Cosmogenic ³He dating of olivine with tightly retained mantle ³He, Volcano Mountain, Yukon

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Abstract. We present a step-heat method for isolating cosmogenic 3 He (3 Hec) from mantle He in olivine xenocrysts to date the eruption of four of the morphologically youngest nephelinite lava flows from Volcano Mountain (VM), the youngest cone in the Fort Selkirk volcanic field in Yukon, Canada. In these olivines, the standard procedure of powdering grains to $<30 \mu m$ failed to effectively remove mantle helium prior to fusion: samples from four different flows yielded unusually high powder fusion 4 He concentrations of 40 to 141 ncc/g with 3 He/ 4 He ratios of 7.9 to 9.6 R_A. When combined with the 3 He/ 4 He ratios obtained by crushing (average 8.1 ± 0.2 R_A), these measurements yield Holocene cosmogenic exposure ages but with very large uncertainties arising from the large mantle 3 He correction. The inability to effectively isolate 3 He $_{c}$ from these samples likely arises from the survival of small ($<<30 \mu m$) fluid inclusions hosting mantle He through the powdering step. The presence of such unusually small fluid inclusions may relate to the origin of the olivines as disaggregated peridotite xenoliths rather than the more commonly analyzed olivine phenocrysts. We circumvented this problem by step-heating powdered olivine in steps of 800, 1000, and 1400°C. Helium isotopic systematics indicate that 70-90 % of 3 He $_{c}$ was released in the low temperature step and the rest was released in the middle temperature step. By the highest temperature step, the released He had a mantle-like 3 He/ 4 He ratio. Combining results from the step heating and crush-fusion methods, we determined that the four Volcano Mountain lava flows erupted approximately coevally, at 10.5 ± 1.7 ka.

1 Introduction

The in-situ production of cosmogenic ³He in olivine has been used to date the surface exposures of lava flows for decades (e.g. Aciego et al., 2007; Kurz et al., 1990; Marchetti et al., 2020). The abundance of mm-size olivine crystals in basalt and the high helium retentivity of olivine (Shuster and Farley, 2004; Trull and Kurz, 1993) make it an ideal candidate for cosmogenic ³He dating. The method is particularly suited to the determination of eruption ages of young volcanics (Heineke et al., 2016; Fenton and Niedermann, 2014; Licciardi et al., 1999) because ³He has one of the highest production/detection limit ratios amongst cosmogenic nuclides (Niedermann, 2002).

Cosmogenic ³He in terrestrial rocks is primarily produced by spallation of target nuclei in the crystal lattice of minerals. At surface temperatures, ³He_c accumulates quantitatively in the matrix of olivine (Kurz, 1986a, b), allowing exposure ages to be obtained from independently established production rates. Unlike cosmogenic radionuclides such as ¹⁰Be and ²⁶Al in which the background nuclide concentration is often negligible, a practical challenge of the ³He method is isolating the cosmogenic component from other ³He components. The most common non-cosmogenic sources of ³He include thermal neutron capture on ⁶Li (Andrews and Kay, 1982; nucleogenic ³He, ³He_{nuc}) and mantle He with ³He/⁴He ratios well above crustal values trapped in fluid inclusions (Kurz, 1986a, b). U and Th-bearing minerals also produce radiogenic ⁴He (⁴He_{rad}) through α decay. Taken together:

$$^{3}\text{He}_{\text{total}} = ^{3}\text{He}_{\text{c}} + ^{3}\text{He}_{\text{m}} + ^{3}\text{He}_{\text{nuc}}$$
 (1)

$$40 ^4He_{total} = ^4He_m + ^4He_{rad} (2)$$

However, concentrations of Li, U, and Th in olivine tend to be low (Woodhead, 1996; Dostal and Capedri, 1975; Kent and Rossman, 2002; Blard and Farley, 2008; Amidon et al., 2009; Blard et al., 2013) such that neutron capture and radioactive decay produces minimal amounts of He in most young lava flows. Thus, in the specific limiting case in which nucleogenic/radiogenic He can be ignored:

$$45 {}^{3}\text{He}_{\text{total}} = {}^{3}\text{He}_{c} + {}^{3}\text{He}_{m} (3)$$

$$^{4}He_{total} = ^{4}He_{m} \tag{4}$$

Equation 3 highlights that cosmogenic ³He dating in young olivine requires the isolation of ³He_c from ³He_m. Past studies (e.g., Kurz, 1986a) achieved sufficient separation of the two components using a two-step process in which the first step is crushing whole olivine grains in vacuum and measuring the ³He/⁴He ratio. Vacuum-crushing olivine grains to <500 μm releases most mantle He in fluid inclusions and yields the crush ratio for a given sample, (³He/⁴He)_{crush} (Kurz, 1986a, 1986b; Puchol et al., 2017; Blard, 2021). In the second step, the crushate is powdered to reduce the number of surviving fluid inclusions, and then fused in vacuum to measure mostly matrix-hosted He. In basalts, crush ratios are often close to the upper mantle ratio of ~8 times the atmospheric ratio (R_A = 1.384 x 10⁻⁶) while fusion ratios ((³He/⁴He)_{fusion}) of cosmic-ray irradiated samples may be several times higher (Kurz, 1986a, b). If ⁴He_{fusion} is assumed to be entirely mantle derived (eq. 4), then the cosmogenic component can be isolated even if some of the He in the fusion is mantle derived:

$${}^{3}\text{He}_{c} = {}^{3}\text{He}_{\text{fusion}} - ({}^{4}\text{He}_{\text{fusion}})({}^{3}\text{He}/{}^{4}\text{He})_{\text{crush}}$$
 (5)

³He_c is then used to calculate an exposure age. The uncertainty on ³He_c depends in part on how large and how well-known the correction for mantle He is. When powdering does not effectively remove the mantle component, the correction can be large compared to the total ³He measurement.

In this study, we attempted the crush-fusion method to obtain exposure ages for the youngest lava flows from Volcano Mountain in Yukon, Canada, however the mantle component was so large and well-retained during powdering that precise ³He_c concentrations could not be obtained. As an alternative we applied a protocol previously used on cosmic-ray exposed peridotites (Swindle et al., 2023) in which step heating successfully isolated cosmogenic ³He from mantle ³He.

65 2 Geologic Setting

The Fort Selkirk Volcanic Complex in Yukon, Canada is comprised of valley-filling volcanics that interacted with Cordilleran ice sheets during the Pliocene and Pleistocene epochs (Jackson et al., 2012; Jackson and Huscroft, 2023). Radiometric ages on the complex range from 0.4 to 4.3 Ma (Jackson and Huscroft, 2023). Volcano Mountain, also called Nelruna in the local First Nations language, is located north of the confluence of the Yukon and Pelly Rivers and is the youngest volcanic center in the complex (Jackson and Stevens, 1992). It rises several hundred meters above the valley fill and is comprised of lava flows, some of very youthful appearance. Among the more recent flows are several that dammed small lakes in the early-mid Holocene, some time before 7300-4200 BP (Jackson and Stevens, 1992; Francis & Ludden, 1990). The youngest flows could be even younger (Jackson and Stevens, 1992; Francis & Ludden, 1990). Jackson and Huscroft (2023) provide detailed field descriptions, photographs, and an eruptive history of Volcano Mountain.

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Volcano Mountain lava flows are nephelinites carrying ultramafic xenoliths (mostly dunite) and olivine phenocrysts and xenocrysts (Francis & Ludden, 1990; Trupia & Nicholls, 1996). The xenocrysts are large (up to 1 cm) and are the dominant population of olivine crystals in the rock. Based on deformation features such as kink-banding and subgrain boundaries, their anhedral/broken shape, and highly magnesian composition (Fo88-90), they are likely disaggregated dunite xenoliths (Francis & Ludden, 1990; Trupia & Nicholls, 1996). These abundant and large xenocrysts are an attractive target for cosmogenic ³He dating.

3. Samples

3.1 Field Sampling

Sampling of Volcano Mountain lavas was intended to date flows of the most youthful appearance and stratigraphic position to determine the age of the most recent eruption (Table 1; Fig.1). Particular attention was paid to the flow within the small summit crater as the most likely to be the youngest. For redundancy purposes, 2-3 samples were collected at each location. Site parameters used to guide sampling included: a relatively flat, stable surface exhibiting primary depositional features (e.g. ropy textures) and if possible, minimal vegetation cover. A gas-powered cutoff saw was used to control sampling by slicing the surface to a uniform depth. A rock hammer and chisel were used to liberate the sample from the prepared area. Approximately 2 kg of sample was collected from each site.

A total of 13 samples were collected on six separate lava flow surfaces. Jackson and Stevens (1992) (see also Jackson and Huscroft, 2023) mapped VM flows and named them based on the direction of the flow (north or south) and its stratigraphic position relative to the other flows based on field observations (0-2 where 2 is assigned to the youngest flows) (Fig. 1). Samples VM-01, 02, and 03 were collected from the stratigraphically highest flow in the cinder cone (flow 2Na). VM-06 is from the fifth south flow (flow 2Sa), VM-08 and 09 are samples from the third south flow (flow 1Sa) and VM-10 and VM-11 are samples from the second north flow (flow 1N). Samples VM-04, 05, 12, and 13 are from the fourth south flow (flow 2Sb, possibly the same age as flow 2Sa, (Jackson and Huscroft, 2023)), but along with sample VM-07 (flow 2Sa), yielded too little olivine to make a cosmogenic ³He measurement. See Table 1 for sample details.

At elevations ranging from ~0.7-1.1 km, the area experiences on average 28.8 cm of snow cover for 5.5 months of the year 100 (Government of Canada, 2007). Vegetation, including lichen and moss, covered a significant surface area of the samples, though it is highly variable, from no coverage at VM-03 to covered in 8 cm thick mat of moss at VM-08.

Table 1

		Latitude	Longitude	Elevation	Depth	Shielding
Flow	Sample	(N)	(W)	(m)	(cm)	Factor
2Na	VM-01	62.92311	137.3775	1052	3	0.96
2Na	VM-02	62.92302	137.3779	1067	3	0.96
2Na	VM-03	62.92302	137.3779	1067	3.5	0.96
2Sa	VM-06	62.90544	137.43741	718	3	0.97
1Sa	VM-08	62.90447	137.4359	725	2.5	0.96
1Sa	VM-09	62.90413	137.43684	725	2	0.96
1N	VM-10	62.95478	137.36848	788	3.5	0.98
1N	VM-11	62.9547	137.36888	785	3.5	0.97

3.2 Sample Selection and Preparation

All hand samples were scrubbed of vegetation, rinsed in water, dried in air, and jaw-crushed to mm-sized grains. The abundance of coarse olivine crystals varied among samples; only samples VM-01, 02, 03, 06 and 08 through 11 had sufficient olivine for cosmogenic ³He measurement. From these, matrix-free, mm-size olivine crystals were handpicked from the crushate based on their distinct lack of cleavage and green color.

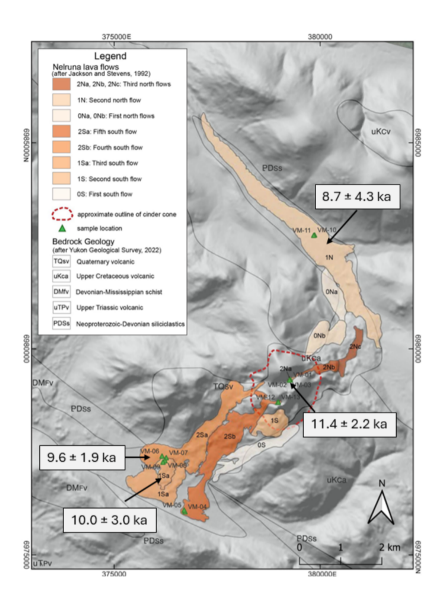


Figure 1. Geologic Map of Volcano Mountain (Nelruna). Map of Volcano Mountain study area after Jackson and Stevens (1992). Samples VM 01,02, and 03 are from the stratigraphically youngest flow on Volcano Mountain, while the other samples all come from flows of very youthful appearance. Numbers in boxes indicate our best estimate of eruption age based on cosmogenic ³He.

4 Analytical Methods

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4.1 Crushing and Powder Fusion Method

Helium was extracted from olivines by crushing under vacuum for 2 min, and then re-crushed for another 5 min. The two steps were combined into a single He result. Previous work with this same crusher design indicates these crush durations will not cause release of matrix-sited ³He (Blard et al., 2008). Prior to dropping samples into the crusher anvil, 2 minute "no sample" crushes were measured and used to blank correct He measurements. Maximum blank crush levels of ³He and ⁴He were 0.07 fcc STP and 0.2 ncc STP, respectively (1 ncc = 26884 Mat; STP are the standard temperature and pressure conditions of 273.15 K and 101.3 kPa where 1 mol of an ideal gas occupies a volume of 22419 cm³).

- 125 For some samples, material recovered from the vacuum crushes was powdered in a mortar and pestle under ethanol and sieved to <30 μm, dried, weighed, and wrapped in Sn foil packets for fusion analysis. For other samples analyzed by fusion, the hand-picked olivines did not undergo the initial vacuum-crush step, only powdering under ethanol. Ethanol was used to circumvent (Cox et al., 2022) trapping of atmospheric He during powdering (Protin et al., 2016).
- Sn-foil wrapped samples for high temperature vacuum extraction analysis were loaded into the dropper arm of a double-wall furnace and evacuated overnight. The dropper arm was not heated to eliminate any possibility of baking-induced loss of ³He. The furnace was outgassed directly into the vacuum pump for 1 hr at 1400°C prior to analysis. Samples were then dropped into a carbon liner in the furnace and heated to ~1200 °C for 25 min. Re-extracts were measured directly after each extraction to ensure the complete release of helium. When the re-extract signal exceeded the pre-analysis hot blank, the re-extract was repeated. All sample gas in excess of blank was combined in the concentrations reported here. While these olivines were not fused during these analyses (the temperature was held sub-melting to minimize He blanks), complete He extraction is ensured by the re-extraction steps. For consistency with prior work, we nevertheless refer to these measurements as "powder fusions". Two olivine samples were fused in a lithium borate flux at ~1000 °C for 25 min in a molybdenum liner. The flux was used in these fusions to reduce blank He levels by lowering extraction temperature (Farley et al., 2020). Hot blanks at ~1200 °C for ³He and ⁴He were 0.8 fcc STP and 0.2 ncc STP respectively. Blanks measured on the tin foil are negligible compared to this hot furnace blank. Blank corrections made to sample gas concentrations were mostly <2% but always <5%.

4.2 Step-heating

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About 750 mg of powdered (<30 μm) sample was wrapped in Sn foil packets and loaded into the vacuum furnace as described above. The Sn packets were then dropped into the furnace, and He successively released at 800 °C, 1000 °C, and 1400 °C each for 30 minutes duration. These temperature steps were selected based on previous studies reporting the release of cosmogenic

³He below 1000 °C (Kurz, 1986a; Swindle et al., 2023). A thermocouple and pyrometer were used to calibrate furnace temperatures as a function of furnace output power, after which heating steps were maintained using constant power output.

Hot blanks were performed on an empty furnace at each of the scheduled temperature steps to use as blank corrections on He quantities at each step. Blank levels were <1.5 % of extracted He for all steps. Re-extracts were measured on the material after step-heating to ensure total helium release. When the re-extract signal exceeded the pre-analysis hot blank, the re-extract was repeated. When the re-extract exceeded the hot blank, the excess He was added to the high temperature step.

4.3 Helium Measurement

Helium was purified and measured as previously described (Horton et al., 2019; Swindle et al., 2023). Gas extracted from the furnace was passed through a charcoal U-trap to remove contaminants such as CO₂, H₂O, and Ar. Remaining reactive gases were removed using a hot SAES NP10 getter and H₂ was removed using a cold SAES NP10 getter. Helium was then cryofocused on charcoal at 14 K and released at 34 K into a Helix SFT mass spectrometer. ³He was measured on a pulse-counting electron multiplier and ⁴He on a Faraday cup. Instrument sensitivity and internal consistency were monitored by running a standard containing ~70 ncc of He (³He/⁴He ratio = 2.05 R_A) throughout our experiments. This standard was created from pure ³He and ⁴He using a capacitance manometer and the amounts delivered are known to better than ±1 %.

4.4 Shielding Correction

Corrections for snow cover, vegetation cover, and self-shielding were considered for every sample. The snow correction factor was calculated using Eq. 3 in Vermeersch (2007) based on the spallogenic neutron attenuation length, average snow thickness, and snow density. A snow density of 0.3 g/cm³ was assumed. Topographic shielding corrections were made for VM-01, 02, and 03 due to their location inside the VM cinder cone (see red outline in Fig. 1). Topographic shielding for the other samples was negligible. Shielding factors are listed in Table 1.

5 Results

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5.1 Crush and Fusion Results

Analytical results for the crushing and fusion experiments are listed in Table 2. The three samples (VM-02, VM-06, and VM-170 09) analyzed by crushing come from three different flows (2Na, 2Sa, and 1Sa) yet have indistinguishable crush ³He/⁴He ratios (7.99 ± 0.19, 8.08 ± 0.19, and 8.23 ± 0.2 R_A). Crush-release He concentrations in these samples are high for basalt-hosted olivines, ranging from 76 to 184 ncc/g.

Six powder fusion analyses and the total helium yield from the three step heats (described below) provide information on the matrix-hosted helium in all four of the investigated lava flows. Across these nine analyses, He concentrations range from 39 to 141 ncc STP/g, similar to the crush results. 3 He/ 4 He ratios ranged from 7.9 ± 0.2 to 9.6 ± 0.2 R_A. For the three lava flows on which both crushing and fusion analyses are available, fusion 3 He/ 4 He ratios ranged from the same within error as the crush result (VM-09 from flow 1Sa), to a maximum of 1.5 R_A higher (VM-06 from flow 2Sa).

Table 2

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		Analysis	Mass	³ He		⁴ He		³ He/ ⁴ He		³ He _c		3 He $_{c}$
Flow	Sample	Type	g	pcc/g	±	ncc/g	±	RA	±	Mat/g	±	%
2Na	VM-02	Crush	0.097	2.04	0.04	183.7	3.7	7.99	0.19			
2Na	VM-02	Fusion	0.246	1.31	0.03	109.0	2.2	8.67	0.25	2.75	1.25	7.8
2Na	VM-01	StepHt	0.734	1.13	0.04	91.0	3.2	8.96	0.26	3.27	1.57	10.8
2Na	VM-03	StepHt	0.800	1.32	0.05	104.9	3.7	9.03	0.25	4.05	1.82	11.4
2Sa	VM-06	Crush	0.630	0.85	0.02	75.6	0.9	8.08	0.19			
2Sa	VM-06	Fusion	0.500	0.51	0.01	38.6	0.8	9.59	0.27	2.19	0.36	15.8
1Sa	VM-09	Crush	0.100	1.22	0.02	106.5	1.4	8.23	0.20			
1Sa	VM-09	Fusion	0.068	0.51	0.04	46.1	1.8	7.94	0.71	-0.51	1.16	
1Sa	VM-08	Fusion	0.320	0.70	0.01	53.4	1.1	9.37	0.27	2.27	0.64	12.1
	Assumes average of all VM crushes:						8.10	0.20				
1N	VM-10	Fusion	0.116	0.55	0.04	47.4	1.0	8.36	0.33	0.46	1.17	3.1
1N	VM-11	Fusion	0.100	1.15	0.02	100.3	2.0	8.25	0.24	0.56	1.15	1.8
1N	VM-11	StepHt	0.657	1.65	0.06	140.6	4.9	8.44	0.41	1.79	2.37	4.0

We used these data to compute cosmogenic 3 He concentrations via equation 1 by assuming that the crush 3 He/ 4 He ratio on a given flow can be applied to all samples obtained from that flow. For flow 1N, for which we did not measure a sample by crushing, we computed cosmogenic 3 He concentrations by assuming the mean crush 3 He/ 4 He ratio of 8.1 ± 0.2 R_A obtained on the other flows. This assumption is supported by the step heat results described below.

Cosmogenic 3 He was confidently detected in five of the nine fusion analyses (Table 3). For flow 2Na we obtained cosmogenic 3 He concentrations of 2.75 ± 1.25 Mat/g (VM-02), 3.27 ± 1.57 Mat/g (VM-01), and 4.05 ± 1.82 Mat/g (VM-03). One analysis each of flow 2Sa (VM-06) and 1Sa (VM-08) yielded 3 He_c of 2.19 ± 0.36 and 2.27 ± 0.64 Mat/g, respectively. For the remaining

four analyses the 1 sigma analytical uncertainty on ³He_c exceeds the measured value, i.e., the cosmogenic ³He concentration is indistinguishable from zero.

The high analytical uncertainties on ${}^{3}\text{He}_{c}$ - typically more than 1 Mat/g - in the VM samples arises from the large corrections required for mantle ${}^{3}\text{He}$ present in the powder fusion analyses (equation 1). For example, as shown in Table 2, the fraction of ${}^{3}\text{He}$ that is cosmogenic in these analyses ranges from a maximum of 15.8% (VM-06) to < 2% (VM-11, VM-09). Stated differently, the ${}^{3}\text{He}/{}^{4}\text{He}$ ratios obtained by fusion are very close to the ratios obtained by crushing, and this similarity at least in part reflects preservation of a high concentration of mantle helium through the powdering step.

Table 3

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		Mass	Step	³ He		⁴ He		3 He/ 4 He		$^{3}\mathrm{He_{c}}$		$^{3}\mathrm{He_{c}}$
Flow	Sample	g	°C	pcc/g	±	ncc/g	±	RA	±	Mat/g	±	%
2Na	VM-01	0.734	800	0.34	0.01	19.8	0.4	12.28	0.35	3.26	0.27	35.9
			1000	0.29	0.01	25.2	0.5	8.35	0.24	0.45	0.30	5.8
			1400	0.50	0.01	46.0	0.9	7.87	0.22			
			Total	1.13	0.04	91.0	3.2	8.96	0.44			
			Total	Cosmog	enic					3.71	0.41	
2Na	VM-03	0.800	800	0.27	0.01	14.8	0.3	13.14	0.37	2.76	0.21	37.9
			1000	0.25	0.01	20.1	0.4	9.00	0.26	0.62	0.25	9.3
			1400	0.79	0.02	70.0	1.4	8.17	0.23			
			Total	1.32	0.00	104.9	0.3	9.03	0.44			
			Total Cosmogenic					3.38	0.33			
1N	VM-10	0.657	800	0.39	0.01	27.6	0.6	10.18	0.29	2.47	0.35	23.5
			1000	0.51	0.01	43.9	0.9	8.38	0.24	0.97	0.52	7.0
			1400	0.75	0.01	69.0	1.4	7.79	0.22			
			Total	1.65	0.06	140.6	4.9	8.44	0.41			
	Total Cosmogenic								3.43	0.63		

5.2 Step-heat

Because the crush-fusion method was not especially successful in isolating the cosmogenic component, we attempted step heating on two samples from the stratigraphically youngest flow (2Na, samples VM-01 and VM-03) and one from the stratigraphically oldest flows (Flow 1N, sample VM-10). Results are shown in Table 3 and Fig 2. In all three samples the ³He/⁴He ratios decline monotonically with extraction temperature, while the ⁴He yields increase. For example, in sample VM-

03, the 800, 1000, and 1400°C steps yielded 3 He/ 4 He ratios of 13.1, 9.0, and 8.2 R_A, and 4 He concentrations of 15, 20, and 70 ncc/g, respectively. Previous work indicates that mantle He in fluid inclusions is released at extraction temperatures >1000°C (Kurz, 1986a, b; Swindle et al., 2023). In the case of the VM samples, the 3 He/ 4 He ratios extracted in the final steps (7.87 \pm 0.22, 8.17 \pm 0.23, 7.79 \pm 0.22 R_A) are indistinguishable from the mean 3 He/ 4 He ratio obtained by crushing (8.10 \pm 0.20 R_A), supporting this conclusion.

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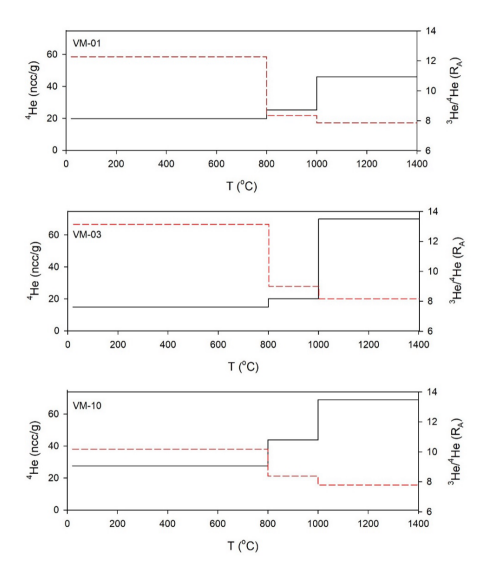


Figure 2. Step heat results on three of the VM samples. Samples were analyzed at three successive temperatures (800, 1000, 1400 °C) held for one half hour each. Horizontal lines indicate the temperature range over which each measurement integrates. Red dashed line indicates ³He/⁴He ratio (right hand axis) while black solid line indicates ⁴He concentration (left hand axis).

6 Discussion

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6.1. Tightly Retained Mantle He in Volcano Mountain Olivine

³He/⁴He ratios of He released by crushing and in the highest temperature step of the step heats are within the typical range of 215 mid-ocean ridge basalts (Graham et al., 1992), suggesting He in Volcano Mountain derives from the upper mantle. Regardless of its origin, the presence of this mantle He substantially complicates determination of cosmogenic ³He exposure ages. While mantle ⁴He concentrations of crushed olivines are sometimes, but not often (Kurz et al., 1990; Farley and Neroda, 1998; Fenton and Niedermann, 2014; Heineke et al., 2016), as high as the ~100 ncc STP/g we obtained on the VM samples, the more 220 important complication arises from the fact that this mantle He is not effectively removed by powdering to <30 um. Our powder fusion analyses indicate that about a third of the mantle component remains in the powdered olivine in every case. Our results suggest a mantle component that survives the powdering step, most likely in <<30 µm fluid inclusions. Evidence to support this interpretation cones from our observations of trails of circular voids (<10 um) present in backscatter electron images of VM olivine xenocrysts. We interpret these to be secondary inclusion trails that carry mantle He. Survival of the 225 mantle component when crushing this fine is not typical, but it is similar to results obtained on Twin Sisters peridotites (Swindle et al., 2023). Another possibility is that the ubiquitous kink banding and sub-grain boundaries observed in VM (Francis & Ludden, 1990) and Twin Sisters olivines (Swindle et al., 2023) could contribute to mantle He retention.

6.2 Component separation by step heating

The step-heat method takes advantage of the different release kinetics of matrix-sited and fluid inclusion-sited He. Diffusion alone is adequate to release matrix-sited helium; in contrast, for fluid inclusion helium to escape the crystal requires it to first dissolve into the olivine, an energetically disfavored process (Trull and Kurz 1993). Three temperature steps allowed us to be flexible about the temperature that best separates these two components. The evolution of the ³He/⁴He ratio with temperature thus can reveal two-endmember mixing between mantle and cosmogenic He (assuming radiogenic and nucleogenic helium are both absent).

Compared to the ³He/⁴He ratios obtained on powder fusions, the ³He/⁴He ratios of the lowest temperature steps in our stepheat experiments are between 1.5 and 4 R_A higher (Tables 2, 3). In contrast, ³He/⁴He ratios of the 1000°C steps are very similar to the powder fusion values, and about 0.7 R_A higher than the highest temperature steps and the associated crush-released ³He/⁴He ratios. These data provide support for the interpretation that all cosmogenic He is released prior to the highest temperature step, while mantle He is released in progressively greater amounts as step temperature increases.

To calculate the cosmogenic ³He concentration from the step heat results for each sample, we assumed the final step represents the mantle component, used equation 1 to compute the cosmogenic ³He yield in the two lower temperature steps, and combined

them to obtain a total. Using this method, we obtained indistinguishable ${}^{3}\text{He}_{c}$ concentrations on the two samples of flow 2Na (3.71 \pm 0.41 and 3.38 \pm 0.33 Mat/g), and 3.43 \pm 0.63 Mat/g on flow 1N. While still uncertain (average \pm 13%), these cosmogenic concentrations are significantly better determined than by the crush/fusion method.

It is possible that a more detailed step heat could yield even greater degrees of separation between cosmogenic and mantle helium components, but at the potential cost of analytical precision associated with averaging multiple analyses and with blank corrections. Were significant amounts of radiogenic and/or nucleogenic helium present in these samples (unlikely given the eruption age of \sim 10 ka, see below), this method would yield inaccurate results in the same way that the crush-fusion method would.

6.4 ³He_c Exposure Ages

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Exposure ages in Table 4 were computed from ${}^3\text{He}_c$ concentrations and corresponding uncertainties using the CRONUS-Earth online calculator (Balco et al., 2008; Phillips et al., 2016; version 3) where the production rate of ${}^3\text{He}_c$ is calibrated using data from Borchers et al. (2016). Exposure ages were obtained for four of the flows. By far the best determined is the exposure age of the youngest flow, 2Na. The two step heat analyses (VM-01, 03) yield indistinguishable ages of 12.3±1.9 and 11.1±1.6 ka. These are in good agreement with the three crush-fusion pairs (VM-01, 02, 03) with ages ranging from 9±4.2 to 13.3±6.1 ka. Assuming independent uncertainties among these analyses, the weighted means of these five ages is 11.4 ± 2.2 ka (MSWD 0.17, 95%), slightly more uncertain than the step heat analyses on their own. Four analyses were performed on flow 1N, with the step heat experiment yielding an age of 14.0±3.0 ka and three crush-fusion pairs yielding ages between 2.3±4.8 and 7.4±9.9 ka. The weighted mean of these analyses is 8.7 ± 4.3 ka (MSWD 2.3, 7.4%), again more uncertain than the step heat analysis by itself. The individual crush-fusion analyses of flows 2Sa and 1Sa yielded exposure ages of 9.6 ±1.9 ka and 10.0 ±3 ka, respectively.

Exposure ages are indicated on the map in Figure 1 using the weighted means when available. All four flows yield the same exposure age within error (weighted mean of all analyses: 10.5 ± 1.7 ka, MSWD 0.92, 51%) and there is no indication that the stratigraphically lower flows (1Sa and 1N, weighted mean 9.2 ± 3.5 ka, MSWD 1.8, 13%) have older exposure than the higher flows (2Na and 2Sa, weighted mean 11.0 ± 1.9 ka, MSWD .28, 93%).

The simplest interpretation of these results is that all the studied VM lava flows erupted at 10.5 ± 1.7 ka. Alternatively, honoring the stratigraphy the flows may have erupted over an age range from about 12.7 ka (1S flows) to about 9.1 ka (2Na and 2 Sa).

Table 4

				³ He _c		Exp Age	
	Flow	Sample	Method	Mat/g	±	ka	±
280	2Na	VM-01	C/F	2.75	1.25	9.0	4.2
		VM-02	C/F	3.27	1.57	10.8	5.3
		VM-03	C/F	4.05	1.82	13.3	6.1
		VM-01	StepHt	3.71	0.41	12.3	1.9
		VM-03	StepHt	3.38	0.33	11.1	1.6
		weighted	mean			11.4	2.2
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	2Sa	VM-06	C/F	2.19	0.36	9.6	1.9
	1Sa	VM-08	C/F	2.27	0.64	10.0	3.0
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270	1N	VM-10	C/F	0.46	1.17	1.9	4.8
		VM-11	C/F	0.56	1.15	2.3	4.8
		VM-11	C/F	1.79	2.37	7.4	9.9
		VM-10	StepHt	3.43	0.63	14.0	3.0
		weighted	mean			8.7	4.3

7 Conclusion

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Volcano Mountain nephelinites contain olivine likely sourced from disaggregated peridotite xenoliths and are thus a unique set of samples compared to the more commonly studied olivines grown in basaltic melts. This olivine contains high concentrations of upper mantle He and powdering does not effectively remove this component. Swindle et al. (2023) were similarly unable to remove mantle He components by powdering olivine from the Twin Sister peridotite. In both cases, this mantle He may be housed in fluid inclusions too small to be released by crushing. Alternatively, ubiquitous kink bands and subgrain boundaries seen in the Twin Sisters and Volcano Mountain flows could be the source of the He that survives powdering. In either case, survival of mantle He in VM powder fusions caused the crush-fusion method to yield imprecise ³He_c concentrations. By using the step heating method described here, we were able to determine that the morphologically youngest lava flows on Volcano Mountain erupted approximately coevally in the earliest Holocene, 10.5 ±1.7 ka.

Data Availability

All data is provided in the figures and tables of this manuscript.

Author Contribution

J. M. Mueller prepared the manuscript with significant contributions from K. A. Farley and J. D. Bond. J. D. Bond and B.C310 Ward completed the field sampling.

Competing Interests

The authors declare that they have no conflict of interest.

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