seen in smear slides.

All layers form distinct tephra horizons separated from each other by several centimeters to decimeters of sediment (Fig. 2). Thicknesses were measured in the opened core. Individual tephra in the cores are assigned working names and numbered from top to bottom (tephra T0–T7). Tephra layers have a white to dark brown appearance, and comprise shards that are mainly colorless or pale yellow to brown. Particle morphologies include highly vesicular pumice, tubular and blocky pumice, bubble-wall shards and non-vesicular platy shards; igneous crystals are rare in the tephra layers (Fig. 3).

Table 1. Positions and ages of tephra layers identified in Lake El'gygytyn sediments. Core abbreviations are as follows: 1 – PG1351; 2 – Lz1024; A – 5011-1A; B – 5011-1B; and C – 5011-1C. Core positions in Lake El'gygytyn as in Melles et al. (2011). Age estimates and composite depth from Nowaczyk et al. (2013).

Tephra	Core	Lab ID Kiel	Lab ID Alberta	Depth in 1, 2 (m)	Depth in A (m)	Depth in B (m)	Composite core depth (m)	Thickness (cm)	Age estimate (ka)	Description
T0	1, 2	T0–4 core 1	UA1936, UA1937, UA1938	2.00–2.01 in 1, 2.45–2.46 in 2			2.54–2.55	1	> 45 (Anderson, Lozhkin, 2002)	Whitish fine ash
T1	1, 2, A, B	T1-2 core 2 T1-3/ET1-3 core B	UA1939, UA1940, UA1941	7.83–7.84 in 1, 7.91–7.92 in 2	5.085–5.095	5.148–5.158	7.88–7.89	1	177	Yellowish fine ash
T2	A, B	T2-1 core A T2-2 core B	UA1942, UA1943		24.415–24.420	24.293–24.298	27.52–27.52	0.5	674	Greyish fine ash
T3	A, B	T3-1 core B T3-2 core B T3-3 core B	UA1944, UA1945, UA1946, UA1947		33.090–33.159	32.971–33.027	36.41–36.47	6.9	918	Fine ash at the bottom and top, coarser ash in the middle
T4	A, B	T4-1 core A T4-4 core B	UA1948, UA1949, UA1950, UA1951		56.565–56.576	56.945–56.960	60.79–60.80	1.1	1411	Fine white ash, folded layer
T5	A, B	T5-1 core A T5-3 core B	UA1952, UA1953, UA1954		58.519–58.559	58.433–58.475	62.04–62.08	4	1434	Dark brown fine ash layer
T6	A, B	T6-1 core A T6-2 core B	UA1955, UA1956		75.092–75.099	75.167–75.176	79.25–79.26	0.7–0.9	1775	White, medium to fine ash
T7	A, B, C	T7-1 core B T7-2 core B	UA2014, UA2015		100.981–101.015	100.715–101.7855	104.93–105.00	7	2225	White-grey fine ash layer

Shards are up to 400 µm in diameter (T7), but the median diameter is 40 µm. All of the tephra layers have distinct individual visual and chemical characteristics. Glass shards were analyzed from all horizons. More than 900 electron micro probe analyses on individual glass shards from the eight tephra layers have been obtained from both laboratories. The eight tephra layers have glass shards that are chiefly rhyolite to dacite ($\text{SiO}_2 > 65$ wt.%); one tephra (T5) has glass shards ranging from rhyolite to basaltic andesite (SiO_2 76–54 wt.%) in composition (Fig. 4). The tephra layers are medium- to high-K subalkaline composition (Figs. 4, 5). All of them show high Cl contents > 0.1 wt.% (Fig. 5). Major-oxide composition of shards from individual tephra layers are homogeneous except T3 and T5, which vary systematically (Figs. 4 and 5). The tephra layers resemble each other compositionally, but can be distinguished by their major-element composition, such as SiO_2 and K_2O concentrations (Fig. 5), trace-element composition (Fig. 6), glass shard morphologies, relative stratigraphic position, and age (Table 1).

Twenty-six trace elements were determined from each of the 8 sampled El'gygytyn tephra layers by LA-ICP-MS (see Pearce et al., 2002, 2004a, b, 2007). There are many similarities between each of the tephra layers in their general trace-element contents. Except for T5, all samples show a distinct anomaly at Sr and Eu (Fig. 6, Table 3a) indicating fractiona-

tion of plagioclase. Using a series of bivariate plots of compatible (in silicic magmas) and incompatible elements, it is possible to readily distinguish among most of the tephra layers (Fig. 6). Particularly characteristic are the elements Rb, Ba, Zr, Y, and the HREE (heavy rare earth elements), which can be combined (e.g., to Ba/Y, Ba/Ho, Zr/Ba, Ba/Th) to better show compositional groupings. Tephra layers T1, T5, and T7 have distinctive trace-element compositions readily separated using Rb, Ba, Y and Zr. Tephra layers T0 and T6 can be separated using Nb and Y, and whilst T2, T3 and T4 have very similar trace-element compositions, they can be distinguished by differences in their Ba/Y ratio (Fig. 6). Each of the tephra horizons represents a distinct eruption event or tephra marker, and are described from youngest to the oldest.

3.1 Tephra 0

The youngest tephra visible in the cores was found in the pilot core Lz1024-5 I as a 1 cm thick very fine whitish ash layer (Fig. 2), as well as cores Lz1029-8 and Lz1039-2 at 1.59–1.61 m and 2.12–2.13 m below lake floor, respectively (Juschus et al., 2007). The tephra consists of platy, highly vesicular, colorless glass shards, up to 160 µm, except in the pilot core, where the shards are finer-grained (< 80 µm) (Fig. 3). The glass is chemically homogeneous with 73 wt.%

Table 2. Major-element geochemistry of Lake El'gygytyn tephra from both the University of Alberta (UA) and Kiel laboratories given as averages. **Both** = data from UA and Kiel combined. Analyses of single glass shards are provided in the Supplement Tables S1 and S3. MAX and MIN values for tephra T3 and T5 show the range in composition.

Lab	Tephra		SiO ₂	TiO ₂	Al ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	P ₂ O ₅	F	SO ₃	Cl	Total
UA	T0	MEAN	72.89	0.50	14.01	2.88	0.13	0.57	2.45	4.65	1.77				0.15	100.00
		SD	0.52	0.06	0.26	0.13	0.04	0.06	0.13	0.15	0.07				0.03	0.00
		<i>n</i>	70	70	70	70	70	70	70	70	70				70	70
Kiel	T0	MEAN	72.64	0.50	13.94	2.86	0.13	0.54	2.36	4.97	1.79	0.08	0.03	0.01	0.15	100.00
		SD	0.72	0.04	0.34	0.14	0.04	0.08	0.21	0.09	0.07	0.02	0.03	0.01	0.01	0.00
		<i>n</i>	19	19	19	19	19	19	19	19	19	19	19	19	19	19
Both	T0	MEAN	72.84	0.50	14.00	2.88	0.13	0.57	2.43	4.72	1.78	0.08	0.03	0.01	0.15	100.00
		SD	0.57	0.05	0.27	0.13	0.04	0.06	0.16	0.19	0.07	0.02	0.03	0.01	0.02	0.00
		<i>n</i>	89	89	89	89	89	89	89	89	89	19	19	19	89	89
UA	T1	MEAN	77.77	0.24	12.38	1.22	0.06	0.22	1.20	3.96	2.80				0.16	100.00
		SD	0.39	0.06	0.22	0.13	0.03	0.03	0.08	0.15	0.11				0.03	0.00
		<i>n</i>	73	73	73	73	73	73	73	73	73				73	73
Kiel	T1	MEAN	77.57	0.22	12.37	1.16	0.06	0.20	1.19	4.21	2.79	0.02	0.03	0.01	0.16	100.00
		SD	0.17	0.02	0.10	0.10	0.04	0.02	0.03	0.12	0.08	0.02	0.04	0.01	0.01	0.00
		<i>n</i>	71	71	71	71	71	71	71	71	71	71	71	71	71	71
Both	T1	MEAN	77.67	0.23	12.37	1.19	0.06	0.21	1.20	4.08	2.80	0.02	0.03	0.01	0.16	100.00
		SD	0.32	0.05	0.18	0.12	0.03	0.03	0.06	0.19	0.10	0.02	0.04	0.01	0.02	0.00
		<i>n</i>	144	144	144	144	144	144	144	144	144	71	71	71	144	144
UA	T2	MEAN	74.14	0.41	13.63	2.52	0.10	0.40	1.81	4.25	2.58				0.16	100.00
		SD	1.20	0.09	0.38	0.36	0.04	0.11	0.35	0.20	0.15				0.03	0.00
		<i>n</i>	52	52	52	52	52	52	52	52	52				52	52
Kiel	T2	MEAN	73.21	0.42	13.78	2.61	0.10	0.42	1.88	4.76	2.53	0.06	0.06	0.01	0.16	100.00
		SD	0.95	0.07	0.27	0.32	0.04	0.09	0.26	0.17	0.11	0.02	0.05	0.01	0.01	0.00
		<i>n</i>	49	49	49	49	49	49	49	49	49	49	49	49	49	49
Both	T2	MEAN	73.69	0.41	13.70	2.56	0.10	0.41	1.84	4.50	2.56	0.06	0.06	0.01	0.16	100.00
		SD	1.17	0.08	0.34	0.35	0.04	0.10	0.31	0.31	0.13	0.02	0.05	0.01	0.02	0.00
		<i>n</i>	101	101	101	101	101	101	101	101	101	49	49	49	101	101
UA	T3	MEAN	69.71	0.75	14.57	3.99	0.15	0.94	3.07	4.62	2.05				0.13	100.00
		SD	1.56	0.09	0.34	0.59	0.03	0.23	0.49	0.27	0.19				0.03	0.00
		<i>n</i>	70	70	70	70	70	70	70	70	70				70	70
		MAX	74.05	1.12	15.61	6.07	0.24	1.82	4.76	5.34	2.56				0.22	
		MIN	64.48	0.63	13.65	2.69	0.07	0.59	1.97	3.59	1.49				0.08	
Kiel	T3	MEAN	69.25	0.73	14.63	3.93	0.14	0.88	2.94	5.05	2.08	0.15	0.05	0.03	0.13	100.00
		SD	1.33	0.07	0.47	0.50	0.05	0.22	0.47	0.23	0.16	0.05	0.04	0.02	0.01	0.00
		<i>n</i>	81	81	81	81	81	81	81	81	81	81	81	81	81	81
		MAX	73.31	0.97	16.88	5.16	0.27	1.51	4.16	5.57	2.78	0.33	0.15	0.07	0.16	
		MIN	65.18	0.56	13.51	2.46	0.03	0.37	1.85	3.97	1.76	0.06	0.00	0.00	0.09	
Both	T3	MEAN	69.47	0.74	14.61	3.96	0.15	0.91	3.00	4.85	2.07	0.15	0.05	0.03	0.13	100.00
		SD	1.46	0.08	0.41	0.54	0.04	0.23	0.48	0.33	0.17	0.05	0.04	0.02	0.02	0.00
		<i>n</i>	151	151	151	151	151	151	151	151	151	81	81	81	151	151
		MAX	74.05	1.12	15.61	6.07	0.24	1.82	4.76	5.34	2.56				0.22	
		MIN	64.48	0.63	13.65	2.69	0.07	0.59	1.97	3.59	1.49				0.08	
UA	T4	MEAN	71.33	0.68	14.23	3.42	0.16	0.64	2.20	4.90	2.31				0.13	100.00
		SD	0.33	0.06	0.11	0.11	0.04	0.04	0.10	0.27	0.06				0.03	0.00
		<i>n</i>	118	118	118	118	118	118	118	118	118				118	118
Kiel	T4	MEAN	70.52	0.69	14.34	3.43	0.16	0.66	2.19	5.34	2.34	0.12	0.05	0.02	0.13	100.00
		SD	0.23	0.02	0.12	0.20	0.04	0.03	0.06	0.12	0.04	0.02	0.05	0.01	0.01	0.00
		<i>n</i>	42	42	42	42	42	42	42	42	42	42	42	42	42	42
Both	T4	MEAN	71.00	0.68	14.29	3.43	0.16	0.65	2.21	5.05	2.32	0.12	0.05	0.02	0.13	100.00
		SD	0.48	0.05	0.12	0.15	0.04	0.03	0.08	0.35	0.06	0.02	0.05	0.01	0.02	0.00
		<i>n</i>	160	160	160	160	160	160	160	160	160	42	42	42	160	160
UA	T5	MEAN	62.06	1.08	15.62	7.24	0.19	2.37	5.57	4.20	1.57				0.10	100.00
		SD	5.90	0.26	0.88	2.45	0.05	1.14	2.19	0.46	0.64				0.04	0.00
		<i>n</i>	67	67	67	67	67	67	67	67	67				67	67
		MAX	75.75	1.44	16.70	11.16	0.28	4.00	8.84	5.21	3.60				0.23	
		MIN	53.95	0.30	13.13	1.55	0.07	0.23	1.03	3.18	0.89				0.01	
Kiel	T5	MEAN	59.03	1.19	15.71	8.27	0.20	2.87	6.36	4.41	1.34	0.35	0.03	0.14	0.08	100.00
		SD	4.06	0.20	1.00	1.90	0.06	0.89	1.63	0.43	0.44	0.12	0.04	0.14	0.02	0.00
		<i>n</i>	45	45	45	45	45	45	45	45	45	45	45	45	45	45
		MAX	70.33	1.43	19.71	11.14	0.36	4.05	9.67	5.36	2.50	0.57	0.15	0.84	0.14	
		MIN	53.76	0.54	12.91	3.54	0.07	0.54	2.21	3.51	0.68	0.10	0.00	0.00	0.05	
Both	T5	MEAN	60.84	1.19	15.71	8.27	0.20	2.87	6.36	4.41	1.34	0.35	0.03	0.14	0.08	100.00
		SD	5.43	0.20	1.00	1.90	0.06	0.89	1.63	0.43	0.44	0.12	0.04	0.14	0.02	0.00
		<i>n</i>	112	112	112	112	112	112	112	112	112	45	45	45	112	112
		MAX	75.75	1.44	19.71	11.16	0.36	4.05	9.67	5.36	3.60				0.14	
		MIN	53.76	0.30	12.91	1.55	0.07	0.23	1.03	3.18	0.68				0.05	
UA	T6	MEAN	70.94	0.65	13.95	4.06	0.13	0.64	2.58	4.48	2.43				0.14	100.00
		SD	0.29	0.05	0.10	0.12	0.03	0.04	0.12	0.31	0.07				0.03	0.00
		<i>n</i>	50	50	50	50	50	50	50	50	50				50	50
Kiel	T6	MEAN	70.26	0.66	13.97	4.07	0.14	0.64	2.61	4.87	2.46	0.11	0.04	0.02	0.14	100.00
		SD	0.28	0.03	0.16	0.19	0.04	0.03	0.09	0.17	0.05	0.02	0.04	0.02	0.02	0.00
		<i>n</i>	40	40	40	40	40	40	40	40	40	40	40	40	40	40
Both	T6	MEAN	70.64	0.66	13.96	4.07	0.13	0.64	2.59	4.65	2.45	0.11	0.04	0.02	0.14	100.00
		SD	0.45	0.04	0.13	0.15	0.03	0.04	0.11	0.32	0.06	0.02	0.04	0.02	0.03	0.00
		<i>n</i>	90	90	90	90	90	90	90	90	90	40	40	40	90	90
UA	T7	MEAN	77.26	0.13	12.87	0.71	0.09	0.11	0.63	3.81	4.25				0.14	100.00
		SD	0.27	0.03	0.10	0.04	0.03	0.02	0.03	0.31	0.22				0.02	0.00
		<i>n</i>	48	48	48	48	48	48	48	48	48				48	48
Kiel	T7	MEAN	77.67	0.12	12.96	0.66	0.08	0.09	0.62	3.58	4.02	0.02	0.03	0.01	0.14	100.00
		SD	0.19	0.02	0.09	0.08	0.04	0.02	0.02	0.11	0.14	0.01	0.03	0.01	0.01	0.00
		<i>n</i>	52	52	52	52	52	52	52	52	52	52	52	52	52	52
Both	T7	MEAN	77.48	0.12												

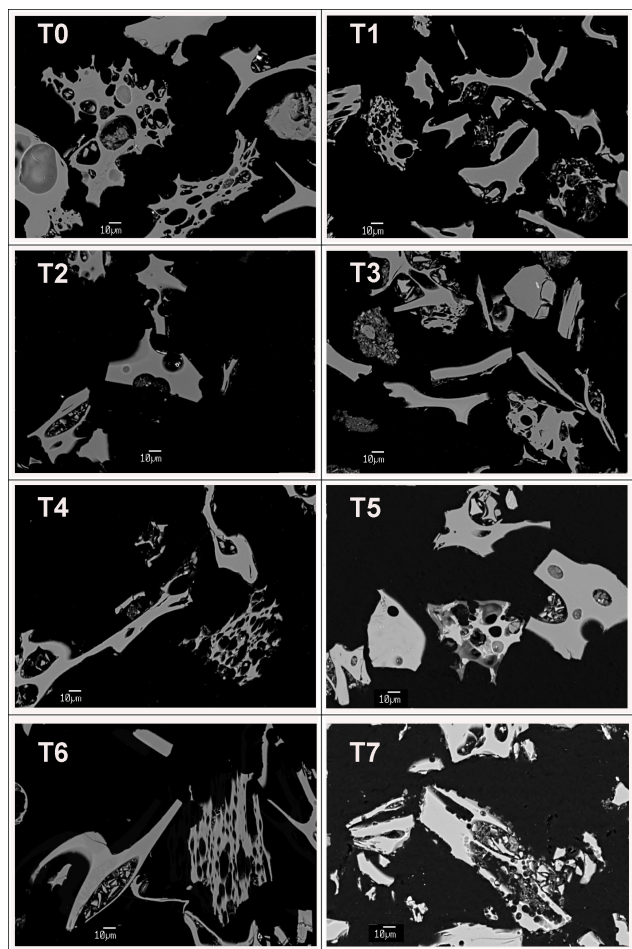


Figure 3. Backscatter electron images of characteristic glass shards from tephra layers T0–T7 from Lake El'gygytyn. Scale bar is 10 µm.

SiO₂ (Fig. 5, Table 2), and with distinct Y and Nb values (Fig. 6d).

3.2 Tephra 1

T1 comprises tephra beds in the pilot core Lz1024-9 and in cores 5011-1B-1H-2 and 5011-1A-1H-3. The tephra is present as an approximately 1 cm thick discrete ash layer of extremely fine to very fine grain size (Fig. 2). It consists mainly of colorless pumiceous shards with a median size of about 40 µm, and some larger (< 100 µm), colorless, highly vesicular shards (Fig. 3). The composition of T1 is homogeneous. With high SiO₂ content of 78 wt.% it is the most silicic tephra (comparable to T7) of the eight layers (Figs. 4, 5; Table 2), and it has characteristic Y, Rb and Ba contents (Fig. 6).

3.3 Tephra 2

T2 occurs in cores 5011-1A-8H-1 and 5011-1B-7H-3 as a 0.5 cm fine greyish ash (Fig. 2). Glass shards are ≤ 100 µm

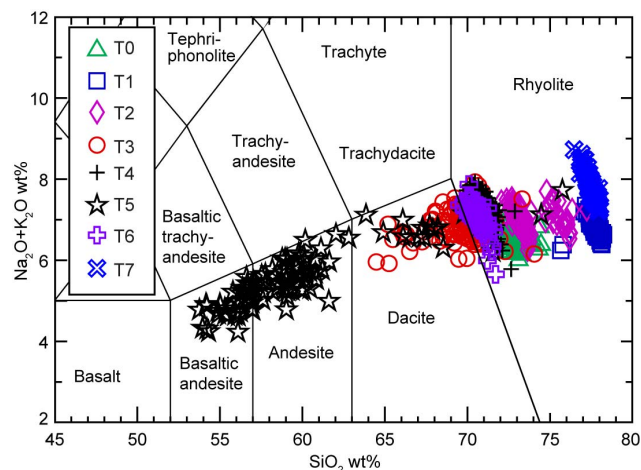


Figure 4. TAS (total alkali silica) classification diagram with fields according to Le Bas et al. (1986). All data are single shard glass analyses normalized to 100 %. Data include both sets analyzed in Edmonton and Kiel.

in diameter, colorless, and platy with round vesicles (Fig. 3). The glass composition is bimodal, comprising rhyolitic to rhyodacitic compositions (77 to 72 wt.% SiO₂; Figs. 4 and 5, Table 2).

3.4 Tephra 3

T3 occurs in cores 5011-1B-10H-2 and 5011-1A-11H-1, and is a 6 cm thick graded ash (Fig. 2). The layer is finer grained at the bottom and top, and coarser in the middle, where shards are up to 150 µm in diameter. It consists of colorless, vesicular shards, with rare brown glass. The shards have variable SiO₂ contents from 74.0 to 64.5 wt.% (Fig. 4). This, and the variation in MgO concentration from 0.4 to 1.8 wt.% and of FeO from 2.5 to 6.1 wt.%, suggests an eruption from a chemically zoned magma chamber (Figs. 4, 5; Table 2).

3.5 Tephra 4

This prominent, white, ~ 1 cm thick tephra layer (Fig. 2) occurs in sections of cores 5011-1B-18H-2 and 5011-1A-19H-1/18H-3, which are interpreted as representing a sequence of multiple turbidites, subsequently folded by a larger mass movement (Melles et al., 2012; Sauerbrey et al., 2013). T4 is reworked within these deposits and appears up to three times in the core. This tephra layer consists of colorless tubular pumice and pumiceous and vesicular glass shards (Fig. 3). The median grain size is about 60 µm; the largest shards are up to 250 µm. T4 is chemically homogeneous, with a rhyolitic composition of 71 wt.% SiO₂ (Fig. 4, Table 2).

Table 3. Average trace-element composition of single glass shards from Lake El'gygytyn tephra collected by LA-ICP-MS in Aberystwyth. All concentrations are in ppm, unless otherwise stated.

Sample number	Eruption ID	Crater diameter µm	Int'l std – SiO ₂ , wt %	Rb	Sr	Y	Zr	Nb	Cs	Ba	La	Ce	Pr	Nd	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu	Hf	Ta	Pb	Th	U																													
UA1936 avg. s.d. n = 21	T0	All 20 µm	72.79	37.2 3.3	210 21	70.0 8.6	244 24	4.27 0.45	1.61 0.16	588 49	15.8 2.1	32.5 2.4	5.07 0.50	26.4 2.9	8.20 1.80	1.86 0.37	10.2 1.1	1.78 0.31	11.6 1.6	2.58 0.40	7.89 1.02	1.17 0.19	1.17 1.01	7.75 0.92	1.29 0.77	7.91 1.01	0.43 0.11	15.5 6.1	3.56 0.54	1.57 0.33																												
UA1939 avg. s.d. n = 24	T1	All 20 µm	77.78	52.8 11.2	154 4.5	37.9 28.0	253 371	4.41 6.87	1.80 2.39	891 895	19.4 23.4	34.9 44.7	4.61 6.96	20.1 33.4	5.03 8.84	0.85 1.72	5.69 9.90	0.86 1.72	6.06 1.09	1.39 2.55	4.64 7.56	0.78 1.09	0.84 8.54	1.92 1.12	0.84 1.16	0.71 0.71	32.4 32.4	8.83 8.83	3.50 3.50																													
UA1942 avg. s.d. n = 16	T2	All 20 µm	74.17	58.7 11.4	203 32	68.0 8.1	371 60	6.87 0.64	0.59 0.63	895 98	23.4 4.3	44.7 5.4	6.96 1.00	33.4 6.1	8.84 2.10	1.72 0.32	9.90 2.29	1.72 0.40	2.7 0.55	1.80 2.20	0.90 0.72	0.26 0.92	0.36 1.15	0.15 0.12	0.49 0.51	0.25 0.12	26.8 26.8	3.61 3.61	1.30 1.30																													
UA1943 avg. s.d. n = 9	T2	All 20 µm	74.11	56.7 6.7	186 27	63.0 5.4	344 6.7	6.67 2.43	868 0.51	868 47	22.5 2.0	48.4 10.8	6.41 0.69	5.4 1.24	32.8 0.33	8.58 1.71	10.3 1.03	1.56 0.23	9.95 1.37	2.50 0.52	7.29 0.86	1.07 1.12	1.22 0.96	1.22 0.86	1.08 0.66	0.62 0.14	22.4 22.4	7.81 7.81	3.09 3.09																													
T2 (Overall average) s.d. n = 25			58.0 13.0	197 45.5	66.2 82.1	361 15.0	6.80 1.42	2.40 0.66	885 166	23.08 16.6	46.0 10.7	6.76 1.55	33.2 8.29	8.75 2.40	1.71 0.47	10.1 2.72	1.66 0.49	10.9 3.05	2.53 0.77	7.46 1.95	1.08 0.25	7.46 1.95	1.08 0.25	8.24 2.76	1.15 0.31	11.3 3.32	0.69 0.24	28.8 21.4	8.47 5.23	3.35 1.22																												
UA1944 avg. s.d. n = 20	T3	20/14 µm	68.51	40.6 6.28	272 63.6	72.9 8.45	333 40.5	6.74 1.35	1.74 0.38	768 81.0	23.0 4.21	44.3 4.94	7.43 2.79	34.1 3.50	9.82 2.49	2.40 0.40	11.0 1.90	1.86 0.32	12.80 2.89	2.77 0.53	8.91 1.59	1.30 0.31	1.30 1.04	7.97 0.93	1.31 1.72	0.59 0.24	15.20 2.32	5.27 0.86	1.86 0.33																													
T3-2 avg. s.d. n = 10	T3 (Kiel Mount)	All 20 µm	68.82	45.7 3.49	240 30.7	71.6 7.85	317 28.6	5.99 0.20	1.74 0.19	740 58.5	20.1 1.63	43.6 2.61	6.77 0.56	34.9 3.59	9.58 2.05	2.03 1.24	1.91 1.94	1.91 0.16	11.7 1.70	2.64 0.30	7.54 0.67	1.24 0.92	7.64 1.02	1.19 0.92	9.33 1.07	0.50 0.08	13.9 17.7	4.21 0.54	1.55 0.11																													
T3 (Overall average) s.d. n = 30			42.9 6.58	264 55.0	73.38 7.99	33.8 37.3	6.57 1.13	1.76 0.34	768 76.9	22.3 3.67	44.6 4.21	7.30 2.29	34.8 3.46	9.86 2.02	2.04 0.36	11.2 2.92	1.90 0.28	12.6 2.53	2.76 0.46	8.54 1.43	1.29 0.69	0.92 0.28	0.92 1.00	0.15 0.28	1.07 1.60	0.08 0.20	14.9 2.47	4.97 0.89	1.78 0.30																													
UA1947 avg. s.d. n = 26	T4	20/14 µm	71.39	36.7 4.7	287 22	72.3 5.8	285 22	7.11 0.83	1.57 0.65	802 46	24.9 3.8	47.0 4.6	7.48 0.82	36.5 3.6	9.63 1.49	2.10 0.36	11.5 2.2	1.78 0.28	12.1 1.73	2.58 2.35	7.98 7.34	1.16 1.12	7.98 7.30	1.25 1.02	9.26 8.55	0.68 0.61	16.7 17.0	4.67 3.58	2.05 1.96																													
T4-1 avg. s.d. n = 10	T4 (Kiel mount)	All 20 µm	71.39	38.4 1.96	281.0 20.5	66.4 5.77	286 24.1	6.91 0.36	1.38 0.14	808 50.5	21.6 1.92	46.1 3.04	7.05 0.41	36.9 3.34	9.39 1.20	2.22 0.27	11.1 1.59	1.73 0.24	11.0 1.09	2.35 0.26	7.34 0.81	1.12 0.15	7.30 0.96	1.02 0.14	8.55 0.90	0.61 0.11	17.0 1.30	3.58 0.44	1.96 0.15																													
T4 (Overall average) s.d. n = 36			37.19 4.20	285 21.5	70.6 6.33	285 22.2	70.6 0.56	1.52 0.42	804 46.3	24.0 3.65	46.8 4.24	7.36 3.65	36.6 3.51	9.56 2.13	2.13 1.41	11.4 2.05	1.77 0.27	11.8 1.63	2.52 0.32	7.80 1.15	1.15 0.23	0.93 0.11	0.71 0.20	1.47 0.20	1.06 0.18	0.66 0.20	16.8 2.69	4.36 1.40	2.02 0.52																													
UA1952 avg. s.d. n = 27	T5	20/14 µm	variable	19.1 3.26	483 93.4	39.3 6.50	158 39.0	4.36 0.93	0.73 0.22	502 71.7	15.4 3.85	31.7 4.24	4.91 0.70	24.8 3.08	7.18 1.29	1.84 0.36	7.65 1.11	1.22 0.29	7.67 1.40	1.66 0.35	4.76 0.91	0.74 0.19	4.95 1.16	0.74 0.20	5.04 1.21	0.35 0.17	11.1 1.57	2.02 0.42	1.09 0.23																													
UA1954 avg. s.d. n = 23	T5	20/14 µm	variable	17.4 81.8	512 5.00	36.3 23.8	148 0.71	0.92 0.34	0.12 0.56	470 56.1	13.0 2.06	28.1 3.52	22.7 0.78	64.9 3.05	9.38 1.49	0.98 0.30	7.52 1.19	1.08 0.25	6.95 1.19	1.51 0.29	4.27 0.88	0.67 0.14	0.67 0.82	0.46 0.18	4.61 1.22	0.30 0.13	9.73 1.33	1.88 0.39	1.00 0.22																													
T5 (Overall average) s.d. n = 50			18.3 3.16	496 89.6	37.9 6.06	153 33.4	4.25 0.85	0.72 0.28	486 71.1	14.3 3.40	30.1 4.74	23.8 3.76	6.86 3.25	1.90 1.43	2.38 1.35	7.59 0.34	1.16 0.11	1.77 0.28	11.8 1.36	2.52 0.33	7.80 0.93	1.15 0.21	0.93 0.19	0.71 0.12	1.45 1.24	0.45 0.16	1.61 1.05	1.96 0.41	1.05 0.23																													
T6-2 avg. s.d. n = 25	T6 (Kiel Mount)	20/10 µm	70.97	49.8 1.8	248 28	64.4 7.9	341 51	9.53 0.90	1.81 0.14	760 70	23.1 2.8	48.6 4.2	7.30 0.79	35.2 5.3	9.25 1.39	1.86 0.24	1.98 1.53	1.65 0.25	10.8 1.8	2.32 0.36	7.06 1.15	1.05 0.19	1.05 1.15	1.11 0.17	9.64 1.77	0.73 0.16	23.4 4.3	4.98 0.97	2.33 0.36																													
UA2014 avg. s.d. n = 15	T7	All 20 µm	77.17	94.6 2.9	80.4 5.5	28.2 2.1	123 12	6.93 0.35	3.75 0.31	1089 43	28.8 2.0	51.8 2.2	6.50 0.44	25.0 2.8	5.17 0.84	0.71 0.20	4.43 0.73	0.75 0.14	4.53 0.84	1.06 0.14	3.17 0.52	0.50 0.11	0.50 0.64	0.62 0.21	5.29 0.75	0.69 0.11	22.9 3.3	10.9 1.0	4.16 0.36																													
Reference material analyses																																																										
ATHO-G, this study, 20 µm analyses s.d. (n = 10)																																63.5 2.2	93.9 4.2	95.4 6.7	506 27	62.6 2.2	1.01 0.11	537 3.2	54.2 3.2	122 0.4	14.5 3.2	62.8 0.9	13.9 0.3	2.69 0.1	15.7 1.4	15.7 0.13	2.46 1.4	16.5 1.1	3.47 0.32	10.3 0.8	1.54 1.0	10.2 0.7	1.58 0.15	13.5 0.20	3.87 0.35	7.51 0.40	2.41 0.14	
ATHO-G reference conc. GeoRem s.d																																65.3 3	94.1 2.7	94.5 3.5	512 20	62.4 2.6	1.08 0.11	547 16	55.6 1.5	121 4	14.6 0.4	60.9 2	14.2 0.4	2.76 0.1	15.3 0.7	2.51 0.2	2.51 0.08	16.2 0.7	3.43 0.43	10.3 0.5	1.52 0.07	10.5 0.4	1.54 0.05	13.7 0.3	3.9 0.2	5.67 0.62	7.4 0.27	2.37 0.12
Selected data for widespread Alaskan tephra																																																										
Aniakchak 1643 BC																																71.4	251	44.9	227	13.0	n.d.	725	30.9	57.0	8.25	35.3	8.61	2.30	9.29	1.45	8.95	1.91	5.04	0.78	5.55	0.85	0.85	6.07	0.70	13.2	9.01	2.56
Aniakchak 1643 BC																																66.5	199	46.3	267	15.5	3.11	861	26.4	57.4	6.96	30.6	7.66	1.71	6.54	1.27	7.74	1.82	4.84	0.73	4.78	0.74	n.d.	1.03	n.d.	6.13	2.84	
UT-1																																103	138	34.7	259	n.d.	4.37	923	24.7	52.8	5.83	23.7	5.87	0.87	4.83	0.90	5.34	1.27	3.44	0.56	3.60	0.57	7.00	n.d.	n.d.	9.81	4.28	
UT613																																99.3	151	29.9	233	8.95	4.47	1003	26.2	46.4	6.66	25.5	5.50	1.07	5.83	0.94	5.47	1.20	3.49	0.55	3.84	0.62	7.46	0.76	13.3	10.2	5.13	
Dawson tephra																																85.0	73	39.6	299	7.31	5.13	920	19.7	40.6	5.86	23.4	5.82	0.95	6.33	1.07	6.76	1.46	4.31	0.76	4.64	0.80	8.43	0.50	n.d.	7.50	4.02	

Table 4. Average Pb isotopic values collected on single glass shards from Lake El'gygytyn tephra. Avg. = average; SD = standard deviation; n.r. = not reported; RSD% = relative standard deviation given as percentage (+ std dev/mean \times 1000).

Samples	208 Pb avg. cps	Avg. Pb ppm	206/ 204 avg	207/ 204 avg.	208/ 204 avg.	206/ 208 avg.	207/ 208 avg.	206/ 204 SD	207/ 204 SD	208/ 204 SD	206/ 208 SD	207/ 208 SD
T0, 30 μ m	25941	15,5	19,063	16,026	39,518	0,4826	0,4055	0,870	0,846	2,024	0,0071	0,0071
RSD %, $n = 19$								4,6 %	5,3 %	5,1 %	1,5 %	1,8 %
T1, 20 μ m	16757	18,4	18,750	15,760	38,683	0,4847	0,4075	1,570	1,275	3,252	0,0065	0,0056
RSD %, $n = 15$								8,4 %	8,1 %	8,4 %	1,3 %	1,4 %
T2, 20 μ m	19016	22,4	18,951	16,090	39,299	0,4822	0,4093	1,495	1,303	2,905	0,0089	0,0091
RSD %, $n = 19$								7,9 %	8,1 %	7,4 %	1,8 %	2,2 %
T3, 30 μ m	25596	21,1	18,281	15,451	37,877	0,4825	0,4080	1,317	1,029	2,550	0,0067	0,0052
RSD %, $n = 20$								7,2 %	6,7 %	6,7 %	1,4 %	1,3 %
ET3, 30 μ m	44817	21,1	18,472	15,679	38,436	0,4806	0,4079	0,617	0,554	1,260	0,0045	0,0044
RSD %, $n = 20$								3,3 %	3,5 %	3,3 %	0,9 %	1,1 %
T4, 20 μ m	20107	15,9	18,116	15,236	37,553	0,4824	0,4059	1,172	0,899	2,426	0,0063	0,0058
RSD %, $n = 17$								6,5 %	5,9 %	6,5 %	1,3 %	1,4 %
T5, most 30 μ m, some 20 μ m	24498	8,76	19,193	16,107	39,548	0,4853	0,4073	1,022	0,773	1,961	0,0072	0,0026
RSD %, $n = 15$								5,3 %	4,8 %	5,0 %	1,5 %	0,6 %
T6, most 20 μ m, some 30 μ m	59358	23,4	19,090	15,752	38,884	0,4909	0,4052	1,216	0,877	2,211	0,0126	0,0056
RSD %, $n = 19$								6,4 %	5,6 %	5,7 %	2,6 %	1,4 %
T7, 20 μ m	26639	22,9	18,133	15,286	37,376	0,4854	0,4090	0,867	0,957	2,016	0,0116	0,0145
RSD %, $n = 19$								4,8 %	6,3 %	5,4 %	2,4 %	3,5 %
Aniakchak UT2011, 20 μ m	18554	12,0	19,012	15,646	38,545	0,4933	0,4060	1,368	1,106	2,770	0,0081	0,0077
RSD %, $n = 18$								7,2 %	7,1 %	7,2 %	1,6 %	1,9 %
Reference material analyses												
GeoReM BCR-2G avg. (95 % CL)		11 \pm 1	18,819	15,692	38,699	0,4863	0,4054	0,007	0,006	0,02	0,0005	0,0005
BCR-2G this study 30 μ m	27293		18,739	15,571	38,783	0,4832	0,4015	0,830	0,707	1,771	0,0027	0,0051
RSD %, $n = 10$								4,4 %	4,5 %	4,6 %	0,6 %	1,3 %
Accuracy this study/GeoReM			0,42 %	0,77 %	−0,22 %	0,63 %	0,95 %					
GeoReM TB1-G (avg. $n = 2$)		18.4 \pm 1.6	18,333	15,552	38,611	0,4748	0,4028	0,001	0,001	0,004	n.r.	n.r.
TB-1G this study 30 μ m	78931		18,374	15,640	38,681	0,4750	0,4043	0,287	0,295	0,572	0,0034	0,0034
RSD %, $n = 10$								1,6 %	1,9 %	1,5 %	0,7 %	0,8 %
Accuracy this study/GeoReM			−0,23 %	−0,57 %	−0,18 %	−0,05 %	−0,38 %					

3.6 Tephra 5

Tephra 5 occurs in cores 5011-1A-19H-2 and 5011-1B-19H-1. This distinctive 4 cm thick brownish tephra (Fig. 2) is extremely fine-grained with median and maximum shard sizes of 40 and 80 μ m, respectively. It is comprised of brown to greenish platy tricusate glass shards, and highly vesicular, colorless shards (Fig. 3). The tephra has the broadest compositional range of all the tephra beds, with SiO₂ spanning from 76 to 54 wt.% (Figs. 4, 5). The variation in MgO (0.2–4.0 wt.%) and FeO (1.6–11.1 wt.%) content suggests that this tephra was erupted from a chemically zoned magma chamber. T5 has the flattest REE profile with the least pronounced Eu anomaly (Fig. 6).

3.7 Tephra 6

Tephra 6 occurs in cores 5011-1A-25H-1 and 5011-1B-25H-2. It is a 7–9 mm thick (Fig. 2), medium to fine ash with shards up to 300 μ m that is comprised of colorless glass shards ranging in morphology from stretched and blocky pumice to platy shards (Fig. 3). Blocky pumice have numerous small, round, evenly sized vesicles. Few brown blocky shards were seen, but all displayed the same morphology as the colorless shards. T6 has a homogeneous major-element composition with SiO₂ at 70.6 ± 0.5 wt.% and relatively high Na₂O content of 4.7 ± 0.3 wt.%. It also has the highest Nb

and Ta concentrations of the eight tephtras, and shows the highest HREE and a relatively flat REE profile, similar to T2, T3 and T4 (Fig. 5)

3.8 Tephra 7

Tephra 7 is present in the three cores 5011-1A-35H-1, 5011-1B-36H-1 and 5011-1C-5A-1. Shards in this 7 cm thick white to grey tephra horizon (Fig. 2) are up to 500 μ m in diameter. They are colorless and comprised of stretched and blocky pumice. Blocky shards contain many evenly distributed vesicles that are generally ~ 1 μ m in diameter. The glass margins show extensive scalloping along the edges. The composition of T7 is homogeneous, with high SiO₂ content (77.5 ± 0.3 wt.%), and with the highest K₂O content (4.0 ± 0.1 wt.%) of the El'gygytyn tephtras. T7 is also distinct in its trace-element composition, with the highest Ba, Rb, Th, and LREE (light rare earth elements), and the lowest Sr and HREE, giving it the steepest REE patterns (Fig. 6).

4 Age of tephra layers

The three parallel sediment cores from Lake El'gygytyn were spliced into a composite core, and a high-resolution age model was developed using a multi-proxy approach (Nowaczyk et al., 2013). Magnetostratigraphy provides the

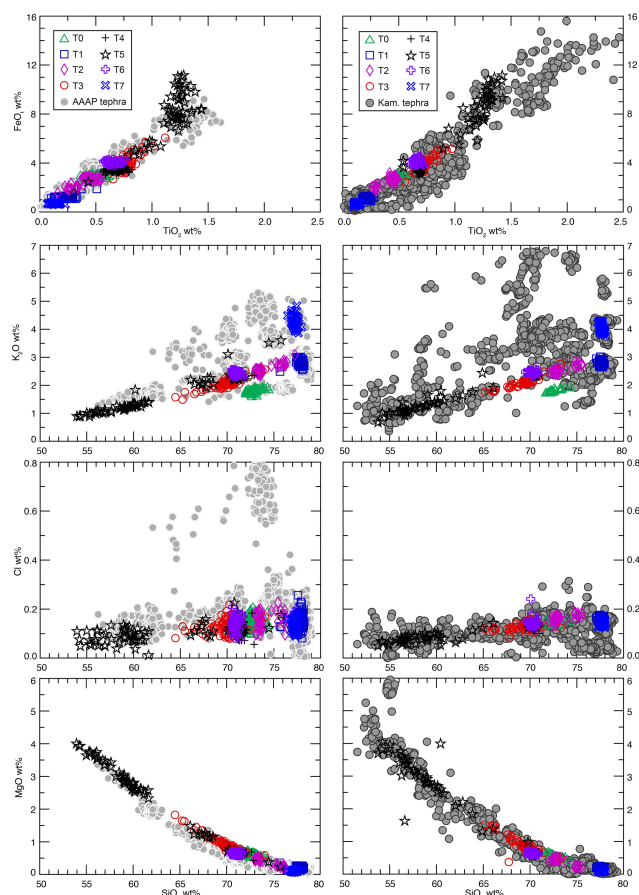


Figure 5. Harker diagrams for major-element geochemistry of T0–T7 in comparison to tephra from Kamchatka and the AAAP (Aleutian Arc–Alaska Peninsula). All data are single shard glass analyses normalized to 100 %. Plots on the left compare data collected in Edmonton to a data set of AAAP tephra that have higher similarity coefficients (> 0.90) to the tephra examined in this study. Compiled AAAP data were analyzed in Edmonton and are from the internal database, but are limited to tephra that have previously been reported in the literature. Plots on the right compare the major-element geochemical analyses collected in the Kiel laboratory to Pleistocene Kamchatkan eruptions. Although T0–T7 all plot within the fields designated by the AAAP and Kamchatkan data, they tend to plot more clearly with Kamchatkan data. This is particularly the case for the mafic oxides (e.g., MgO).

tie points for the basic age model (Haltia and Nowaczyk, 2013). To further refine the age model, nine additional stratigraphic parameters were synchronously tuned (Nowaczyk et al., 2013). Ages of the tephra beds in this study were determined from the resulting age–depth model (Nowaczyk et al., 2013) (Table 1).

Independent ages for the tephra layers are not yet available. Due to the age and composition of the tephra layers (low to medium K and datable phenocrysts being absent) direct dating of the tephra layers is difficult. Vesicular glass shards of 63 μm and smaller have been dated us-

ing a laser Ar–Ar technique; however, analyses carried out on tephra layers from deep sea cores using this approach have been problematic (e.g., Hall and Farrell, 1995). An attempt to date the glass shards of some of the tephra layers from Lake El'gygytyn using the glass fission-track method is currently being carried out and may in the future provide independent age constraints (J. A. Westgate, personal communication, 2013).

5 Potential sources of the tephra layers

5.1 Continental volcanoes versus island arcs

Based on major-element geochemical distribution, the tephra layers from Lake El'gygytyn are characterized by a sub-alkaline composition (Fig. 4), and likely originate from a subduction-related tectonic setting rather than from intra-continental vents. Even though intracontinental volcanoes in Chukotka are present ~ 300 km from Lake El'gygytyn (Fig. 1), much of their activity was limited to the Cretaceous (Akinin and Miller, 2012); only few centers have been active over the Quaternary (Pevzner et al., 2011). Moreover, these volcanoes are associated with mafic lava fields, dykes and/or cinder cones (Laverov et al., 2006), and are thus unlikely sources for the El'gygytyn tephra.

Trace-element geochemistry of the Lake El'gygytyn tephra layers rather suggests that the eruptions originate from subduction-related volcanoes. Magmas formed in non-subduction-related settings (continental and oceanic hot-spots and rifts) are expected to have low ratios of LILE (large ion lithophile elements, e.g., K, Rb, Ba), Th, U and LREE (e.g., La, Ce) to HFSE (high-field-strength elements, e.g., Ta, Nb, Zr) (Gill, 1981; Pearce and Parkinson, 1993). For example, typical characteristics of non-subduction-related rocks are $\text{Th}/\text{Nb} < 0.01$, $\text{La}/\text{Nb} < 2$, and $\text{U}/\text{Nb} < 0.03$ (Pearce and Parkinson, 1993). The trace-element data for the Lake El'gygytyn tephra show strong relative enrichment in LILE (Rb, Ba, Cs) in comparison to similar incompatible LREE (La, Ce) and HFSE (Nb, Ta). Therefore, the major- and trace-element composition of these tephra beds suggests an island arc origin (Figs. 4, 5, 6).

The most likely source for these tephra beds are the many volcanoes that collectively comprise the Kurile–Kamchatka–Aleutian Arc and Alaska Peninsula, more than 1000 km away from Lake El'gygytyn (Fig. 1). Similarly, the Pb isotope data for these tephra beds fall within the compositional envelopes defined by known Pb isotopic values from Kamchatkan and Alaskan volcanoes (Fig. 7). Collectively, the major, trace and isotope geochemical data suggest that the Kurile–Kamchatka–Aleutian Arc and Alaska Peninsula volcanoes are the most likely sources for the eruptions, and these areas have been active for more than 2 million years (e.g., Siebert and Simkin, 2002; Preece et al., 1999).

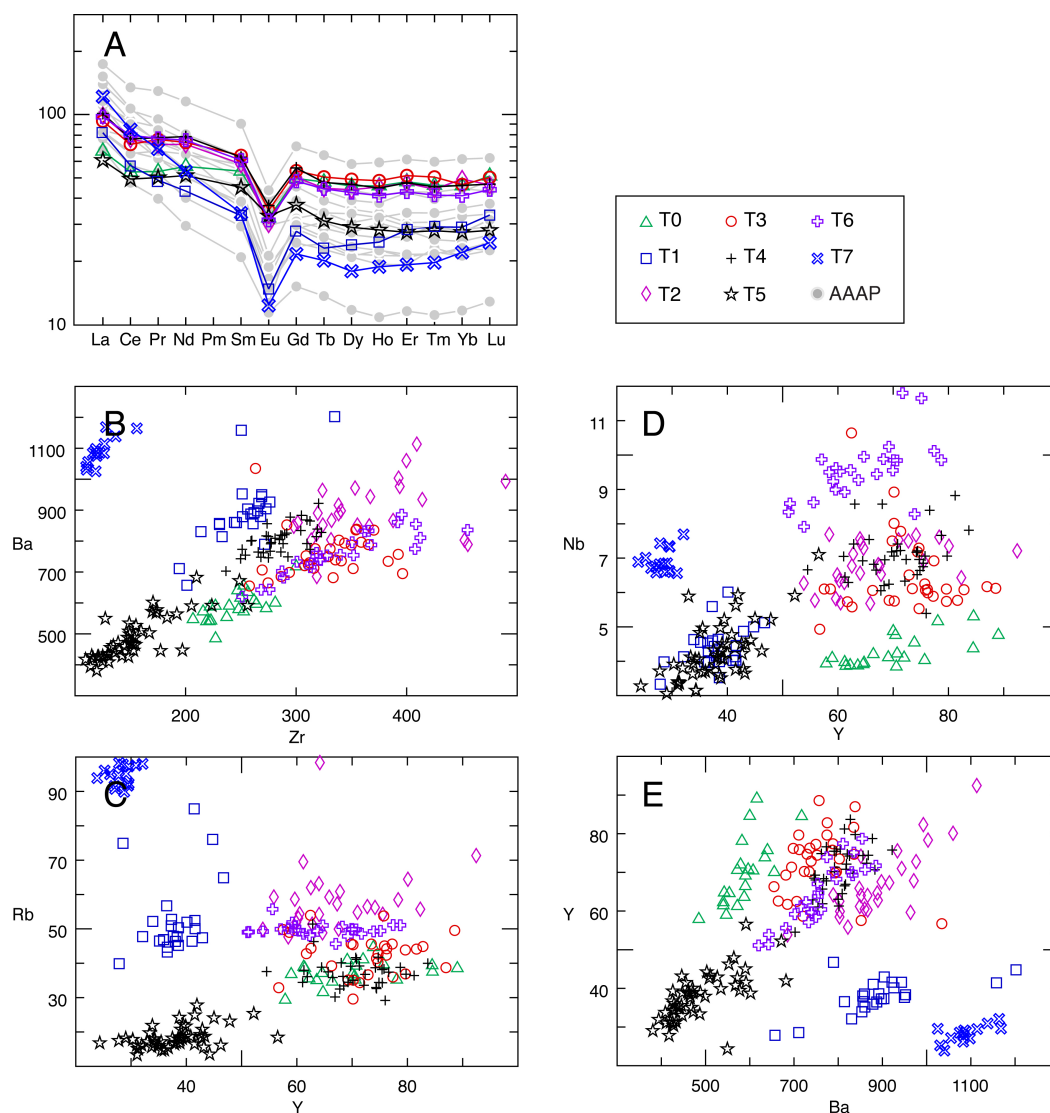


Figure 6. Plots of the trace-element composition of single glass shards for T0–T7. (A) Spider diagram comparing the average REE abundances of each tephra to known AAAP (Aleutian Arc–Alaska Peninsula) Pleistocene tephra (Pearce et al., 2004b; Preece et al., 2011a, b; Jensen et al., unpublished data). Data are normalized to chondrite after Sun and McDonough (1989). (B–E) Plots of trace elements that allow differentiation between each tephra. (B) Zr–Ba, (C) Y–Rb (D), Y–Nb (E) Ba–Y. Tephra T1, T5 and T7 are compositionally distinct, and can be clearly separated based on a combination of Ba, Rb, Zr and Y (B and C). T0 and T6 form separate clusters when compared on their Y and Nb concentrations (D). T2, T3 and T4 are more similar, but T3 shows a lower Zr/Ba ratio than T2 and T4 (B), and these 3 samples all display different Ba/Y ratios (E), which can be used to separate them.

5.2 Island arc: Alaska versus Kamchatka volcanoes

The predominantly explosive volcanism of the NW Pacific area is related to the subduction of the Pacific Plate under the Eurasian and North America plates, forming the Kamchatka–Kurile–Aleutian Arc, and the Alaska Peninsula. The major- and trace-element geochemistry of Lake El'gygytyn tephra falls within the compositional range of both Alaskan and Kamchatkan tephra beds (Figs. 5, 6).

5.2.1 Alaskan volcanoes

Alaska is home to over 130 volcanoes, most of which are located in the Aleutian Arc–Alaska Peninsula (AAAP) and the Wrangell volcanic field (e.g., Nye et al., 1998; Fig. 1). Although the Wrangell volcanic field has been active over the last ~ 25 My (Richter et al., 1990), and the AAAP for ~ 40 My (Jicha et al., 2006), most eruptions from these volcanoes are known only by their widespread tephra because most of their proximal deposits became glacially eroded over the past ~ 3 My.

Much of this distal tephra record is preserved on the terrestrial landscape within widespread loess deposits of eastern Beringia (e.g., Westgate et al., 1990). To date, well over 100 widely distributed distal tephra beds have been identified and characterized from the interior of Alaska and Yukon (e.g., Preece et al., 1999, 2000, 2011a; Jensen et al., 2008, 2013). Although distances from AAAP volcanoes to Lake El'gygytyn are great, generally between 1600 and 2000 km, tephra with comparable distributions have been documented from these volcanoes in the past (e.g., Jensen et al., 2011; Preece et al., 2011b). This suggests that the AAAP could be a potential source for tephra at the study site. With the Wrangell volcanic field downwind and over 2000 km from Lake El'gygytyn, it seems unlikely that visible tephra from these volcanoes would be present in the lake. However, they were not excluded from geochemical comparisons.

Comparison of major- and minor-element geochemistry of Alaskan tephra to the tephra recovered from Lake El'gygytyn was two-fold. Firstly, all El'gygytyn tephra were run through the University of Alberta Tephrochronology Lab searchable database to query for potential correlations. The database uses a similarity coefficient (SC; Borchart et al., 1972) as the basis for its search. Since some of the tephra beds have large geochemical trends, where averages can be misleading, the standard deviation of SiO_2 wt.% is also used in this search. For tephra beds that may have more than one mode, the average of each population is run through the database independently. Generally, a SC of > 0.95 demonstrates some similarity between two samples though does not necessarily indicate a correlative bed. Tephra beds with SCs > 0.90 were further examined visually through Harker diagrams. When the Lake El'gygytyn tephra were run through the database, no values over 0.93 were obtained for the Alaskan tephra except for T1, which had SCs of 0.94 to 0.96 with the Alyeska Pipeline, Gold Run, P13 and other similar, high SiO_2 wt.% rhyolites (e.g., Westgate et al., 2009; Jensen et al., 2013; Supplement Table S2).

The detailed comparison of major-element geochemistry between the Alaskan and Lake El'gygytyn tephra suggests there are no obvious correlations. However, to test the database results and explore the overall characteristics of the unknowns in comparison to tephra recovered in Alaska, a series of prominent widely distributed tephra beds – including Aniakhak, Dawson, VT, Old Crow, Gold Run, GI, PAL, HP and PA – were plotted with the El'gygytyn tephra (e.g., Kaufman et al., 2012; Preece et al., 1999, 2011a, b; Demuro et al., 2008; Westgate et al., 2009; Jensen et al., 2013; Fig. 5). These tephra beds were chosen based on their widespread distribution; they represent the most widely distributed tephra known in Alaska, and some have chemistries broadly similar to the unknowns (e.g., Supplement Table S2). These beds also allow to query the Alaskan tephra to capture variability in terms of age (ca 2 My to 3.6 ky old), and major-element geochemistry (basaltic-andesite to rhyolite). Results indicate that the major-element geochemistry of Lake

El'gygytyn tephra tend to fall within the compositional range exhibited by this selection of Alaskan tephra (Fig. 5). Some slight systematic differences appear to be present: at less than ~ 75 SiO_2 wt.%, MgO, FeO and CaO tend to be higher at any given SiO_2 wt.% than the Alaskan beds, while K_2O and Al_2O_3 wt.% tend to be a little lower. In conclusion, while the major-element geochemistry shows some similarities between the Alaskan and Lake El'gygytyn tephra, it provides no conclusive evidence to support an Alaskan source for the beds. Trace-element geochemistry comparisons between Alaska tephra and T0–T7 are also inconclusive, showing complete overlap (Fig. 6).

The lead isotope data (^{206}Pb , ^{207}Pb and ^{208}Pb) offer the most in terms of delineating (or precluding) a possible Alaskan source. The isotope data are broadly similar for all of the tephra beds except T6, which differs from the other Lake El'gygytyn tephra by having the highest $^{206}\text{Pb}/^{208}\text{Pb}$ and the lowest $^{207}\text{Pb}/^{208}\text{Pb}$ ratios. When compared with recent Pb isotope data compiled from Kamchatka and Alaska volcanics, the majority of samples sit close to or within the envelope of $^{206}\text{Pb}/^{208}\text{Pb}$ vs. $^{207}\text{Pb}/^{208}\text{Pb}$ for published data for recent Kamchatka volcanic rocks. T6, however, plots firmly within the Pb isotope field of recent Alaskan volcanic rocks, suggesting a source from further afield may be possible for this bed (Fig. 7). A plot of known and dated Alaska tephra beds in comparison to T0–T7 indicates that frequent eruptions from these volcanoes occurred during deposition of most of the Lake El'gygytyn tephra (Fig. 8).

5.2.2 Kamchatka volcanoes

The Kurile–Kamchatka volcanic arc has a total length of 2000 km, is about 400 km wide on the Kamchatka Peninsula, and is one of the most active arc segments on earth (Siebert and Simkin, 2002). It has had numerous explosive eruptions, and the largest number of collapse calderas per unit of arc length (Hughes and Mahood, 2008). In the Holocene, more than 37 large volcanic centers and hundreds of monogenetic vents were active in Kamchatka and about 48 volcanoes are known from the Kurile Islands (Gorshkov, 1970; Siebert and Simkin, 2002; Ponomareva et al., 2007). On the Kurile Islands, the eruption history has been reviewed (Ostapenko et al., 1967; Gorshkov, 1970), although the research mainly focused on recent explosive eruptions (e.g., Nakagawa et al., 2002; Belousov et al., 2003; Hasegawa et al., 2012). In Kamchatka, where many explosive eruptions resulted in widespread dispersal of tephra-fall deposits, the Holocene eruptions and their sequences are well known (Braitseva et al., 1987, 1995, 1997, 1998; Kir'yanov et al., 1990; Bazanova and Pevzner, 2001; Gusev et al., 2003; Ponomareva et al., 2004, 2007; Kyle et al., 2011). Many decades of land mapping, correlation of tephra layers, and determination of ages and magnitudes of eruptions have led to a detailed tephrostratigraphic framework with more than 40 known large explosive eruptions to date. Some Holocene

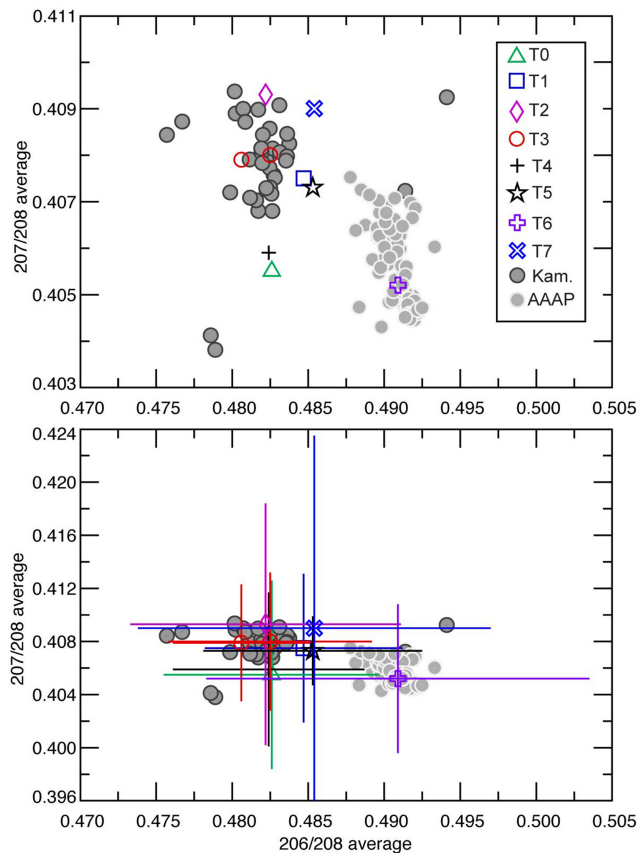


Figure 7. Lead isotope composition of El'gygytyn tephra layers (T0–T7). Top: average $^{206}\text{Pb}/^{208}\text{Pb}$ vs. $^{207}\text{Pb}/^{208}\text{Pb}$ for samples from El'gygytyn core. Bottom: average $^{206}\text{Pb}/^{208}\text{Pb}$ vs. $^{207}\text{Pb}/^{208}\text{Pb}$ for samples from El'gygytyn core including 2σ error bars. Data are plotted with previously published Pb isotope compositions of volcanic rocks from the AAAP (Aleutian Arc–Alaska Peninsula) and Kamchatka (Myers and Marsh, 1987; Kepezshinskas et al., 1997; Bindeman et al., 2010; George et al., 2004; Jicha et al., 2004).

Kamchatka tephra have also been identified off Kamchatka in terrestrial and marine environments on the Bering and Karaginsky Islands (Kyle et al., 2011), in the Sea of Okhotsk cores (e.g., Nürnberg and Tiedemann, 2004; Ponomareva et al., 2004; Derkachev et al., 2012), and as far as mainland Asia (Gorbarenko et al., 2002a, b; Ponomareva et al., 2004).

The volcanic history of the early Holocene and the Pleistocene, however, is less well known in Kamchatka and the Kurile Islands. Older Pleistocene tephra have mostly been removed by glaciation and occur only in isolated outcrops. Geological exposure is poor, erosion rates are high, and the vegetation cover is heavy. Only a few widespread ash layers are known from this time. Among these beds, one of the largest explosive eruptions was sourced from Plosky volcano, Kamchatka, at 10.2 ky cal. BP (e.g., Braitseva et al., 1995; Ponomareva et al., 2013a). The K2 tephra at 27 354–30 325 cal. BP from the northern part of the Kurile Islands

most likely originates from Nemo Caldera (Braitseva et al., 1995; Melekestev et al., 1997; Derkachev et al., 2012). The source of the 177 ka Rauchua tephra is unknown, but it likely comes from the Karymsky volcanic center (Ponomareva et al., 2013b). The Plio-Pleistocene calderas on Kurile Islands and Kamchatka were far larger than the Holocene calderas, and include much thicker proximal pyroclastic deposits (Bindeman et al., 2010), suggesting that the associated tephra layers could be more widely distributed compared to Holocene eruptions. However, the record of these eruptions is poorly known.

The high concentration of tephra layers in marine sediment cores in the adjacent seas supports the notion of a high eruption frequency on Kamchatka in the Plio-Pleistocene. From more than 40 records from the Sea of Okhotsk, spanning the last 350 ky, a sequence of about 22 tephra layers has been documented, including at least 9 widespread ash layers (e.g., Gorbarenko et al., 1998, 2000, 2002a, b; Derkachev et al., 2004, 2012; Nürnberg and Tiedemann, 2004). Western North Pacific deep-sea sediments document numerous explosive eruptions over the last 5 million years with an increase in the number and thickness of volcanic ash layers from 2.6 Ma (Cao et al., 1995: ODP Leg 145, sites 881, 882, 883 and 884; Prueher and Rea, 2001: ODP 887, 883, 882). Cores taken on the SO201 Leg 2 cruise of the RV *Sonne* recovered sediment records spanning the last 180 ky with 14 tephra layers in the Bering Sea and 38 layers in the NW Pacific in Kronotsky Bay and Meiji Seamount (Dullo et al., 2009; Derkachev et al., 2011). High-quality geochemical data on glasses are, however, unavailable for most of the Pleistocene ignimbrites and tephra layers.

Comparison of Pb isotopes, and major- and minor-element compositions of the known Plio-Pleistocene volcanic eruptions to tephra T0–T7 show an obvious affinity to Kamchatkan tephra (potentially excluding T6). However, with the exception of T1, there are no obvious correlations to known beds from Kamchatka. To further explore these connections, we look at the activity of the Kamchatka–Kurile arc based on the ages of major caldera-forming eruptions (from Bindemann et al., 2010) relative to the Lake El'gygytyn beds (Fig. 8). These data suggest that indeed caldera-forming eruptions were taking place at the time of deposition of the Lake El'gygytyn tephra beds.

The lower K_2O wt.% geochemistry of tephra layer T0, with an inferred age of ca 45 ky, shows similarities to the composition of tephra and ignimbrites from southern Kamchatka, such as ca 440 ky old Golygin ignimbrites belonging to Pauzhetka Caldera (Bindeman et al., 2010) and to Kuril Lake ignimbrites (Ponomareva et al., 2004). Although T0 glasses have a slightly more mafic composition, their similarity to tephra deposits related to the Kurile Lake caldera, formed 7.6 ka, is intriguing. The tephra from this eruption was dispersed over an area of ≤ 3 million km^2 , and was found at a distance of 1700 km from the source (Anderson et al., 1998). During this eruption about $\sim 170 \text{ km}^3$ of

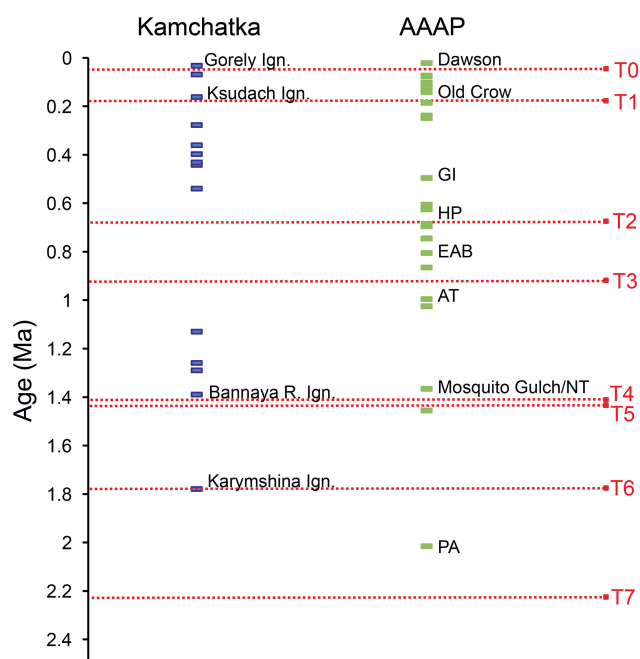


Figure 8. Lake El'gygytyn tephra ages compared to ages of dated caldera eruptions in Kamchatka and ages of published distal tephra from Aleutian Arc–Alaska Peninsula (AAAP; after Preece et al., 1999; 2011a, b; Westgate et al., 2001, 2011b; Demuro et al., 2008; Bindeman et al., 2010). It is important to note that the frequency of eruptions in Alaska is much greater, but we limited the tephra included in this plot to ones that have been directly dated.

tephra were produced, and the eruption ranks among the largest Holocene explosive eruptions on Earth (Ponomareva et al., 2004). The chemical similarity of the T0 glasses to this source suggests that this may represent an unknown older eruption from this volcano.

Tephra layer T1 in core 5011 has been correlated to the second layer in the pilot core Lz1024. The sample from the pilot core is chemically indistinguishable from Rauchua tephra from a terrestrial site in the Far East Russian Arctic, 330 km NW of Lake El'gygytyn (Ponomareva et al., 2013b), and correlates to tephra layers identified in the Bering Sea and Pacific sediment cores SO 201-2-81/85 (Dullo et al., 2009). The distribution of the tephra thicknesses suggests that these beds likely originate from an eruption in the Karymsky volcanic center in the Kamchatka volcanic arc, although the tephra has not been identified on land (Ponomareva et al., 2013b).

Tephra T2, T3, T4 and T6 have dacitic to rhyolitic (with moderate K_2O) compositions, and similar trace-element compositions, which only vary slightly in their ratios of highly incompatible elements (e.g., Zr/Nb , Nb/Y); however, these beds clearly separate by differences in their Ba/Y ratios. T6 has a Pb isotope composition that differs from T2–T4 and may suggest T6 has an Alaskan source. Overall, the compositions are typical for Kamchatka and partic-

ularly for centers in the volcanic front around the Karymsky Caldera (Kyle et al., 2011; Ponomareva et al., 2013a). This area has a high density of nested calderas that might be plausible sources for the tephra. Few of the ignimbrite flows related to these calderas have been dated so far, and no exact match in age and composition between published data and these tephra exists at present. One possible candidate could be the dacitic eruption related to Stena-Soboliny Caldera at 1.13 Ma (Bindeman et al., 2010), which could be correlated to the dacitic T3 tephra with an inferred age of ca 0.9 My old, although the age difference of 200 ky makes this correlation questionable. T6 has an identical inferred age of ca 1.78 My old with ignimbrites related to Karymshina Caldera (Bindeman et al., 2010). However, the composition of the ignimbrites produced by this super eruption shows a much more evolved geochemical composition than the glass of T6 (Bindeman et al., 2010).

Dacite-andesitic glasses of tephra T5 imply an andesitic composition for the bulk tephra. Only a few large andesitic eruptions are known in Kamchatka, and all are dated to the Pliocene (3.5–5.7 Ma; Bindeman et al., 2010). These eruptions occurred in the eastern volcanic zone of Kamchatka and in Sredinny Range. None of these eruptions can be directly correlated to T5 tephra, which is dated to 1.4 Ma.

The oldest tephra, T7, has a unique composition and a somewhat uncertain age due to possible sediment disturbance in the cores. This makes its correlation to a potential source problematic. Nevertheless, judging from its geochemistry alone, and assuming a source in Kamchatka, it should be located in a small region of the volcanic arc where compositions of tephra are more alkalic, enriched in LILE and depleted in HREE, as exemplified by Opala Volcano in southern Kamchatka (Kyle et al., 2011).

6 Conclusions

The eight tephra layers preserved in Lake El'gygytyn sediments provide a unique opportunity to correlate terrestrial and marine climate archives in the Bering Sea and North Pacific region. They also indicate that there is great potential to find distal ash in western Beringian sediments, which would allow a dated framework for these archives.

Geochemical data indicate that the source of these tephra beds lies in the subduction zone volcanoes that form the Kurile–Kamchatka–Aleutian Arc and Alaska Peninsula. While major-, minor- and trace-element geochemistry is generally inconclusive in determining the potential sources for the beds, the Pb isotopic data show that the Kurile–Kamchatka arc is the most likely source for seven of the eight tephra. For tephra T6, an Alaskan source is possible, but there is presently no known tephra that correlates to this layer.

The geochemical composition of the seven other tephra suggests a Kamchatka source, and indeed one, T1, has been correlated to the Rauchua tephra, which likely has a

Kamchatkan source (Ponomareva et al., 2013b). Even though there is evidence for numerous highly explosive eruptions, as seen in nested calderas in the eastern volcanic zone in Kamchatka, the sequence and age of those explosive events are not well constrained. Only a small number of Pleistocene ignimbrites in Kamchatka have been studied with modern analytical tools, and the distal record is limited. Therefore, due to a restricted database available for geochemically fingerprinted Kamchatkan material, none of the remaining Lake El'gygytyn tephra were successfully correlated, although potential source volcanoes and caldera complexes can be identified. Additionally, tephra layers described from RV *Sonne* 201, 202, ODP or ICDP cores from the Bering Sea and North Pacific lack sufficient glass geochemistry data needed to compare with Lake El'gygytyn tephra.

Even though the source volcanoes for tephras T0–T7 are not known at this time, they still offer a unique potential for future correlation of terrestrial and marine sites. The thickness of the Lake El'gygytyn tephra, over 1500 km from potential sources, suggest they are widely distributed, and thus, are likely present in numerous depositional settings. The geochemistry presented here will allow for the future identification of these tephras, and their linking to other terrestrial and marine deposits and eventually to their source volcanoes. Ultimately these tephra beds from Lake El'gygytyn provide the basis for a robust tephrostratigraphic framework for western Beringia that we predict will develop quickly now that the presence of tephra beds in this region is known.

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Appendix A: Appendix methods

A1 Petrographic description

All visible tephra layers in the cores were sampled and described petrographically, prior to chemical analysis for major, trace and Pb isotope composition by EPMA and LA-ICP-MS.

Tephra layers were identified in the cores directly after opening. Smear slides were taken from every change in lithological composition and checked for volcanic glass shards. All tephra layers were sampled and freeze-dried. Petrologic characteristics of the tephra layers were described from smear slides and polished sections (grain size, mineralogical content). Subsamples for chemical investigations of glass shards were carefully washed in distilled water to sieve off finest grains ($< 6\mu\text{m}$) and dried at 40°C , mounted in epoxy, polished and carbon-coated for EPMA. Backscatter images of glass shards were done at the electron microprobe in Kiel.

A2 Electron probe microanalyses

Glass shards from the tephra layers were analyzed for their major and minor element composition by electron microprobe to gain information about the provenance and enable a robust correlation to existing databases of possible source volcanoes. Analyses were performed using two different electron microprobes, one at the University of Alberta, Edmonton, and a second at GEOMAR, Kiel. These two laboratories host databases for volcanic ash layers from different regions (North America and Kamchatka, respectively), which are relevant to the correlation of the El'gygytyn tephra layers. The INTAV (INQUA international focus group on tephrochronology) intercomparison of tephrochronology laboratories confirmed the high quality of the microprobe results from both instruments (Kuehn et al., 2011), with both instruments producing data sets that are identical within statistical deviations. In this way interlaboratory calibration issues could be excluded.

A2.1 EPMA – University of Alberta, Canada

At the University of Alberta, the major-element geochemical composition of single glass shards was analyzed with a JEOL 8900 superprobe. Analytical conditions were a 15 kV accelerating voltage, 6 nA beam current, and $10\mu\text{m}$ beam diameter, with a 20 s peak and 10 s background counting time on all elements. Na, Si and Al were analyzed first to minimize the effects of Na-loss. Standards used for calibration are Lipari rhyolitic obsidian (UA 5831: Si, Al, Na, K) and minerals (tugtupite: Cl; diopside: Ca; willemite: Mn; pyrope: Fe, Mg; ilmenite: Ti). Matrix effects are corrected for using the ZAF correction factor. Internal glass standards ID 3506 (a Lipari obsidian) and Old Crow tephra were analyzed at the beginning, after every four samples (i.e., ~ 100 shards),

and at the end of each analytical session to monitor quality of the calibration (e.g., Kuehn et al., 2011). Routine internal standard analyses also allow for the comparison of data collected over the course of days to years by providing a means to correct for any small offsets that may be present due to variations in probe conditions over time. A minimum of 25 shards per sample were analyzed to capture the compositional range, and all totals under 90 % were discarded. Data are normalized to 100 % on a volatile-free basis and are presented as weight percent (wt.%) in Supplement Table S1. All El'gygytyn tephra layers have been compared to the University of Alberta Alaskan tephra database, with similarity coefficients being calculated for the most similar known Alaskan tephra, and these are presented in Supplement Table S2.

A2.2 EPMA – GEOMAR, Kiel, Germany

At GEOMAR the chemical composition of single glass shards – major elements, S, Cl and F – was analyzed with a JEOL JXA 8200 electron microprobe equipped with five wavelength dispersive spectrometers including three high-sensitivity ones (2 PETH and TAPH). Analytical conditions were 15 kV accelerating voltage; 6 nA beam current and 10/10 s (peak/background) counting time for Na; 20/10 s for Si, Al, Fe, Mg and Ca; 30/15 s for K, Ti, Cl and S; and 40/20 s for Mn and F. Analyses were performed with a $5\mu\text{m}$ electron beam. Standards used for calibration were natural reference material from the Smithsonian collection (rhyolitic glass USNM72854 VG-568 for Si, Al, Na, and K; basaltic glass USNM 113498/1 VG-A99 for P, Fe, Mg, Ti and Ca; and scapolite USNM R6600-1 for S and Cl) (Jarosewich et al., 1980). Sodium was analyzed first during all sessions; sodium loss was not observed. The reference materials and internal standard glasses (KN18, Lipari obsidian) were measured in triplicate at the beginning, after every 60 analyses and at the end of each analytical session. Initial data reduction was done using CITZAF correction (Armstrong, 1995) integrated in JEOL software. Correction for instrumental drift (if any) was done based on deviation of the data for reference samples from their recommended values. The corrections for instrumental drift applied were minor ($< 5\%$ relative to all elements) but allowed the best possible accuracy and reproducibility over long time spans. Only totals above 93 % were accepted. Data were screened to avoid analyses contaminated by micro crystals. To cover the whole range of geochemical variations, at least 20 glass shards in every sample were analyzed. All analytical data presented in Supplement Table S3 are given in wt.% oxides and are normalized to 100 % totals.

A3 Laser ablation ICP-MS analyses

Laser ablation (LA) ICP-MS analyses were performed on a range of individual glass shards for their trace element and Pb isotope composition in the Department of Geography and Earth Sciences, Aberystwyth University. All analyses were performed using a Coherent GeoLas ArF 193 nm excimer laser ablation system coupled to a Thermo Finnegan Element 2 sector field ICP-MS. The Aberystwyth laboratory provides established analytical methods for determining the chemical composition of fine-grained pumiceous glass shards with a median size of less than 40 μm , and allows for the analysis of a pure glass phase at a spatial resolution of 10–20 μm . In total, 226 single shards were analyzed for their glass trace element concentrations and 190 for their Pb isotopes. The database for rhyolitic tephtras from Alaska/Yukon was largely developed with analyses on this instrument (see for example Preece et al., 2011a).

A3.1 Trace element analysis

Trace element data were collected for individual shards using 20 and 14 μm diameter ablation craters at a laser energy of 10 J cm⁻² and a repetition rate of 5 Hz over a 24 s acquisition. Crater size was varied because of the size of individual glass shards to be analyzed. The minor ²⁹Si isotope was used as the internal standard, and the average concentration of SiO₂ for each sample (determined by EPMA) was used to calibrate each analysis, after normalization to an anhydrous basis. The NIST 612 reference glass was used for calibration, with concentrations given in Pearce et al. (1997), and a fractionation factor was applied to the data to account for analytical bias introduced related to the different matrices of the reference standard and the sample (see Pearce et al., 2011). Further details of the setup procedures and discussion of the precisions and accuracies, etc., can be found in Pearce et al. (1996, 1999, 2002, 2004a, 2007, 2011), and ICP-MS and laser operating conditions are summarized in Supplement Table S4. The MPI-DING reference glass ATHO-G (Jochum et al., 2006; Jochum and Stoll, 2008) was analyzed as an unknown under the same operating conditions at the same time, and these analyses are given in Supplement Table S5. Analytical precision is typically within ± 5 –10 % (depending on concentration and crater diameter), and accuracy is typically around ± 5 %, when compared with the published GeoReM concentrations for ATHO-G. All individual shard trace element data are presented in Supplement Table S6.

A3.2 Lead isotopes

Pb isotope compositions were determined by LA-ICP-MS from individual glass shards for all seven samples at Aberystwyth University. For the Pb isotope analysis the Thermo Finnegan Element 2 sector field ICP-MS was operated in low resolution mode, and He was used as the sample gas

(0.825 L min⁻¹) made up with Ar (~ 0.7 L min⁻¹). The instrument was optimized to give the greatest sensitivity for ²⁰⁸Pb from the NIST 612 reference glass. Instrument operating conditions are given in Supplement Table S4. Where possible, analyses were performed using a 30 μm diameter laser beam which ablated to a depth of about 20 μm , although for some samples a 20 μm diameter beam had to be used to avoid ablation of mounting resin.

Calibration was achieved from 10 repeat analyses of NIST 612 at the start and end of each set of analyses using the Pb isotopic ratios from GeoReM (see Jochum et al., 2005; Jochum and Stoll, 2008). Gas blanks (5 repeats) were measured before and after each of the standard analyses, and also between samples. Twenty individual glass shards were analyzed in each unknown, and only two unknowns were analyzed between the NIST reference materials in each set of analyses. This process ensured that reference materials (for calibration and to monitor any instrumental drift) were analyzed at intervals of typically around 1.5 h. Spectra obtained from the NIST 612 glass (30 μm ablation craters) gave about 270 000 cps for ²⁰⁸Pb (38.57 ppm Pb), and from the unknown glass shards (with an average of ~ 12 ppm Pb) count rates were lower, averaging ~ 29 000 cps, a result of poorer ablation characteristics of natural glasses and the need to use a 20 μm diameter crater for many analyses. For each analysis the masses 201, 204, 206, 207, and 208 were measured sequentially and averaged over 100 scans. ²⁰⁴Pb suffers interference from ²⁰⁴Hg, and thus ²⁰¹Hg was measured to allow a correction for this. Mass bias in the ICP-MS was calculated from the ²⁰⁶Pb/²⁰⁸Pb isotope ratio of the NIST glass after blank subtraction (< 100 cps on 208Pb), and using this, and the relative isotope abundances of Hg (Rosman and Taylor, 1998), the calculated contribution of ²⁰⁴Hg (based on the ²⁰¹Hg intensity) was subtracted from the intensity of the mass 204 peak to leave only ²⁰⁴Pb. Mercury is present as a trace contaminant in the Ar used in ICP-MS, as well as being present in the NIST standard and in natural volcanic glasses; thus any attempt to measure ²⁰⁴Pb in glasses must compensate for this Hg interference to the ²⁰⁴Pb signal. Once the correction for ²⁰⁴Hg had been applied, calibration was achieved using the Pb isotope ratios given by Jochum et al. (2005), and included correction for any instrument drift during each run. Further considerations of this technique are given in Westgate et al. (2011a). Supplement Table S7 shows the determined Pb isotope ratios for two USGS glass reference materials, TB-1G and BCR-2G, with the Pb isotope composition taken from GeoReM (see Jochum and Stoll, 2008). The Pb isotope ratios determined by LA-ICP-MS analyses are typically within ± 0.3 % of the reference isotope ratios, and analytical precision for ratios between ²⁰⁶Pb, ²⁰⁷Pb and ²⁰⁸Pb is generally better than ± 1 %. Ratios to ²⁰⁴Pb however show worse precision, at about ± 1.5 % for TB-1G and about ± 4.5 % for BCR-2G, despite good accuracy, and this reflects the noise on the Hg signal, which makes a significant contribution to the ²⁰⁴Pb signal, particularly at low concentrations

and count rates. In the case of BCR-2G, the ^{204}Hg correction accounts for almost 70 % of the signal at mass 204, leaving only a few hundred ^{204}Pb counts. Considering these issues, the size of the ablation craters, and the speed with which analyses can be performed, these data are useful and appropriate for reconnaissance studies of Pb isotope ratios in tephra deposits (see Westgate et al., 2011a).

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