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Petrology of a 2.41 Ga remarkably fresh komatiitic basalt lava lake in Lion Hills, central Vetreny Belt, Baltic Shield

Received: 8 December 1995/Accepted: 26 March 1996

Abstract In the central Vetreny Belt, southeastern Baltic Shield, an areally extensive 110 m deep lava lake is exposed consisting of remarkably fresh differentiated komatiitic basalt. During eruption, the liquid had a temperature of 1380–1400 °C and contained $\sim 15\%$ MgO. The lava ponded in a large topographic depression soon after eruption. The differentiation of the lava lake was controlled by settling of transported olivine and chromite phenocrysts and caused the origin of prominent internal layering. The last portions of the trapped liquid crystallized at temperatures of 1250-1070 °C. A Sm-Nd isochron of 2410 + 34 Ma for whole rock samples, olivine, augite and pigeonite separates from the lava lake provides a reliable estimate for the time of formation of the uppermost sequences in the Vetreny Belt. This age is in good agreement with the Sm-Nd and Pb-Pb isochron ages of 2449 ± 35 and 2424 + 178 Ma for the volcanic rocks from the same stratigraphic level in the northwestern Vetreny Belt. Modeling of Nd-isotopes and major and trace elements shows that the komatiitic basalts at Lion Hills may have had a komatiite parent depleted in highly incompatible elements. It can be shown that this initial liquid

was contaminated by 7–9% of Archaean upper crustal material from the adjacent Vodla and Belomorian Blocks *en route* to the surface thus acquiring the observed geochemical and isotope signatures including relative enrichment in Zr, Ba, and LREE, negative Nb-and Ti-anomalies and ε Nd(T) of -1.

Introduction

The results from integrated studies of mafic-ultramafic volcanic suites are generally regarded as the basis for modelling the evolution of the composition and physical state of the Earth's mantle and crustal reservoirs (e.g., Arndt 1986a; Chase and Patchett 1988; Condie 1989; Dupré and Arndt 1990; Galer and Goldstein 1991; Jochum et al. 1991; Nisbet et al. 1993; Abbott 1994). The ambiguity of interpretation of these results with respect to Archean settings is mainly caused by the inadequate state of preservation of the object of investigation due to the effects of hydrothermal alteration and polyphase metamorphism.

All komatiitic and associated mafic volcanic rocks are more or less altered. In some cases the primary minerals are replaced pseudomorphically, and the important details of morphology of original minerals can still be recognized precisely and thus the precursor igneous minerals can be readily identified. But in most cases, recrystallization becomes complete and little remains of the primary texture. This makes it difficult to infer even the original nature of the premetamorphic protoliths of these rocks. When studying the chemical composition of such rocks, it is necessary to penetrate the veil of serpentinization or evev worse, the dense fog of amphibolite and granulite facies metamorphism. Uniquely fresh by Archean standards, the komatiites from Alexo, Ontario (Arndt 1986b) and especially from Zwishavane, Zimbabwe (Nisbet et al. 1987; Renner et al. 1994) can be indeed regarded as the window that

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Editorial responsibility: J. Hoefs

allows a glance into the depths of the Archean mantle and helps us to understand the mechanisms of formation of the Earth's early crust.

In the present paper we report the results of detailed field, petrographic, geochemical and Nd-isotope studies of the remarkably well preserved early Proterozoic high-magnesian volcanic rocks from the Vetreny Belt in the southeastern Baltic Shield. The objectives of the research include: (a) field study of a large differentiated body of komatiitic basalt in order to reconstruct its original nature; (b) detailed examination of mineralogy, petrography and chemistry of minerals and main rock varieties across the complete section in order to deduce the composition of the initial and erupted magma and the factors that controlled its differentiation after emplacement; (c) Nd-isotope studies of bulk rocks and primary minerals in order to raise further the curtain concealing the enigmatic problem of the age of the Vetreny Belt voleanism.

Geological background

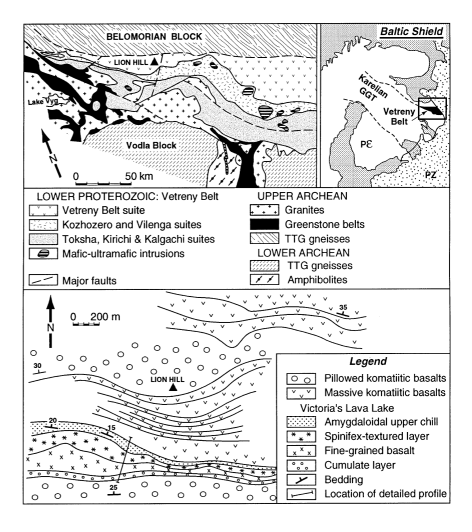
In the southeastern Baltic Shield, three large early Precambrian structural and lithological units are juxtaposed. They include Ar-

chean amphibolite-gneiss-migmatite and granite-greenstone units, and a Lower Proterozoic volcanic-sedimentary unit. The Archean units are volumetrically dominant and constitute the Karelian granite-greenstone terrane, which occupies a total area of approx. 350000 km² (Fig 1).

About 2.5 Ga ago, the Archean continental crust of the Karelian granite-greenstone terrane underwent tectono-magmatic reactivation during the Karelian stage (2.5–2.1 Ga). This episode of reactivation was initiated by continental rifting, as evidenced by the deposition of conglomerates, quartzites and arkoses. Subsequently areally extensive mafic volcanic rocks of the Sumi-Sariola group erupted, followed by the emplacement of mafic-ultramafic layered intrusions and rapid sedimentation in fault-bounded shallow-water basins (Gaál and Gorbatschev 1987; Gorbatschev and Bogdanova 1993). Most of the Sumian-Sariolian rocks occur in NW-trending volcanosedimentary belts, their deposition and magmatism being controlled by NW-trending rift-formed fault systems. One of these large belts the Vetreny Belt (Fig. 1) is the subject of this study. This belt can be traced from lake Vyg southeastward over a distance of more than 250 km; its width increases from 15 to 85 km. In the southeast, the Vetreny Belt plunges under the Paleozoic cover of the Russian platform, and extends further on to the southeast (Kulikov 1988). In the northeast, the Belt is separated from the Belomorian Block by the Northern deep fault zone. The southwestern contact of the Vetreny Belt with the Sumozero-Kenozero greenstone belt and the Vodla Block is also tectonic in character.

The early Proterozoic rocks of the Vetreny Belt are subdivided into six major suites (Fig. 1). The lowermost part of the sequence is represented by the terrigenous (quartzites) Toksha suite, andesites

Fig. 1 Geological sketch map of the Vetreny Belt. Geologic-petrographic map of the Lion Hills area



and basalts of the Kirichi suite and polymict conglomerates and sandstones of the Kalgachi suite. These are overlain by the terrigenous sediments and basalts of the Kozhozero suite, and further upward in the succession, by the tuffaceous and terrigenous Vilenga suite and the Vetreny Belt suite; the latter is composed entirely of komatiitic basalts. The total thickness of the early Proterozoic sequences in this area varies between 4 and 8 km (Kulikov 1983; Sokolov 1987).

This study focuses on the high-magnesian volcanic rocks from the Vetreny Belt suite in the Lion Hills area (Fig. 1). These rocks were recognized and described as high-magnesian basalts by Kulikov (1971) and Slyusarev and Kulikov (1973). Kulikov and Kalinin (1971) reproduced the chilled textures observed in these basalts experimentally and made an attempt to explain the mechanisms responsible for these spectacular textures. Almost at the same time these textures became referred to as spinifex textures (Nesbitt 1971).

In the Lion Hills area, lava flows are dipping consistently to the NNE at angles of $20 \text{ to } 40^{\circ}$, and range in thickness from a few meters to several tens of meters. The metamorphic grade in the central part of the area surrounding Lion Hill does not exceed the prehnite-pumpellyite facies. The rocks in this area are well exposed and are characterized by the superb state of preservation of primary minerals. The northern slope of the hill is made up of massive and pillowed komatititic basalts. On the southern side, regular lava flows of the same type are also exposed in addition to a thick differentiated

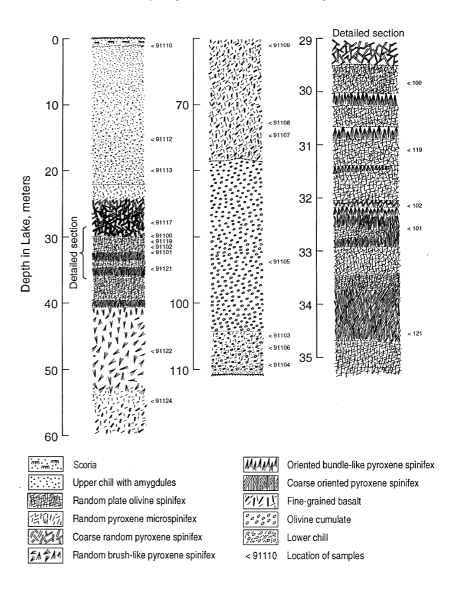
lava unit. Based on field ralationships and petrologic observation this lava unit is interpreted as the remnants of an areally extensive and deep lava lake. After one of its disoverers we name this unit Victoria's lava lake.

This ancient lava lake was documented on the nearly vertical walls of a little canyon that cuts the lake across strike. The lake is gently plunging to the north at 15 to 30°, and can be traced along strike over a distance of more than 2.5 km; in the west and east it is covered by Quaternary glacial deposits. Its thickness decreases from 150 m in the western part down to 60 m in the eastern part of the hill; in the best exposed part within the canyon it is around 110 m deep. The rocks that make up this lava lake are characterized by a wide variety of structures and textures (Fig. 2). In general, the lake consists of four main units (from top downward): (1) scoria; (2) upper chilled margin; (3) spinifex zone, which includes the spinifex subzone proper and the subzone of a fine-grained basalt; (4) cumulate zone.

Petrography of the lava lake

Before describing the petrography of the lava lake, it is necessary to make a few introductory statements concerning the terminology used throughout this paper. To describe crystals having a substantially larger size than those in the groundmass, the term

Fig. 2 Detailed profile through Victoria's lava lake



phenocryst (>0.5 mm) and microphenocryst (<0.5 mm) are adopted. Individual olivine, pyroxene and chromite morphology was described using the terminology introduced by Lofgren et al. (1974) and Donaldson (1976, 1982). Among the olivines, three morphological types were distinguished: (a) solid euhedral equant to tabular crystals; (b) equant to elongate hopper (externally euhedral to subhedral but internally hollow and incomplete) crystals, sometimes with perceptible edge growth; (c) plate olivine. The pyroxene was documented as: (a) prismatic and blocky crystals; (b) columnar and acicular hollow crystals; (c) chain crystals; (d) plumose and feather-like crystals.

Scoria

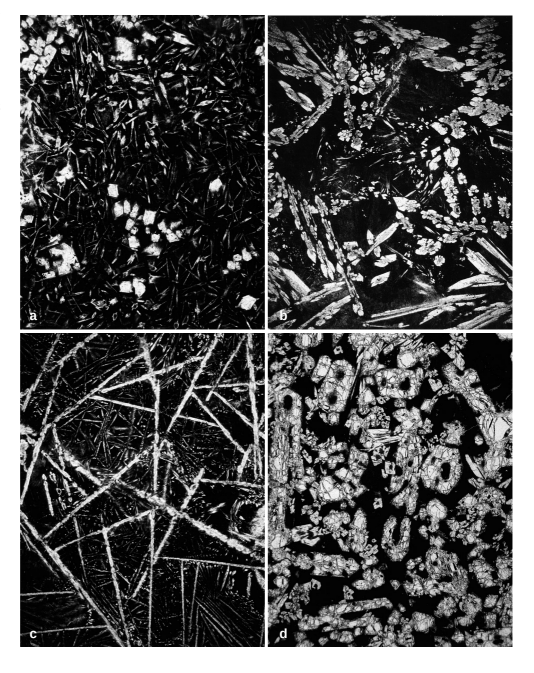
The scoria is a thin ($\sim 0.5 \, \mathrm{m}$) layer containing rounded and amoeboid to subangular lighter-colored clasts up to 2 cm in diameter cemented by cryptocrystalline basaltic material. The clasts

consis of devitrified volcanic glass containing tiny solid euhedral and cruciform (euhedral core and snowflake-like overgrowths) chromite grains and serpentinized olivine crystals. The latter are subdivided into two groups: (1) small (0.1–0.3 mm) solid and incomplete equant euhedral microphenocrysts in clusters of 7–10 grains that make up about 15% of the rock by volume; (2) larger (0.6–2.5 mm) solid and rarely incomplete equant to subequant phenocrysts constituting some 10% of the rock.

Upper chilled margin

In the upper chilled margin, olivine is almost completely serpentinized and is represented by both pheno- and microphenocrysts and a tiny mesh of hopper bladed crystals 0.1–1 mm in length, set in a brownish matrix after glass and constituting the olivine microspinifex texture (Fig. 3a). The chilled margin reveals a heterogeneous

Fig. 3a–d Photomicrographs of representative types of rocks, plane-polarized light: a upper chilled margin, sample 91110, 5.8 mm across the photograph; b coarse random pyroxene spinifex layer, sample 91117, 11.6 mm across the photograph; c plate olivine spinifex layer, sample 91118, 11.6 mm across the photograph; d Olivine cumulate layer, sample 91105, 5.8 mm across the photograph. See text for explanations



texture owing to the presence of thin layers variably enriched in chlorite-serpentine-feldspar amygdules, 1–1.5 mm in diameter.

Spinifex zone

The upper part of the spinifex zone is largely made up of a layer of coarse random pyroxene spinifex. It is characterized by development of long (up to 30 mm) and ~ 1 mm thick hollow columnar and chain zoned pyroxene crystals, and euhedral to subhedral hopper olivine crystals 2-7 mm long in a matrix of devitrified glass and plumose and chain pyroxene (Fig. 3b). The plate olivine spinifex is composed of thin (0.1–0.3 mm) plate olivine crystals up to 25 mm in length oriented nearly orthogonal to each other and imparting a cell-like structure to the rock (Fig. 3c). The cells are filled with devitrified glass and plumose and chain augite. The plate olivine spinifex layer contains thin (12-25 cm) bands of bundle-like pyroxene spinifex and thicker layers of coarse oriented pyroxene spinifex. In the bundle-like spinifex, acicular pyroxene crystals up to 70 mm in length are arranged in bundles, or sheaves nucleating at the upper contact of the layer and splaying downward. The groundmass consists of devitrified glass, plumose pyroxene and cruciform chromite. The oriented coarse pyroxene spinifex ("string beef" spinifex after Arndt et al. 1977) is characterized by the development of up to 12 cm long and 0.5-1 mm thick columnar zoned pyroxene crystals arranged roughly parallel to each other and subperpendicular to the top of the lake. The interstices are filled with devitrified glass, plumose pyroxene, cruciform chromite and tiny plagioclase

The random brush-like pyroxene spinifex layer consists of columnar and chain pyroxene dominating over rare olivine platelets up to 15 mm in length immersed within aggregates after devitrified glass. Toward the bottom of the layer, olivine disappears, and the abundance and grain size of pyroxene decreases, the rock thus displays a microcrystalline texture and grades down into fine-grained basalt. The latter consists of tiny (0.5–2 mm) plagioclase laths and short-prismatic clinopyroxene. Cumulate zone.

The contact between the spinifex and cumulate zone is abrupt, with dramatic changes in mineralogy and textural relationships. The cumulate zone is subdivided into the olivine cumulate subzone proper and a basal subzone. The olivine cumulate is composed of two morphological types of olivine. The first type, which constitutes 3/4 of the whole olivine population, is represented by solid rounded, polyhedral or elongated crystals 0.2-2.5 mm in size. Often they reveal a lighter-colored core and darker edges. The second type is present in the form of hollow or incomplete crystals 0.4–3 mm in size. The internal parts of the crystals are filled with glass (Fig. 3d). Pyroxene occurs in the form of small $(0.2 \times 1 \text{ mm})$ incomplete longprismatic and blocky, sometimes zoned crystals. Small euhedral chromite crystals are present as either inclusions in olivine, or dispersed in glass. The crystalline phases constitute 50-60% of the rock volume. The brownish glass is mostly isotropic indicating a very good state of preservation.

The basal subzone is distinct in having a crystalline matrix consisting of fine-grained pyroxene, tiny plagioclase laths and disseminated equant chromite. Embedded within the groundmass are solid rounded and euhedral olivine phenocrysts 0.4–2 mm in size (15–20% of the rock volume) and small (0.2–0.4 mm) skeletal or larger (0.5–2 mm) irregular pigeonite and augite grains, many of which contain rounded olivine as poikilitic inclusions.

Analytical techniques

Mineral fractions were separated at the Geological Institute in Petrozavodsk. About 150 mg of pure high-Ca pyroxene and 300–500 mg of low-Ca pyroxene and olivine separates were then obtained at IGEM in Moscow by handpicking the $\sim 99\%$ pure concentrates.

Microprobe studies were carried out at the Mineralogisch-Petrographisches Institut, University of Cologne, using a Cameca Camebax electron microprobe. Operating conditions were $15\,\mathrm{kV}$ accelerating voltage, $20\,\mathrm{nA}$ beam current and integration time of $10\,\mathrm{s}$ per element. To obtain information on compositional zoning, traverses across mineral grains in steps of several tens of $\mu\mathrm{m}$ were performed, and $5\,\mathrm{to}$ $10\,\mathrm{points}$ were analyzed on some grains.

Chemical studies were undertaken on 150 g aliquots from 1–3 kg of fresh sample. Major element abundances were determined by X-ray fluorescence on fused glass pellets using a Philips PW-1404 at the Johannes Gutenberg University in Mainz; trace elements were analyzed on pressed powder pellets on the same XRF-machine. The REE concentrations were determined at IGEM in Moscow by isotope dilution mass-spectrometry using the method described by Thirlwall (1982) and modified by Zhuravlev et al. (1989). Accuracies are estimated as follows: major elements – 2 per cent for elements present in concentrations greater than 0.5%; trace elements – 5 per cent for Zr, Cr, V, Co, Ni and Ba; 10 per cent for Y and Sr; 20 per cent for Nb and Rb. Multiple analyses of the BCR-1 standard established the accuracy for La to be 2%, and for the remaining REE as better than 2%.

The Sm-Nd isotope investigations were carried out at the Max-Planck Institut für Chemie in Mainz following techniques described by Chauvel et al. (1985). Both Nd and Sm were run on a MAT-261 Finnigan mass-spectrometer under static mode. The total blank was 0.05 ng for Nd and 0.01 ng for Sm. The effects of fractionation Nd runs were eliminated by normalizing to $^{146}\text{Nd}/^{144}\text{Nd} = 0.7219$. The $^{143}\text{Nd}/^{144}\text{Nd}$ isotope ratio measurements of the La Jolla standard during the period of data collection yielded a value of 0.511831 ± 20 ($2\sigma_{pop}$; n=21). All Nd-isotope ratios were bias corrected to the La Jolla standard 143 Nd/ 144 Nd = 0.511860. Regression analyses followed the method of York (1966), and the error on the initial ϵ_{Nd} value was calculated using the method of Fletcher and Rosman (1982). The analytical uncertainties used in the regression calculations were 0.2% for $^{147}Sm/^{144}Nd$ and for $^{143}Nd/^{144}Nd$, the $2\sigma_{pop.}$ value for the La Jolla standard corresponding to its external reproducibility. All errors on ages and initial isotopic ratios are quoted at 2σ , or 95% confidence level. The Sm-Nd age was calculated using a decay constant for $^{147}{\rm Sm}$ of $6.54\times10^{-12}\,a^{-1}.$

Results

Composition of minerals

Representative microprobe analyses of minerals and volcanic glass are given in Tables 1 to 4.

Olivine

In the upper chilled zone, only a few relict olivine phenocrysts remain unaltered. In the cumulate zone, however, olivine is remarkably well preserved and was therefore studied in more detail. In general, olivine composition is dependent on the size and morphological type of the crystals. Larger solid phenocrysts have a slightly more magnesian composition than microphenocrysts. Hopper crystals show the lowest forsterite content. Cores of larger phenocrysts are characterized by rather uniform compositions and a decrease in Fo content within thin outer rims; microphenocrysts and especially hopper grains reveal continuous normal zoning from the center toward the edges of the crystals.

Table 1 Representative microprobe analyses of olivines from Victoria's lava lake. (*Hopp* hopper, *L* large, *S. Ph* small polyhedral olivine grains respectively)

Sample no. Morphology Location	91112 Hopp Core	91112 Hopp Core	91105 L. Ph Core	91105 L. Ph Core	91105 L. Ph Rim	91105 L. Ph Rim	91105 S. Ph Core	91105 S. Ph Core	91105 Hopp Core	91105 Hopp Core	91105 Hopp Rim
SiO ₂	40.77	40.28	39.94	39.70	40.33	40.01	40.62	40.27	39.57	40.04	40.07
TiO_2	0.030	0.000	0.025	0.002	0.012	0.000	0.008	0.026	0.000	0.002	0.000
Al_2O_3	0.012	0.040	0.036	0.025	0.058	0.067	0.060	0.073	0.040	0.031	0.046
Cr_2O_3	0.140	0.200	0.098	0.130	0.096	0.072	0.096	0.091	0.073	0.061	0.078
FeO	10.91	10.67	11.79	12.34	12.51	13.79	12.49	12.16	13.76	14.05	15.86
MnO	0.130	0.177	0.199	0.178	0.170	0.177	0.140	0.191	0.164	0.204	0.237
NiO	0.310	0.280	0.282	0.290	0.288	0.220	0.286	0.291	0.304	0.214	0.211
MgO	48.06	48.07	45.94	45.78	45.89	45.36	46.02	46.07	45.18	45.40	43.33
CaO	0.370	0.300	0.243	0.236	0.237	0.246	0.260	0.246	0.260	0.233	0.282
Total	100.7	100.0	98.54	98.68	99.59	99.94	99.98	99.42	99.35	100.2	100.1
% Fo	88.7	88.93	87.42	86.87	86.74	85.43	86.79	87.11	85.40	85.21	82.96
Sample no.	91105	91103	91103	91103	91106	91106	91106	91106	91106	91106	91106
Morphology	Hopp	L. Ph	L. Ph	L. Ph	L. Ph	L. Ph	L. Ph	L. Ph	L. Ph	L. Ph	L. Ph
Location	Rim	Core	Core	Rim	Core	Core	Rim	Core	Core	Core	Core
SiO ₂	39.63	40.38	40.91	40.14	40.00	40.51	40.26	40.19	40.07	39.80	40.12
TiO_2	0.002	0.000	0.019	0.028	0.000	0.002	0.000	0.012	0.007	0.000	0.000
$Al_2\tilde{O}_3$	0.028	0.056	0.040	0.039	0.017	0.029	0.018	0.052	0.060	0.038	0.063
Cr_2O_3	0.088	0.077	0.073	0.064	0.096	0.048	0.043	0.067	0.101	0.111	0.116
FeO	16.40	11.67	11.75	13.59	10.81	11.22	12.22	10.98	11.15	11.26	11.47
MnO	0.283	0.191	0.106	0.181	0.143	0.180	0.176	0.141	0.179	0.112	0.181
NiO	0.225	0.304	0.332	0.283	0.223	0.222	0.249	0.300	0.247	0.341	0.288
MgO	42.49	46.71	47.42	45.18	47.46	47.91	47.72	47.86	46.73	47.22	47.76
CaO	0.271	0.211	0.236	0.237	0.220	0.158	0.191	0.229	0.259	0.238	0.229
Total	99.42	99.59	100.9	99.74	98.98	100.3	100.9	99.83	98.80	99.12	100.2
% Fo	82.21	87.71	87.80	85.56	88.67	88.39	87.44	88.60	88.19	88.20	88.13

In the upper chilled margin, hopper crystals have compositions of $Fo_{88.7-88.9}$. Due to the very small size of the relict grains, it was not possible to investigate the compositional zoning in the crystals. In the olivine cumulate subzone proper, cores of larger polyhedral grains have compositions varying from $Fo_{87.4-86.3}$ to $Fo_{85.4}$ at the margins. Small phenocrysts have a rather constant forsterite content of $Fo_{87.1-86.3}$. The compositions of hopper crystals vary from $Fo_{85.4-85.2}$ in the cores to $Fo_{81.7}$ at the rims. In the basal chill subzone, cores of large phenocrysts have the most magnesian compositions recorded in the cumulate zone $(Fo_{88.7})$ and are identical to the hopper grains in the upper chilled zone. Small phenocrysts show a wide range from $Fo_{87.8-86.7}$ in the cores to Fo_{82} at the rims.

Pyroxenes

The pyroxenes are compositionally similar to those from komatiite complexes elsewhere (Arndt et al. 1977; Arndt and Fleet 1979; Arndt 1986b; Renner et al. 1994). Skeletal blocky crystals from the groundmass of the olivine cumulate subzone proper are unzoned and cor-

respond to high-Al (5.4–8.0% Al $_2O_3$) augite Wo $_{45}$ En $_{44}$ Fs $_{11}$ –Wo $_{38}$ En $_{46}$ Fs $_{16}$. In the basal subzone, skeletal crystals of both low-Al (1.7–2.5% Al $_2O_3$) augite Wo $_{42}$ En $_{49}$ Fs $_{10}$ –Wo $_{34}$ En $_{52}$ Fs $_{14}$ and magnesian pigeonite Wo $_{4}$ En $_{75}$ Fs $_{21}$ –Wo $_{19}$ En $_{64}$ Fs $_{17}$ were documented. Tiny crystals of chain skeletal pyroxene from the plate olivine spinifex layer are, as a rule, unzoned and correspond to high-Al (5.4–7.1% Al $_2O_3$) augite Wo $_{45}$ En $_{41}$ Fs $_{14}$ –Wo $_{36}$ En $_{51}$ Fs $_{13}$. In contrast, large columnar hollow crystals from the coarse random and oriented pyroxene spinifex layers are always zoned with pigeonite (Wo $_{11}$ En $_{76}$ Fs $_{13}$ –Wo $_{8}$ En $_{78}$ Fs $_{14}$) cores, and outer portions composed of moderately Al-rich (2.2–5.2% Al $_2O_3$) augite Wo $_{30}$ En $_{60}$ Fs $_{10}$ –Wo $_{40}$ En $_{50}$ Fs $_{10}$.

Chromite

Two morphological types of chromite were documented. Cruciform skeletal crystals mainly occur in the spinifex zone, while solid equant euhedral grains are typical of the cumulate zone. Both types of chromite are similar and correspond to alumochromites. In the Cr_2O_3 versus Al_2O_3 and TiO_2 diagrams, the chromites

Table 2 Representative microprobe data for clinopyroxenes from Victoria's lava lake (Aug augite, Pig pigeonite)

Sample no.	91105	91105	91104	91104	91106	91106	91106	91106	91106	91106
Morphology	Prism									
Composition	Aug	Aug	Aug	Pig	Pig	Aug	Aug	Pig	Aug	Pig
SiO ₂ TiO ₂ Al ₂ O ₃ Cr ₂ O ₃ FeO MnO NiO MgO	49.85	49.40	51.99	54.22	53.45	51.32	52.83	53.70	51.61	53.16
	0.594	0.796	0.639	0.235	0.310	0.759	0.395	0.242	0.551	0.239
	5.54	5.89	2.17	1.30	1.26	1.90	1.70	0.91	1.92	1.13
	0.111	0.244	0.437	0.251	0.186	0.108	0.255	0.205	0.356	0.126
	7.23	6.23	7.90	12.14	11.78	8.89	8.92	13.40	9.04	14.36
	0.164	0.145	0.180	0.313	0.300	0.193	0.216	0.270	0.252	0.319
	0.032	0.070	0.043	0.048	0.050	0.070	0.066	0.084	0.036	0.101
	15.04	15.00	17.28	25.54	24.19	17.37	21.47	26.37	18.37	25.08
CaO	19.79	20.44	18.67	5.46	7.88	18.16	13.32	3.48	16.97	4.08
Na ₂ O	0.181	0.221	0.220	0.022	0.113	0.242	0.135	0.053	0.189	0.070
Total	98.53	98.43	99.53	99.51	99.53	99.02	99.30	98.72	99.30	98.66
Wo	42.7	44.3	38.2	10.8	15.5	36.9	26.6	6.9	34.2	8.1
En	45.1	45.2	49.2	70.4	66.3	49.0	59.6	72.5	51.5	69.5
Fs	12.2	10.5	12.6	18.8	18.1	14.1	13.9	20.7	14.2	22.3
Sample no.	91118	91118	91118	91117	91117	91123	91123	91101	91101	91101
Morphology	Chain	Chain	Chain	Colum						
Composition	Aug	Aug	Aug	Pig	Aug	Aug	Aug	Aug	Aug	Pig
		Chain	Chain	Colum						

occupy the low-Cr and high-Al and high-Ti corners of the komatiite field distinguished by Zhou and Kerrich (1992), Herbert (1982) and Jan and Windley (1990), while in the Cr₂O₃ versus FeO plot they are almost outside the range of FeO contents observed in komatiites and in continental layered intrusions at given chromium abundances. Compared to chromites from the more magnesian counterparts elsewhere they are therefore enriched in Al₂O₃ and FeO and depleted in MgO.

Volcanic glass

Glass was analyzed from the cumulate subzone proper and from the plate olivine spinifex layer in places where it shows the least degree of alteration and therefore represents the closest approximation to the composition of the latest portions of trapped liquid. As can be seen in Table 4, it displays a wide compositional range with SiO₂ varying between 49.3–55%, and MgO from 2.7–7.9%.

Chemical compositions of the rocks

For chemical and Nd-isotope studies, 18 samples were collected along the detailed profile (Fig. 2). Major, trace, and rare earth element data normalized to 100% on a volatile-free basis are listed in Table 5 and shown in the variation diagrams Figs. 4 and 5. The MgO content ranges from 26.1% in the cumulate zone to 7.3% in the fine-grained basalt layer. In the upper part of the lake, the MgO abundances are rather uniform and change slightly from 14.7% at the top to 13.7% at the bottom of the upper chilled margin. The spinifex zone proper contains 10.3–8.8% MgO.

Most components show a strong correlation with MgO (r = 0.99-1.00 for Ti, Al, Zr, Y, V, Ni, and REE, and 0.97–0.98 for Ca and Sr) and vary in a manner entirely consistent with olivine fractionation. When plotted against wt% MgO, the analytical data fall on olivine control lines (Fig. 4), which intersect the MgO axes at $47.9 \pm 0.7\%$ ($2\sigma_{\text{mean}}$). Sample 91113 from the bottom of the upper chilled margin contains abundant

Table 3 Representative microprobe data for chromites from Victoria's lava lake. All crystals analyzed are solid equant grains. (ND not detected)

Sample no.	91105	91105	91105	91105	91105	91106	91117
SiO ₂	0.14	0.76	0.08	0.10	0.09	0.04	0.06
TiO_2	0.430	0.497	0.439	0.435	0.450	0.909	0.519
Al_2O_3	13.16	13.62	13.95	13.17	13.73	8.86	14.99
Cr_2O_3	46.46	46.25	47.25	46.63	47.40	45.89	43.67
FeO	35.21	34.05	32.81	33.48	32.70	38.36	36.18
MnO	ND						
NiO	0.047	0.011	0.028	0.047	0.066	0.020	0.045
MgO	2.29	2.83	2.77	2.55	3.01	1.67	0.54
CaO	0.09	0.01	0.00	0.04	0.04	0.07	0.08
Total	97.83	98.03	97.33	96.45	97.49	95.82	96.08
Si ⁴⁺	0.0047	0.0265	0.0026	0.0034	0.0033	0.0015	0.0022
Ti ⁴⁺	0.0113	0.0130	0.0115	0.0116	0.0118	0.0250	0.0140
A1 ^{3 +}	0.5430	0.5570	0.5749	0.5496	0.5644	0.3815	0.6322
Cr ³⁺	1.2858	1.2687	1.3062	1.3059	1.3069	1.3257	1.2357
Fe^{2+}	0.8755	0.8531	0.8546	0.8623	0.8401	0.9059	0.9669
Fe ³⁺	0.1551	0.1348	0.1047	0.1295	0.1137	0.2663	0.1159
Ni ²⁺	0.0013	0.0003	0.0008	0.0013	0.0019	0.0006	0.0013
Mg^{2+} Ca^{2+}	0.1197	0.1463	0.1446	0.1348	0.1566	0.0907	0.0288
Ca ²⁺	0.0035	0.0003	0.0000	0.0016	0.0015	0.0027	0.0030
Cr/(Cr + Al)	0.703	0.695	0.694	0.704	0.698	0.777	0.662
Mg/							
(Mg + Fe)	0.120	0.146	0.145	0.135	0.157	0.091	0.029

Table 4 Representative microprobe data for volcanic glass (*ND* not detected)

Sample no. Location	91105 Matrix	91105 Matrix	91105 Matrix	91105 Hollow olivine	91105 Matrix	91118/1 Matrix	91118/1 Matrix
SiO ₂	49.25	51.59	49.64	53.16	51.66	55.16	51.89
TiO_2	1.890	0.826	0.954	0.882	0.517	0.782	0.797
Al_2O_3	16.98	17.08	15.70	17.97	18.32	20.24	18.18
Cr_2O_3	0.02	0.04	0.05	0.02	0.02	0.02	0.03
FeO	11.12	9.02	11.71	9.60	9.21	5.34	10.07
MnO	0.456	0.200	0.243	0.204	0.150	0.090	0.068
NiO	0.032	ND	ND	0.011	0.043	0.015	ND
MgO	6.74	5.79	7.93	4.96	4.88	2.66	4.00
CaO	8.52	9.94	8.32	8.72	9.78	8.41	10.16
Na ₂ O	3.49	3.68	2.80	4.03	4.10	5.40	3.05
K_2O	0.11	0.24	1.10	0.18	0.11	0.05	0.18
Total	98.61	98.41	98.46	99.74	98.78	98.17	98.43

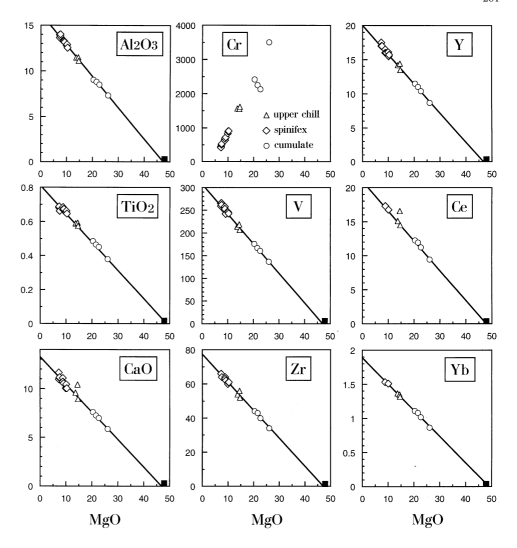
amygdules. It lies well above the olivine control lines and was excluded from the following discussion and petrologic modeling. Nickel shows a behavior similar to that of MgO, revealing a strong negative correlation with the above components (r=0.99-1.00). When plotted against Ni, the olivine control lines give intercepts at 2410 ± 70 ppm ($2\sigma_{\rm mean}$); this value corresponds to the average Ni content in the liquidus olivine (Table 1). Chromium is positively correlated with MgO, which suggests that along with olivine, chromite was a liquidus phase over the whole compositional range of the crystallization of the lake. The alkalis show either poor correlation with MgO (r=0.82 for Na₂O), or mostly irregular behavior (K₂O, Rb), and are regarded as having been mobile during the low-grade metamorphism.

Ratios CaO/Al₂O₃ (0.82 \pm 0.01), Al₂O₃/TiO₂ (19.6 \pm 0.3), and Ti/Y (249 \pm 4, 2 σ_{mean}) are close to

chondritic values of 0.80, 22, and 275, respectively (Hart and Zindler 1986; McDonough and Sun 1995) while Zr/Y ratios (3.86 \pm 0.04) are substantially higher, and Ti/Zr (64.4 \pm 0.8) lower than those in chondrites (2.5 and 110), implying significant enrichment of the rocks in Zr relative to Ti and Y.

Shown in Fig. 5 are patterns for selected major, trace and rare earth elements normalized to primitive mantle values of Hofmann (1988). The patterns are strongly parallel, with abundances reversely correlated with MgO content. In line with the enrichment in Zr, the rocks are also enriched in Sr, Ba, and LREE [chondrite normalized (La/Sm)_N = 2.1–2.3] and reveal strong negative Nb- and Ti-anomalies [(Nb/Ba)_N = 0.12 \pm 0.01, (Nb/La)_N = 0.26 \pm 0.02]. Europium shows slight negative anomalies (Eu/Eu* = 0.94–0.97).

Fig. 4 Variation diagrams of selected major and trace elements for Victoria's lava lake. The calculated and measured liquidus olivine composition is shown by *solid square*



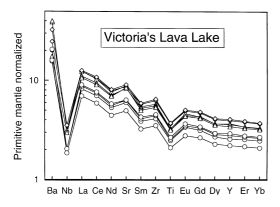


Fig. 5 Abundances of selected major, trace and rare earth elements normalized to primitive mantle values of Hofmann (1988) in Victoria's lava lake. Legend same as in Fig. 4

Sm-Nd isotope data

Sm-Nd isotope data are listed in Table 6 and plotted on the conventional Sm-Nd diagram (Fig. 6). Bulk rock samples representing the whole section through the lake from top to bottom as well as olivine, augite and pigeonite mineral separates define a well-constrained [mean standard weighted deviates MSWD = 1.1] isochron with a slope corresponding to an age of 2410 ± 34 Ma and a negative initial ϵ Nd(T) value of -0.9 ± 0.2 .

Discussion

Lake, flow or sill?

In this paper, the thick differentiated unit containing olivine cumulate layer was referred to as a lava lake. The most convincing evidence that it is an extrusive body and not a sill comes from such crucial indicators of volcanic nature as the asymmetric structure typical of komatiitic lavas (e.g., Arndt et al. 1977; Arndt 1986b; Renner et al. 1994) and the presence of the upper breccia, or scoria – a prominent rock consisting of vitreous blocky fragments of lava. In contrast to polygonal jointing, these fragments show displacement

Table 5 Major, trace and rare earth element data for Victoria's lava lake. The analyses are normalized to 100% on an anhydrous basis (*LOI* loss on ignition).

Sample no.	91100	91101	91102	91103	91104	91105	5 9110	06 911	107 91	108 91	109 91	110 911	112 91113
SiO2	51.6	52.4	51.7	48.2	47.9	46.0	47.8	52.9	52.6	52.1	50.3	50.5	49.3
TiO ₂	0.646	0.672	0.670	0.485	0.464	0.379	0.451	0.662	0.662	0.681	0.575	0.589	0.593
Al_2O_3	12.5	13.2	13.3	9.0	8.8	7.3	8.5	13.8	14.0	14.0	11.1	11.5	11.5
Fe_2O_3	11.5	11.1	11.0	11.9	11.7	12.3	11.7	10.6	10.6	11.2	12.1	11.6	11.3
MnO	0.194	0.191	0.193	0.170					0.194	0.204	0.185	0.173	
MgO	10.3	8.85	8.85	20.4	21.6	26.1	22.6	7.54	7.55	7.29	14.6	13.7	14.5
CaO	10.0	10.7	10.6	7.62	7.26	5.88	7.00	11.1	11.2	11.7	8.96	9.59	10.4
Na ₂ O	2.90	2.21	3.07	1.89	1.68	1.62	1.28	2.79	2.73	2.67	1.63	2.03	1.90
K_2O	0.17	0.47	0.48	0.26	0.32	0.22	0.39	0.29	0.35	0.20	0.43	0.35	0.26
P_2O_5	0.08	0.09	0.09	0.08	0.07	0.06	0.07	0.11	0.11	0.11	0.08	0.09	0.09
LOI	0.70	0.19	0.17	1.22	0.69	2.04	0.89	0.22	0.25	0.16	2.50	1.50	2.42
Cr	909	688	628	2417	2255	3507	2131	502	525	447	1619	1570	1552
V	243	253	258	176	167	137	161	266	262	265	207	214	219
Co	45	47	39	87	91	107	96	35	39	34	67	56	58
Ni	216	119	106	801	874	1138	952	68	70	61	408	348	386
Zr	61	63	64	44	43	34	40	64	64	66	52	54	56
Nb	2.2	2.2	2.2	1.3	1.3	1.1	1.3	2.6	2.6	2.6	1.8	2.0	2.0
Y	15.7	16	16	11.5	11	8.7	10.1	17	17	17.5	13.5	14.2	14.4
Sr	164	165	172	116	115	90	109	164	175	176	154	152	161
Rb	5	8	7	5	7	4	9	6	7	4	10	9	5
Ba	190	197	180	97	125	95	123	180	184	190	106	242	236
La		7.75		5.54	5.38	4.27	5.05				6.52	6.78	7.60
Ce		17.3		12.3	11.9	9.5	11.2				14.5	15.1	16.6
Nd		9.63		6.91	6.68	5.30	6.34				8.24	8.45	9.09
Sm		2.28		1.65	1.59	1.26	1.50				1.94	2.03	2.10
Eu		0.734		0.533	0.508	0.404	0.494				0.634	0.672	0.646
Gd		2.47		1.75	1.71	1.36	1.61				2.13	2.19	2.21
Dy		2.63		1.88	1.83	1.46	1.73				2.26	2.34	2.32
Er		1.62		1.15	1.12	0.896	1.06				1.37	1.42	1.42
Yb		1.53		1.11	1.08	0.865	1.02				1.32	1.37	1.35
Mg/(Mg + Fe)	0.639	0.612	0.614	0.773	0.785	0.808	0.793	0.584	0.585	0.564	0.706	0.700	0.717
Ti/Zr	63.5	64.0	62.8	66.1	64.7	66.9	67.5	62.0	62.0	61.8	66.3	65.4	63.5
Zr/Y	3.89	3.94	4.00	3.83	3.91	3.91	3.96	3.76	3.76	3.77	3.85	3.80	3.89
(La/Nb) _N		3.6		4.4	4.3	3.7	4.0				3.6	3.5	3.9
(La/Sm) _N		2.14		2.11	2.13	2.14	2.12				2.12	2.10	2.28
(Gd/Yb) _N		1.30		1.27	1.27	1.27	1.28				1.30	1.30	1.32
Eu/Eu*		0.945		0.957	0.943	0.946					0.954	0.974	

Table 6 Nd-isotopic data for Victoria's lava lake (WR bulk rock samples, OI olivine, Aug augite, Pig pigeonite mineral separates respectively)

Sample no.	Sm, ppm	Nd, ppm	$^{147} Sm/^{144} Nd$	$^{143}Nd/^{144}Nd$	εNd(2410)
91103 WR	1.631	6.923	0.14238	0.511739 ± 8	- 0.71
91103 Aug	1.325	3.724	0.21508	0.512884 ± 8	-0.90
91103 Ol	0.1738	0.625	0.16819	0.512156 ± 9	-0.57
91104 WR	1.603	6.808	0.14233	0.511710 ± 7	-1.3
91104 Aug	1.397	3.976	0.21238	0.512826 ± 7	-1.2
91104 Pig	0.563	1.917	0.17751	0.512284 ± 9	-0.97
91106 WR	1.522	6.381	0.14420	0.511751 ± 7	-1.0
91106 Aug	1.265	3.699	0.20681	0.512753 ± 8	-0.90
91106 Ol	0.0492	0.168	0.17708	0.512291 ± 11	-0.69
91101 WR	2.306	9.731	0.14322	0.511737 ± 7	-1.0
91105 WR	1.248	5.209	0.14475	0.511764 ± 4	-0.96
91117 WR	2.255	9.516	0.14324	0.511743 ± 8	-0.90

relative one another, i.e., it is a true breccia. The fragments are interpreted to have been detached from the continuously solidifying foundered roof and incorporated within the upper part of flowing lava. Both ancient lava lakes and their modern equivalents were shown to contain scorias (Arndt 1986c; Wright et al. 1968; Helz et al. 1989). We interpret this volcanic unit as a lava lake owing to its large thickness and high-Mg composition. High-magnesian liquids have a very low viscosity (Arndt 1976; Huppert et al. 1984), which prevents them

91117	91119	91121	91122	91124
51.3	51.2	51.4	51.5	52.5
0.662	0.651	0.684	0.673	0.688
12.9	12.8	13.2	13.2	13.6
11.7	11.5	11.6	11.4	11.0
0.201	0.204	0.194	0.205	0.203
10.1	9.94	8.81	9.16	7.30
10.4	10.0	11.1	10.6	11.0
2.52	3.41	2.36	2.81	3.14
0.13	0.19	0.52	0.45	0.53
0.09	0.09	0.12	0.10	0.09
0.18	0.32	0.40	0.39	0.21
898	837	653	721	417
246	247	250	242	259
53	55	52	53	42
174	193	112	122	54
61	60	63	62	66
2.2	2.2	2.2	2.2	2.2
15.5	16.1	16.5	16	17
158	172	171	166	169
3	5	5	5	8
150	160	205	187	190
7.59				
16.8				
9.43				
2.23				
0.716				
2.41				
2.58				
1.58				
1.51				
0.632	0.632	0.601	0.613	0.568
65.1	65.1	65.1	65.0	62.5
3.94	3.73	3.82	3.88	3.88
3.5	5.75	2.02	2.00	2.00
2.14				
1.29				
0.943				
0.5 1.5				

from forming thick continuous lava bodies unless they are ponded. As can be seen in Fig. 1, the studied unit represents an unusually thick body in this area, which probably formed under uniquely favorable conditions. These conditions were obviously not met for the other lava units, which are much thinner and virtually undifferentiated. Field relations suggest that the lake is getting thicker to the west and is gradually pinching out to the east. It might well have been that its shape was controlled by a large topographic low or a previously formed trough-like structure.

Composition and evolution of the erupted liquid

According to the petrographic observations, the only minerals present as phenocrysts or cumulus phases are olivine and chromite. It is very likely therefore, that the evolution of the primary magma after eruption was

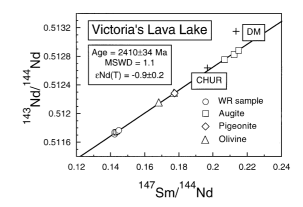


Fig. 6 Conventional Sm-Nd diagram for Victoria's lava lake. The present-day parameters of the chondrite uniform reservoir (*CHUR*) were adopted from Jacobsen and Wasserburg (1980, 1984) to be the following: $^{147} \mathrm{Sm}/^{144} \mathrm{Nd} = 0.1967, ^{143} \mathrm{Nd}/^{144} \mathrm{Nd} = 0.512638.$ The parameters of the model depleted mantle reservoir (*DM*) are $^{147} \mathrm{Sm}/^{144} \mathrm{Nd} = 0.2135, ^{143} \mathrm{Nd}/^{144} \mathrm{Nd} = 0.513151$ and correspond to the $\epsilon \mathrm{Nd}_{\mathrm{(DM)}}(\mathrm{T}=0) = +10$ and the time of fractionation from CHUR 4.6 Ga ago

controlled by fractionation of these minerals only. This conclusion is also supported by the analysis of the variation diagrams, in which all elements but Ni and Cr are reversely correlated with MgO.

In order to deduce the liquid compositions, several independent approaches were used. The amygdule-free chilled sample 91110, containing 14.7% MgO, is essentially a glassy rock, consisting of phenocrysts in a quenched groundmass. Obviously, it did not undergo any post-eruption differentiation and is interpreted to have a bulk composition very similar to that of the liquid from which it has formed. The second approach to estimate the composition of the primary lava was to calculate the liquid compositions using olivine data. The most magnesian olivines (Fo_{88.7-88.9}) were documented in the upper and lower chilled margins of the lake. If a $K_{D(ol-liq)}^{Mg-Fe^{-1}} = 0.31$ (Beattie et al. 1991) is adopted, then, taking into account the Mg/Fe relationships in the studied komatiitic basalts, these olivines can be shown to have been in equilibrium with a liquid containing 14.8–15.2% MgO. The liquid and equilibrium olivine compositions can also be inferred using an indirect approach based on the general differentiation trends established for the lava lake. In the plots MgO versus an element that is excluded from the olivine crystal lattice $[K_{D(01-lig)} \leq 1]$, the analytical data for samples containing different proportions of olivine crystallized from a common parental melt fall on the mixing lines (olivine control lines). The intercept of these regression lines with the MgO axis corresponds directly to the average composition of liquidus olivine. For the studied suite of samples, this value is equal to 47.9 + 0.7% MgO. Recalculating for the olivine stoichiometry, this gives an olivine composition of Fo_{88.7}, in excellent agreement with the results of the olivine microprobe studies. Thus three independent lines of evidence indicate that komatiitic basalts from the Lion Hills area erupted with liquid compositions of $\sim 15\%$ MgO.

Eruption temperature, thermal history and internal differentiation of the lava lake

In order to calculate the eruption temperature of the komatiitic basalt melt, the procedures described by Abbott et al. (1994) and Nisbet et al. (1993) are particularly suitable. The chemical analyses of the upper chilled margin were used to calculate the eruption temperature on the basis of the well-known assumption that the Fe/Mg ratio in a primitive magma is directly related to its liquidus temperature (Roeder and Emslie 1970; Hanson and Langmuir 1978). Calculations show that the komatiitic basalt magma erupted onto the surface with liquidus temperatures of 1380–1400 °C (Table 7). The liquidus temperatures for the samples from the spinifex zone vary from 1320–1310 °C for the plate olivine spinifex layer to 1310-1280 °C for the pyroxene spinifex and to 1250 °C for the fine-grained basalt layer. In order to obtain an estimate for the solidus temperatures of crystallization of the lava lake, we applied the above procedure to the analyses of the fresh volcanic glass (Table 4), which represents the last portions of the trapped liquid. Calculated temperatures vary over a wide range between 1270–1070 °C depending on the composition of the glass. The crystallization interval for the lake is therefore estimated to be between 1400-1070 °C.

The main liquidus phases in the lava lake are olivine and chromite. The partitioning of Fe and Mg between these two minerals also can be utilized to estimate the eruption temperature of the magmas. We used Lehmann's equation to calculate the liquidus temperatures (Lehmann 1983). In order to avoid disequilibrium bias, we studied only intergrown or adjacent olivine and chromite grains. All studied samples come from the cumulate zone. The temperatures obtained range from 970 to 1040 °C (Table 8) and are hence lower than those calculated from the MgO content in the primary liquid or inferred from the experimental studies (Arndt 1976; Thy 1995), while being similar to crystallization temperatures for the last and most evolved portions of the trapped liquid.

The temperature estimates based on the two-pyroxene thermometer are listed in Table 8. The selected samples come from the basal chilled margin and the middle part of the spinifex zone, where skeletal augite and pigeonite grains are found to occur in close spatial association. The estimated temperatures range from 1185 to 1225 °C using the thermometer of Lindsley (1983). These temperatures are slightly lower than the liquidus temperatures of the pyroxene spinifex-textured rocks, but agree well with the experimental results of Kinzler and Grove (1985) and Thy (1995) and are interpreted to correspond to the crystallization temperatures of the augite-pigeonite pairs.

From the field relationships and the petrographic observations it can be concluded that a magnesian (15% MgO) komatiitic basalt liquid, containing 15 to 20% olivine phenocrysts during the eruption filled a large deep topographic depression, located distinctly to the side of the main lava channel. As a result, a large volume of magma ponded and stopped flowing. This in turn allowed efficient olivine settling to form the layered structure of the lake. Judging from the continuous, undisturbed structure of the lake, it was probably filled to its present depth in a large single pulse or several smaller but almost contemporaneous pulses of magma. Many of the polyhedral olivine grains in the lava lake are regarded as being transported by the lava as phenocrysts and crystallized before emplacement. Only a smaller part of the whole population, mostly hopper crystals and overgrowths on the polyhedral grains, were formed in situ. This is suggested by the uniform and close compositions of phenocryst cores both in the upper chilled margin and cumulate zone, implying their slow equilibrium crystallization from a large volume of a magma prior to eruption.

After phenocrysts were removed from the part of the lake immediately below the upper chilled margin, the spinifex zone started to form proceeding from the top downward. The wide textural and mineralogical variability of this zone at largely constant chemical composition is attributed mainly to a different degree of supercooling attained in each successive layer in the crystallizing liquid, and to its boundary composition (from 10 to 9% MgO), at which olivine is replaced by clinopyroxene on the liquidus (Arndt 1976). The reverse sequence of crystallization, when clinopyroxene is replaced by olivine on the liquidus, as evidenced by the intercalation of olivine and pyroxene spinifex layers, can be explained by temperature increase at the front of crystallization caused by the heat released during very fast spinifex pyroxene nucleation. Besides,

Table 7 Results of liquidus temperatures calculations for Victoria's lava lake

Sample	91110	91112	91113	91100	91101	91102	91107
Tliq(°C)	1398	1381	1393	1317	1280	1287	1247
Sample	91108	91109	91117	91119	91121	91122	91124
Tliq(°C)	1248	1239	1309	1313	1282	1293	1247
Sample	Glass 1	Glass 2	Glass 3	Glass 4	Glass 5	Glass 6	Glass 7
Tliq(°C)	1230	1200	1274	1170	1168	1068	1127

Table 8 Calculated temperatures from olivine-chromite and augite-pigeonite pairs

Sample	91105 (1)	91105 (2)	91105 (3)	91105 (4)	91105 (5)
			Chromite		
A1 ^{3 +}	0.5430	0.5570	0.5749	0.5496	0.5644
Cr ³⁺	1.2858	1.2687	1.3062	1.3059	1.3069
Fe ²⁺	0.8755	0.8531	0.8546	0.8623	0.8401
Fe ³⁺	0.1551	0.1348	0.1047	0.1295	0.1137
Mg^{2+}	0.1197	0.1463	0.1446	0.1348	0.1566
J			Olivine		
Fe ²⁺	0.2606	0.2937	0.2852	0.303	0.2895
Mg^{2+}	1.7233	1.6915	1.699	1.6808	1.6939
T(°C)	969	1038	1018	1025	1035

Sample	91104 (1)	91104 (2)	91106 (1)	91106 (2)	91117
			Augite		
Si ⁴⁺	1.9251	1.9192	1.9062	1.9061	1.9330
Ti ⁴⁺	0.0130	0.0177	0.0212	0.0153	0.0096
Al ^{IV}	0.0749	0.0808	0.0832	0.0835	0.0670
Al ^{VI}	0.0336	0.0136	0.0000	0.0000	0.0853
Cr ³⁺	0.0118	0.0127	0.0032	0.0104	0.0088
Fe ³⁺	0.0149	0.0348	0.0657	0.0664	0.0000
Fe ²⁺	0.1823	0.2093	0.2105	0.2129	0.2452
Mn^{2+}	0.0053	0.0056	0.0061	0.0079	0.0063
Mg^{2+} Ca^{2+}	0.9270	0.9510	0.9616	1.0113	0.9861
Ca ²⁺	0.8005	0.7382	0.7227	0.6716	0.6742
Na +	0.0114	0.0158	0.0175	0.0135	0.0151
			Pigeonite		
Si ⁴⁺	1.9763	1.9620	1.9493	1.9505	1.9613
Ti ⁴⁺	0.0087	0.0064	0.0090	0.0066	0.0031
Al ^{IV}	0.0237	0.0380	0.0507	0.0490	0.0387
Al ^{VI}	0.0348	0.0175	0.0028	0.0000	0.0349
Cr ³⁺	0.0091	0.0072	0.0061	0.0037	0.0131
Fe ³⁺	0.0000	0.0022	0.0289	0.0377	0.0000
Fe ²⁺	0.3950	0.3651	0.3355	0.4028	0.2836
Mn^{2+}	0.0091	0.0096	0.0109	0.0099	0.0063
Mg^{2+}	1.2737	1.3776	1.3194	1.3715	1.5100
Ca ²⁺	0.2984	0.2115	0.2806	0.1604	0.1577
Na+	0.0036	0.0016	0.0051	0.0050	0.0028
T(°C)	1185	1200	1215	1230	1225

unstable pyroxene crystallization could also occur from a liquid with > 9% MgO owing to the high degrees of supercooling attained (Arndt and Fleet 1979; Campbell and Arndt 1982). The fine-grained basalt layer immediately overlying the cumulate zone and having a remarkably uniform composition represents the residual liquid and crystallized at the final stages of formation of the lava lake as indicated by the lowest liquidus temperatures (1240–1250 °C) and MgO content (7.3%) obtained for this layer. This layer is similar to the lowermost part of spinifex-textured zones in more magnesian komatiite lavas elsewhere (e.g., Arndt 1986b; Renner et al. 1994)

The observed mineral morphology and temperature estimates can be used to place some constraints on the thermal history of the lava lake. As shown by experimental data of Lofgren et al. (1974) and Donaldson (1976; 1982), the change in shape of olivine and

pyroxene crystals as a function of cooling rate and degree of supercooling is systematic and common to all melts investigated. From our study of crystal morphology in the lava lake it can be inferred that the cooling rates and the degree of supercooling were highly variable, from $40-80\,^{\circ}\text{C/h}$ ($\Delta T = 50-80\,^{\circ}\text{C}$) in the spinifex zone to $1.2-15\,^{\circ}\text{C/h}$ ($\Delta T = 0-30\,^{\circ}\text{C}$) in the cumulate zone. It is apparent, however, that the cooling rates became much slower after a large proportion of cumulus olivine had settled and the grains thus formed a self-supporting framework. Convection of the liquid was then mostly inhibited and cooling proceeded dominantly by means of conduction.

Turner et al. (1986) and Gole et al. (1990) noted that the center of thick extrusive bodies remains at a temperature close to the liquidus for some tens of years depending on their thickness, while the top cools rapidly to the ambient surface temperature. These conclusions are also supported by direct cooling rate measurements from the Kilauea Iki lava lake, Hawaii (Helz et al. 1989). This lava lake formed in 1959 as a large pond of picritic basalt (15% MgO) and has cooled and crystallized as a self-roofed magma chamber approximately 115 meters deep. All processes of large-scale internal differentiation occurring in the lake ceased sometime between 1979 and 1981. If the crystallization time interval for Victoria's lava lake, which has about the same depth and primary liquid composition, was of the same order of magnitude, it becomes clear that the central part of the body cooled very slowly. Therefore it is not unexpected that cation exchange in the olivine-spinel pairs ceased late and this system was finally frozen at nearly solidus temperatures.

Sm-Nd isotope data and the age of the lava lake

The Sm-Nd isochron age of 2410 \pm 34 Ma represents a reliable estimate of the crystallization age of the lava lake in particular and, more generally, the timing of accumulation of the Vetreny Belt suite. This in turn provides an estimate for the final stages of magmatism in the Vetreny Belt. The whole rock samples reveal a very narrow range in Nd-isotope compositions and Sm/Nd ratios throughout the lava lake, implying that the erupted melt was chemically very homogeneous and no contamination occurred after emplacement of the lava onto the surface. Within analytical uncertainty, this age is in good agreement with the Sm-Nd and Pb-Pb isochron ages of 2449 + 35 and 2424 + 178 Ma, respectively, obtained for volcanic rocks from the same suite in Golets Hills, northwestern Vetreny Belt (Puchtel et al. 1991).

As can be seen in Table 6, the olivine, pigeonite and augite separates are characterized by relatively high Nd

and Sm concentrations. Assuming that abundances of these elements in sample 91110 correspond to those in the erupted melt, it is possible to calculate theoretical contents of these elements in the studied olivine and low- and high-Ca pyroxenes using published partition coefficients. As can be seen from Table 9, the measured values for olivine are 30–250 times, for pigeonite 8–15 times, and for augite 1.5–3 times higher than the theoretical abundances. These discrepancies are likely to be ascribed to the presence of tiny glass inclusions in the mineral separates, mostly inside hopper crystals. Calculations show that some 1–2% of these inclusions would be enough to account for all the excess Nd. Although these inclusions cannot shift the isotope equilibrium between the crystals and the liquid, and, therefore, do not alter the interpretation of the isochron, they have substantial effect on the Sm/Nd ratio in the mineral separates. As seen in Table 9, the measured ¹⁴⁷Sm/¹⁴⁴Nd in the olivine is 2.5 times, in the pigeonite - almost 2 times, and in augite - 7–30% lower than the calculated values.

Composition of parental magmas and isotope characteristics of the mantle source region

Despite the obvious compositional differences between the continental crust and mantle, the effect of crustal contamination is not so easily recognizable. This is ascribed to the fact that, at least in some instances, the influence of crustal contamination is almost indistinguishable from the geochemical effects on basaltic magmas derived from a depleted asthenospheric source and then mixed with a component of the ancient continental lithosphere (Glazner and Farmer 1992; Brandon et al. 1993). Ryabchikov et al. (1988) suggested that komatiitic basalt magmas of the Vetreny Belt were

Table 9 Comparison of calculated and measured Nd and Sm concentrations in mineral separates

Mineral separate	Nd, ppm	Sm, ppm	$^{147} Sm/^{144} Nd$	Reference
	F	Partition coeffcice	ents used	
Olivine	0.0003	0.0009		Green 1994
Low-Ca Pyroxene	0.015	0.035		Green 1994
Ca-Pyroxene	0.15	0.30		Green 1994
-	0.19	0.30		Hart and Dunn 1993
	0.30	0.54		Mc Rae and Russel 1987
C	oncentrations a	and ratios calcula	ited (1) and measur	red (2)
Melt (2)	8.24	1.94	0.142	
Olivine (1)	0.0025	0.0017	0.411	Green 1994
91103 Olivine (2)	0.625	0.174	0.168	
91106 Olivine (2)	0.168	0.0492	0.177	
Low-Ca Pyroxene (1)	0.124	0.0679	0.332	Green 1994
91104 Pigeonite (2)	1.92	0.563	0.178	
Ca-Pyroxene (1)	1.24	0.582	0.284	Green 1994
-	1.57	0.582	0.224	Hart and Dunn 1993
	2.47	1.05	0.256	Mc Rae and Russel 1987
91103 Augite (2)	3.72	1.33	0.215	
91104 Augite (2)	3.98	1.40	0.212	
91106 Augite (2)	3.70	1.27	0.207	

derived from the enriched continental lithospheric mantle because of the apparent large-scale homogeneity of the lavas. Our more extensive studies indicate however that these lavas are very variable in terms of trace element and isotope characteristics (Puchtel IS, Haase KM, Hofmann AW, Chauvel C, Kulikov VS, Garbe-Schönberg D, submitted). We argue that the evolution of magmas parental to the Vetreny Belt komatiitic basalts was mainly controlled by assimilation of crustal rocks *en route* to the surface.

Due to very high liquidus temperatures and low viscosities, komatiitic melts are very susceptible to contamination by crustal rocks (e.g., Arndt 1986a; Arndt and Jenner 1986; Cattell 1987; Lesher and Arndt 1995). Contamination normally occurs during ascent or eruption (Huppert et al. 1984; Huppert and Sparks 1985) and results in a sharp increase in abundances of the elements concentrated in the upper continental crust, e.g., Ba, Zr, and LREE, in primary asthenospheric melts, but will have little effect on the concentrations of elements Ti, Nb and HREE. On average, this causes the origin of negative Nb- and Ti-anomalies in crustally contaminated rocks (Jochum et al. 1991). In contrast, the subcontinental mantle lithosphere is enriched in Nb relative to La (McDonough 1990). The studied volcanic rocks exhibit pronounced negative Nb- and Tianomalies relative to neighboring elements with a comparable degree of incompatibility, i.e., La, Ba and Zr $[(Nb/Ba)_N = 0.12 \pm 0.01, (Nb/La)_N = 0.26 \pm 0.02, (Ti/Zr)_N]$ = 0.59 ± 0.01 , $2\sigma_{\text{mean}}$]. These relationships are most consistent with contamination of parental depleted komatiite magma with felsic material of the upper continental crust.

The upper crust in the Vodla and Belomorian Blocks consists of TTG-gneisses and amphibolites metamorphosed under amphibolite to granulite facies conditions and have ages of 3.1–3.2 and 2.7–2.8 Ga, respectively (Kulikov et al. 1990; Lobach-Zhuchenko et al. 1993; Bogdanova and Bibikova 1993; Timmerman and Daly 1995; Bibikova et al. in press). Trace element concentrations for these rocks, however, are not available and in order to prove or reject the assimilation hypothesis and to estimate the extent of contamination of the initial magma, the trace element composition of

the bulk felsic Archean granulite reported by Rudnick and Fountain (1995) were used. The composition of this bulk analysis provides the closest approximation to the average composition of the upper crustal rocks in the Vodla and Belomorian Blocks area in terms of major elements. The average Nd-isotopic composition for these rocks (143 Nd/ 144 Nd = 0.510746) was adopted from the data of Kulikov et al. (1990), Lobach-Zhuchenko et al. (1993), Timmerman and Daly (1995), and Bibikova et al. (in press). As a possible equivalent to the primitive melt giving rise to the Lion Hills lavas, the composition of the 2.7 Ga old komatiite from the Belingwe greenstone belt (Nisbet et al. 1987; Jochum et al. 1991; Arndt et al. 1993) was chosen. This komatiite has 17.6% MgO, 143 Nd/ 144 Nd = 0.513197, ϵ Nd(T) = +2.5, $(La/Sm)_N = 0.8$ and was suggested to have the composition close to that of the primary magma for the Vetreny Belt lavas based on major, trace element and olivine data (Puchtel IS, Haase KM, Hofmann AW, Chauvel C, Kulikov VS, Garbe-Schönberg D, submitted).

The results of the calculations are shown in Fig. 7. We used the ratios of selected elements, which are insensitive to fractional crystallization but are crucial indicators of assimilation processes. As can be seen in the diagrams Fig. 7, the erupted magma could have been derived from mixing of such an incompatible element depleted komatiite melt with upper crustal rocks. The inferred degrees of contamination are very consistent for different elemental ratios and vary slightly between 7 and 9%. Such contamination would have had only little effect on the major element composition of the initial magma (Table 9). The negative Eu-anomalies observed in the rocks were also shown to have been the results of crustal contamination. Based on these data therefore, little can be deduced with respect to the Nd-isotope composition of the mantle source region of the komatiitic basalts from the lava lake.

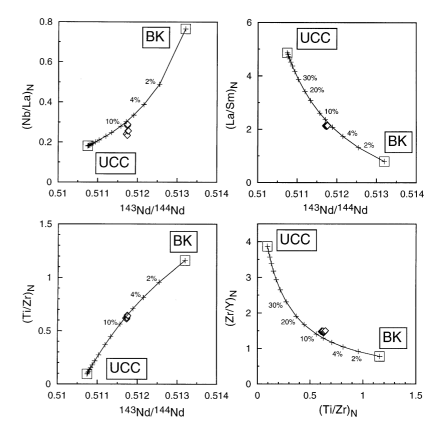
Concluding remarks

1. Remarkably fresh by early Precambrian standards, komatiitic basalts from the uppermost part of the Vetreny Belt provide a sound source of information on the

Table 10 Results of mixing calculations for major elements. Numbers correspond to percentage contamination. [BK Belingwe komatiite (Nisbet et al. 1987; Jochum et al. 1991), UCC felsic upper continental crust after Rudnick and Fountain (1995)]

	BK	2%	4%	6%	8%	10%	20%	30%	40%	UCC
SiO ₂ TiO ₂ Al ₂ O ₃ FeO MgO CaO	49.65	50.07	50.48	50.90	51.32	51.74	53.82	55.91	57.99	70.50
	0.430	0.429	0.429	0.428	0.428	0.427	0.424	0.421	0.418	0.400
	9.01	9.12	9.23	9.33	9.44	9.55	10.09	10.63	11.17	14.40
	11.74	11.57	11.39	11.22	11.05	10.88	10.01	9.15	8.28	3.10
	17.59	17.26	16.93	16.60	16.27	15.94	14.29	12.64	10.99	1.10
	8.79	8.67	8.55	8.43	8.31	8.19	7.59	6.99	6.39	2.80
$\begin{array}{c} \mathrm{CaO/Al_2O_3} \\ \mathrm{Al_2O_3/TiO_2} \end{array}$	0.976	0.951	0.927	0.903	0.880	0.858	0.753	0.658	0.573	0.194
	20.95	21.23	21.51	21.80	22.08	22.36	23.79	25.24	26.71	36.00

Fig. 7 Diagrams (Nb/La)_N, (La/Sm)_N, (Ti/Zr)_N versus ¹⁴³Nd/¹⁴⁴Nd and (Zr/Y)_N versus (Ti/Zr)_N showing the compositions of Victoria's lava lake komatiitic basalts (diamonds) and mixing lines between an inferred komatiite parent and a crustal contaminant. (BK Belingwe komatiite, UCC upper continental crust). Numbers next to tick marks give percentage contaminant



origin of high-magnesian volcanic rocks in general, and in this volcanic pile in particular.

- 2. The differentiated body of komatiitic basalt studied represents an areally extensive 110 m deep lava lake. During eruption, the initial liquid had a temperature of 1380–1400 °C, contained ~ 15% MgO and ponded in a large topographic depression soon after emplacement. Evolution of the erupted magma was controlled by settling of transported olivine and chromite phenocrysts and resulted in the prominent internal layering. The last portions of the trapped liquid crystallized at 1250 to 1070 °C. As evidenced by two-pyroxene and olivine-spinel thermometry, these mineral systems were frozen between 1185–1225 and 970–1040 °C, respectively.
- 3. The internal Sm-Nd isochron of 2410 \pm 34 Ma provides a reliable estimate for the time of formation of the uppermost sequences in the Vetreny Belt. This age is in good agreement with the Sm-Nd and Pb-Pb isochron ages of 2449 \pm 35 and 2424 \pm 178 Ma for the volcanic rocks from the same stratigraphic level in the Golets Hills area, northwestern Vetreny Belt.
- 4. Modeling of isotopes, major and trace elements can be used to show that the komatiitic basalts at Lion Hills may have had an incompatible trace element-depleted komatiite parent. This initial liquid was shown to have undergone 7–9% contamination by upper crustal rocks *en route* to the surface thus acquiring the observed geochemical and isotope signatures

[relative enrichment in Zr, Ba, and LREE, negative Nb- and Ti-anomalies and negative $\varepsilon Nd(T)$ value].

Acknowledgements The handpicking of mineral separates was carefully and patiently done by V.Pavlichenko and N.Ashikhmina. I.S.P. thanks Prof. H.A. Seck for support during microprobe work at the Mineralogisch-Petrographishes Institut in Cologne. Discussions with Dallas Abbott and Svetlana Bogdanova were most beneficial. Critical comments of Heinz Stosch, Peter Sorjonen-Ward and an anonymous reviewer substantially improved the initial draft of the manuscript. I.S.P. acknowledges the support of the Alexander von Humboldt Foundation.

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