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Experimental constraints on pre-eruption conditions of pantelleritic magmas: Evidence from the Eburru complex, Kenya Rift

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Abstract

The phase relationships and compositions of a pantellerite from the Eburru complex in the Kenya Rift Valley have been determined at 150 MPa and under reducing conditions, 2 log units below the Ni–NiO solid buffer. The effects of temperature and melt water content on phase relationships have been explored. Alkali feldspar and quartz crystallise alone at temperatures above 700 °C, irrespective of melt water content. Below 700 °C, sodic amphibole and clinopyroxene also crystallise; the amphibole being the liquidus phase under water-rich conditions. The coexistence of amphibole phenocrysts with alkali feldspar and quartz in a crystal-poor pantellerite implies temperatures below 700 °C and melt water contents higher than 4 wt.%, possibly up to 5–6 wt.%. Pantellerites have lower liquidus temperatures than associated comendites, which supports a parent–daughter relationship between the two magma types. The melts produced in the experiments extend the compositional trend displayed by the natural rock series, and reproduce some extreme compositions occasionally observed in alkaline volcanic series, with FeO contents above 12 wt.% and Na₂O contents approaching 10 wt.%. Pantellerites are therefore the true near-minimum melt compositions of alkaline oversaturated magma series.

Keywords: Peralkaline; Rhyolites; Pantellerite; Kenya; Volatiles

1. Introduction

Pantelleritic magmas are strongly peralkaline rhyolites having a $(\text{Na}_2\text{O} + \text{K}_2\text{O}) / \text{Al}_2\text{O}_3$ mol ratio (or agpaitic index, hereafter referred to as AI) higher than 1.4. Together with comendites, the less strongly peralkaline rhyolite variety ($1 < \text{AI} < 1.4$), pantellerites usually occur in extensional tectonic settings, from oceanic islands (Ascension Island) to continental rift zones (Ethiopia, Kenya) and back arc settings (Mayor Island, New Zealand), generally in close spatial and temporal association with transitional basalts. Recent analytical studies have emphasised that peralkaline rhyolites, once thought to be relatively anhydrous (Bailey and Macdonald, 1987), may have in fact up to several wt.% dissolved water under pre-eruptive conditions (Harris, 1986, Kovalenko et al., 1988, Lowenstern and Mahood, 1991, Webster et al., 1993, Wilding et al., 1993 and Barclay et al., 1996). Experimental work on comendites from the Olkaria volcanic complex, Kenya, has confirmed the water-rich nature of peralkaline rhyolites, as well as the usual corollary, the rather low temperatures of equilibration (Scaillet and Macdonald, 2001 and Scaillet and Macdonald, 2003). For the comendites studied, it was found that phase equilibrium constraints demand pre-eruptive temperatures to be lower than 750 °C, and possibly as low as 660 °C. Such low temperatures conflict with the widely held view that peralkaline silicic magmas are among the highest temperature rhyolites, which

would explain certain unusual volcanological features, such as their forming rheomorphic ignimbrites, by allowing extremely fluid behaviour during emplacement.

Although some progress has been made with respect to the volatile contents of peralkaline rhyolitic magmas, little is known, by comparison, of the temperature range over which such magmas evolve and/or are stored. This is largely due to the infrequency of coexisting oxide in such rocks, preventing application of the well-calibrated FeTi oxide thermobarometer (Andersen and Lindsley, 1988 and Ghiorso and Sack, 1991), a feature stressed more than 40 years ago by Carmichael (1962). There has been no detailed work on the phase equilibria of natural pantellerites, other than the exploratory studies of Bailey and coworkers (Bailey, 1974, Bailey et al., 1974 and Bailey and Cooper, 1978). As a consequence, although some insight is being gained through the application of alternative mineral equilibria (i.e. hedenbergite–fayalitic olivine and aenigmatite), the pre-eruption temperatures and water contents of pantellerites worldwide are not well established. This study aims to provide constraints on this issue, using the phase equilibrium approach that has proven useful in assessing the magmatic conditions of comendites (Scaillet and Macdonald, 2001).

We have determined the phase relationships of a pantellerite from the Eburru volcanic complex, which is located north of the Olkaria volcanic field in the Kenya rift on which our previous experimental efforts have been focused (Scaillet and Macdonald, 2001 and Scaillet and Macdonald, 2003). The study is a continuation of on-going work directed at understanding the mechanisms by which peralkaline silicic magmas are produced (e.g. Davies and Macdonald, 1987, Macdonald et al., 1987 and Heumann and Davies, 2002) and the conditions of magma storage in the crust. In our previous work, we systematically explored the effects of pressure, temperature, water content and fO_2 on phase equilibria and phase compositions, including the melts produced by extreme fractionation of marginally to moderately peralkaline rhyolites (comendites). Here we restrict our attention to the role played by variations in both melt water content and temperature in pantelleritic magmatism. We keep total pressure constant at 150 MPa and fO_2 at around NNO-2 (fO_2 is referenced to the Ni–NiO solid buffer, so that NNO-2 means an fO_2 2 log units below NNO at the chosen P and T), the values chosen for these two parameters being in agreement with those commonly accepted for peralkaline rhyolites, i.e. low pressure of magma storage (Mahood, 1984) and somewhat reducing conditions (see Scaillet and Macdonald, 2001), and in line with our previous findings on the P – fO_2 conditions of comendites (Scaillet and Macdonald, 2001). Note that in the present report, as in our previous studies (Scaillet and Macdonald, 2001 and Scaillet and Macdonald, 2003), the parameter which is fixed during the experiments is the hydrogen fugacity (fH_2). As explained below in detail, the fO_2 in our charges varied according to the water activity (variable melt water content). As a result, the phase equilibria reported here encompass a range of fO_2 . However, given the overall reducing conditions investigated (NNO-1 to NNO-4), the variation in fO_2 should be accompanied by minor changes in the Fe^{2+}/Fe^{3+} ratio (i.e. almost all iron in the melt is Fe^{2+}) so variations in experimental fO_2 are expected to have little effect on observed phase equilibria.

2. Geological background and selected sample

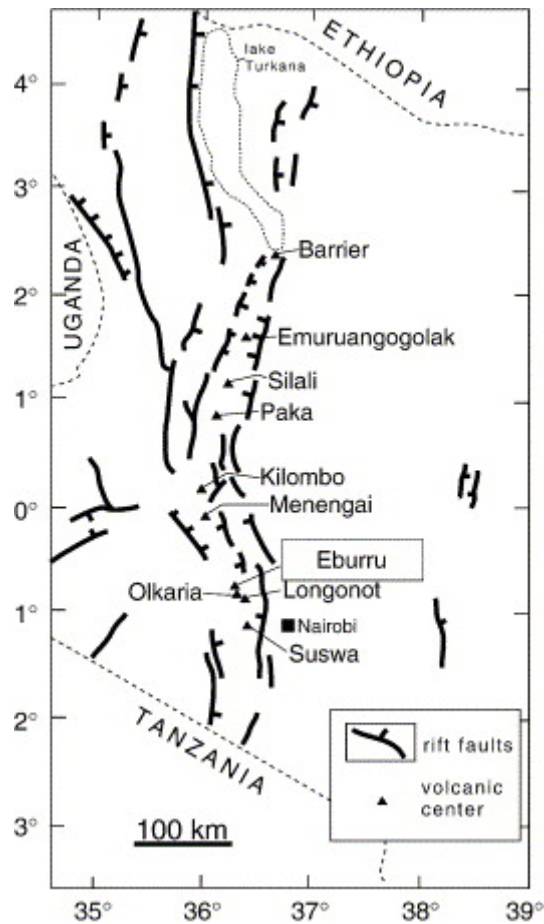


Fig. 1. Locality map of the Eburru and Olkaria rhyolitic complexes and of the nearby trachytic caldera volcanoes Longonot and Suswa. Also shown are the main Quaternary volcanic centres of the Kenya rift

The Eburru Volcanic Complex is located NW of Lake Naivasha (Fig. 1). Its southern boundary coincides with the northern boundary of the Olkaria field. Its age is not known in detail. According to Clarke et al. (1990), all units are thought to be < 0.45 Ma and the youngest eruptives probably no more than a few hundred years old. Ren et al. (2006-this issue), however, report that the complex was initiated prior to 1.2 Ma. Lithologically, Eburru is composed almost entirely of peralkaline trachytes and rhyolites of pantelleritic affinity. Two groups, members of which have been available for eruption throughout much of the history of the complex, have been distinguished by Clarke et al. (1990); the Eburru Trachyte and the Eburru Pantellerite. On an $\text{FeO}^{\square}-\text{Al}_2\text{O}_3$ plot, the Eburru Trachyte straddles the boundary between pantelleritic trachyte and pantellerite, whereas the Eburru Pantellerite plots entirely within the pantellerite field, parallel to, but displaced below, the Eburru Trachyte (see Fig. 7). This feature alone precludes a direct genetic connection between the trachytes and rhyolites.

Table 1. : Composition of the starting material

	Eburru-89^a	This study^b
SiO ₂	70.86	71.25
Al ₂ O ₃	8.48	8.03
FeO	5.02	4.23
Fe ₂ O ₃	2.40	3.00
FeO _{tot}	7.18	6.93
MgO	0.52	0.01
MnO	0.25	0.21
CaO	0.30	0.33
Na ₂ O	6.68	6.54
K ₂ O	4.50	4.60
TiO ₂	0.33	0.30
ZrO ₂	–	0.29
Cl	0.42	0.40
F	0.69	1.10
O = F, Cl	0.39	0.55
Total	100.06	99.44

^a From [Macdonald and Bailey \(1973, Table 3, no. 89\)](#) and [Bailey and Macdonald \(1975\)](#) for F and Cl analyses.

^b Average of 6 EMP analyses.

3. Experimental and analytical techniques

Powdered obsidian glass (20 mg) was loaded into Au capsules (1.5 cm length, wall thickness 0.2 mm and outer diameter 2.9 mm). Prior to the glass powder, distilled H₂O with or without CO₂ (added as silver oxalate) was inserted at the bottom of the capsule (2 mg H₂O + CO₂). Some charges were loaded with only the nominally dry glass powder (which in fact has 0.2 wt.% dissolved H₂O, as determined by several Karl Fischer titrations, plus 0.2 wt.% of adsorbed H₂O which is the water released at 120 °C during the titration). Once fully loaded, the capsules were welded shut using a graphite arc welder. The weight of the capsules was used as a monitor of volatile loss during welding. All charges reported in this work showed, upon welding or after being run at *P* and *T*, no weight changes in excess of 0.2 mg, which is within the analytical uncertainty of the balance. Experiments were performed either with cold seal pressure vessels (CSPV) or with an internally high-pressured vessel (IHPV), both fitted with an H₂ membrane to monitor the H₂ fugacity of the runs. Details of the procedure for the use of the vessels and accompanying furnaces are given in [Scaillet et al. \(1992\)](#) and [Schmidt et al. \(1995\)](#). Uncertainties in temperature reported in this work are estimated to be ± 8 °C for CSPV and ± 5 °C for IHPV.

The pressurizing medium was always an Ar–H₂ mixture, obtained either via sequential loading of H₂ and then Ar at room temperature (IHPV) or via H₂ supply through the H₂-membrane at P and T (CSPV). The uncertainty in total pressure is estimated to be ± 3 MPa, whilst that on H₂ fugacity is 0.01 MPa. Runs performed within CSPV involved 1 to 3 capsules, whilst those in IHPV had 6 capsules. Run durations varied according to temperature between 1193 h (650 °C) and 197 h (800 °C). Runs were terminated by either switching off the power supply and letting the vessel cool down at its own pace (IHPV), or by removing the vessel from the furnace (CSPV). Both quenches resulted in cooling rates on the order of 50–100 °C/min, with the IHPV imposing slightly faster cooling rates than the CSPV. The pressure drop during quench until the glass transition temperature (c. 300–400 °C) was reached was c. 15 MPa for the CSPV and 30 MPa for the IHPV.

Upon completion of the run, capsules were weighed to check for leaks and opened. During opening, evidence for excess fluid was looked for using two criteria: outbursts of water drops and hissing, especially for CO₂-bearing charges. Charges which were loaded with only water at amounts below H₂O saturation (as inferred from H₂O saturated runs, see below) did not show the above features of fluid saturation, as expected. For all capsules, a smell of H₂S was noted, indicating the presence of reduced sulphur in the charges. Polished probe mounts were then prepared using approximately half the run products, the remainder being used for direct optical investigation or additional analyses, if deemed necessary.

Run products were characterised by scanning electron microscopy (SEM) and electron microprobe analysis (EMPA). Analytical conditions for EMPA were accelerating voltage 15 kV, sample current 6 nA, counting time 10 s on peak for all elements and a focused beam for minerals. For glasses, the beam was defocused to 5 μm . Na and K were analysed first and a ZAF correction procedure was applied. A series of hydrous glass standards with known water content, made from the Eburru obsidian, was used to determine the empirical correction factor to be applied to the sodium content of the experimental glasses. This series was prepared with variable H₂O contents in Au capsules and annealed at 200 MPa and 850 °C for a week in a CSPV and drop quenched. Under the above analytical conditions, empirical correction factors varied from 1.05 for melt water contents lower than 2 wt.%, to 1.35 for melt water contents of 6–7 wt.%. Apart from Na, no significant variation in the abundance of any element was noted with variations in H₂O_{melt}.

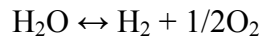
Melt H₂O contents were determined using the by-difference method (see [Scaillet and Macdonald, 2001](#)), and via mass balance constraints, assuming perfectly incompatible behaviour of H₂O (except for CO₂-bearing charges, see below). In the latter case, the amount of residual melt was determined by mass balance, using phase compositions as determined by EMPA. In the mass balance approach, it was assumed that the bulk water content of the charge consisted of the H₂O loaded to the capsule, plus that already dissolved or adsorbed in the natural glass (0.4 wt.% H₂O). We further considered the reduction of Fe₂O₃ to FeO as a possible source of water. Given that the starting glass contains 4.2 wt.% FeO, as determined by wet chemistry, this implies that the 3 wt.% of Fe₂O₃ originally present in the glass can yield 0.37 wt.% additional water, should the reduction of Fe₂O₃ proceed to completion. The H₂O contents of melts determined by both approaches generally agree to within 0.5 wt.% of each other, except for the most crystal-rich charges, for which the by-difference approach yields melt H₂O contents 1–2 wt.% higher than those via mass balance. SEM observations reveal that these crystal-rich charges are often packed with μm -sized bubbles of inferred quench origin ([Scaillet and Macdonald, 2001](#) and [Scaillet and Macdonald, 2003](#)). As a result, broad beam EMPA analyses end up producing exceedingly low totals and thus spuriously

high melt H₂O contents, much higher than permitted by the total amount of H₂O of the system. Melt water contents of H₂O-saturated charges obtained at 800 °C and 750 °C and determined by-difference are 5.8 and 6.1 wt.%, respectively, in reasonable agreement with previous estimates for similar compositions ([Scaillet and Macdonald, 2001](#)). In contrast, H₂O-saturated charges synthesised at 700 and 650 °C display higher H₂O contents (7–8 wt.%). Although we cannot rule out that such an increase is due to a temperature effect, the rate of H₂O increase with falling temperature is much higher than in metaluminous rhyolites ([Johannes and Holtz, 1996](#)). We thus suspect that the H₂O content of low temperature charges is affected by quench processes, as in crystal-rich charges. For simplicity, however, we have assumed a constant melt water content of 6 wt.% at all temperatures, based on the by-difference determination at 750 °C and 800 °C of H₂O-saturated charges. The water contents of H₂O-undersaturated charges with no CO₂ used in this study are those obtained by mass balance. For CO₂-bearing charges, we assumed that the following relationships holds:

$$\text{wt.\% H}_2\text{O} = \text{H}_2\text{O}_{\text{sat}} \times \text{XH}_2\text{O}_{\text{in}}$$

where XH₂O_{in} is the mole fraction of H₂O of the starting fluid composition loaded to the charge. This relationship is equivalent to assuming ideal behaviour of water in the CO₂–H₂O fluid mixture. The method yields maximum estimates for H₂O_{melt}. Overall, melt water contents determined in this study are in the range 0.8–6 wt.%.

For each charge, the determination of prevailing *f*O₂ used the water dissociation reaction:



with thermodynamic data from [Robie et al. \(1979\)](#) for the dissociation constant, *K_w*. For H₂O-saturated charges, in addition to *K_w* we use the *f*H₂ as given by membrane reading and *f*H₂O of pure H₂O at the *P* and *T* of the experiments, as given by [Burnham et al. \(1969\)](#). For H₂O-undersaturated charges, a complete *f*O₂ calculation requires that *f*H₂O be evaluated. Given that the activity–composition relationships of H₂O in strongly peralkaline silicic melts are unknown, we have adopted the following procedure. We have assumed that the solubility of H₂O in pantellerite obeys a power law of the form:

$$f\text{H}_2\text{O} = a(\text{H}_2\text{O}_{\text{melt}})^b$$

where *a* and *b* are fitted parameters and H₂O_{melt} is the wt.% H₂O dissolved in the melt. We fitted this function so that it yields a H₂O solubility of 6 wt.% at 150 MPa and 0.1 wt.% at atmospheric pressure. The resulting values for *a* and *b* coefficients were 52.7386 and 1.7221, respectively. Next, using this empirical equation, *f*H₂O can be calculated for any charge whose H₂O content is known. Thereafter, the corresponding *f*O₂ can be calculated using the water dissociation reaction and *f*H₂, as explained above. The calculated *f*O₂, given as ΔNNO values, ranges from –0.69 at H₂O saturation to –4.38 for the driest charges (< 1 wt.% H₂O_{melt}), with an average of –2.2 ([Table 2](#)). For convenience, we refer in the following to the redox conditions of this experimental study as being at an average *f*O₂ of NNO-2. For H₂O-saturated charges, the uncertainty in *f*O₂ depends primarily on the uncertainty on the H₂ fugacity and is estimated to be better than 0.1 log unit. For H₂O-undersaturated charges, the uncertainty in *f*O₂ is heavily dependent on that in *f*H₂O, which is not well known. However, charges whose H₂O_{melt} exceeds 3 wt.% differ in *f*O₂ from the H₂O-saturated case by less than

0.5 log unit ([Table 2](#)). It is for the driest charges (≤ 1 wt.% $\text{H}_2\text{O}_{\text{melt}}$) that small variations in $\text{H}_2\text{O}_{\text{melt}}$ produce quite significant changes in $f\text{O}_2$ and for these charges we estimate the uncertainty in $f\text{O}_2$ to be of the order of 1 log unit.

4. Results

4.1. General

The experimental conditions and run products of 26 successful experiments are listed in [Table 2](#). Also given are the phase proportions obtained from mass balance calculations ([Albarède, 1995](#)). Calculations were performed using SiO_2 , Al_2O_3 , FeO^\square , MgO , MnO , CaO , Na_2O , K_2O and TiO_2 and take into account the propagation of the average analytical uncertainty of probe analyses. The data needed to determine phase proportions include the compositions of crystal and melt phases, as determined by EMPA, and the starting bulk rock composition ([Table 1](#)). The calculation is performed using matrix procedures, as explained in detail in [Albarède \(1995\)](#). Apart from two charges (#1 and #9), the sum of residuals (r^2) is below 1, indicating that the main phases have been identified. However, in some charges the incorporation of an additional Na-bearing phase significantly decreases the residuals (charges with NaCl in [Table 2](#)). This could imply either an underestimation of the Na correction factor during probe analysis of the residual melts or the possibility that Na has been partitioned into an unidentified phase which could be a fluid, a highly saline immiscible melt or a solid (NaCl). For some charges at H_2O saturation (#1 and #5), the second hypothesis seems plausible and is corroborated by the fact that Cl contents are lower than those calculated assuming perfectly incompatible behaviour, suggesting effective transport of a NaCl compound into a coexisting fluid or immiscible saline melt. However, above-liquidus CO_2 -bearing charges (#6, #7 and #8) have Cl contents similar to that of the starting material (c. 0.39 wt.%). Overall, the fact that Cl in the residual melts shows broadly incompatible behaviour (see below) suggests that, despite the care taken in glass analysis, some may have had their Na_2O content slightly underestimated.

Table 2. : Experimental conditions and run products

Run	P_{tot} (bar)	T (°C)	$f\text{H}_2$ (bar)	$f\text{H}_2\text{O}^{\text{a}}$ (bar)	$\log f\text{O}_2$	$\Delta\text{NNO}^{\text{b}}$	$\text{XH}_2\text{O}_{\text{in}}^{\text{c}}$	$\text{H}_2\text{O}^{\text{d}}$ (wt.%)	CO_2^{e} (wt.%)	$\text{H}_2\text{O}_{\text{melt}}^{\text{f}}$	Duration (h)	r^2	Phases present and abundances (wt.%)
1	1525	650	23	961	-19.03	-1.46	1	9.3	-	6.00	1193	1.02	gl (92.9), vp, amph (5.9), nacl (1.2)
2	1525	650	23	903	-19.08	-1.51	0.94	7.8	1.2	5.64	1193	0.51	gl (87.5), vp, amph (6.7), qz (0.1), af (3.8), nacl (2.0)
3	1525	650	23	817	-19.17	-1.60	0.85	6.2	2.6	5.10	1193	-	gl, vp, amph, aeg, qz, af
4	1525	650	23	740	-19.25	-1.68	0.77	5.3	3.9	4.62	1193	-	gl, vp, amph, aeg, qz, af
5	1510	700	19	1056	-17.33	-1.15	1	10	-	6.00	807	-	gl, vp
6	1510	700	19	875	-17.50	-1.31	0.91	8.2	1.6	5.46	807	-	gl, vp
7	1510	700	19	817	-17.56	-1.37	0.85	6.5	2.7	5.10	807	-	gl, vp
8	1510	700	19	692	-17.70	-1.52	0.72	4.7	4.5	4.32	807	-	gl, vp, amph (tr)
9	1510	700	19	139	-19.10	-2.91	-	-	-	1.76	807	1.85	gl (43.9), vp, amph (7.6), qz (21.3), af (27.2)
10	1584	751	33	98	-18.56	-3.64	-	-	-	1.43	789	0.04	gl (53.8), qz (17.2), af (29)
11	1526	775	33	74	-18.22	-3.86	-	-	-	1.22	240	0.11	gl (63.1), qz (14.7), af (22.2)
12	1526	775	33	245	-17.18	-2.82	-	1.6	-	2.44	240	0.36	gl (97.2), qz (1.8), af (1)
13	1546	801	34	51	-17.97	-4.18	-	-	-	0.99	240	0.07	gl (78.1), qz (9.3), af (12.6)
14	1546	801	34	233	-16.65	-2.87	-	1.6	-	2.37	240	-	gl
15b	1541	825	37	44	-17.65	-4.38	-	-	-	0.90	197	0.23	gl (85.7), qz (6.5), af (7.8)
15	1525	750	13	74	-18.01	-3.07	-	-	-	1.22	286	0.09	gl (63.0), qz (13.4), af (23.6)
16	1525	750	13	247	-16.97	-2.03	-	1.3	-	2.45	286	0.46	gl (84.4), qz (6.3), af (9.3)

Run	P_{tot} (bar)	T (°C)	$f\text{H}_2$ (bar)	$f\text{H}_2\text{O}^{\text{a}}$ (bar)	$\log f\text{O}_2$	$\Delta\text{NNO}^{\text{b}}$	$\text{XH}_2\text{O}_{\text{in}}^{\text{c}}$	$\text{H}_2\text{O}^{\text{d}}$ (wt.%)	CO_2^{e} (wt.%)	$\text{H}_2\text{O}_{\text{melt}}^{\text{f}}$	Duration (h)	r^2	Phases present and abundances (wt.%)
17	1525	750	13	301	-16.79	-1.86	-	1.9	-	2.75	286	0.02	gl (97.1), qz (2.1), nacl (0.8)
18	1525	750	13	403	-16.54	-1.60	-	2.5	-	3.26	286	-	gl, qz (tr), nacl
19	1525	750	13	695	-16.07	-1.13	-	3.7	-	4.47	286	-	gl
20	1525	750	13	1154	-15.63	-0.69	-	6.2	-	6.00	286	-	gl, vp
21	1550	800	13	37	-17.45	-3.65	-	-	-	0.81	208	0.07	gl (95.2), qz (3.4), af (1.4)
22	1550	800	13	59	-17.04	-3.23	-	0.3	-	1.07	208	-	gl, qz (tr), nacl
23	1550	800	13	213	-15.92	-2.12	-	1.5	-	2.25	208	-	gl, qz (tr), nacl
24	1550	800	13	344	-15.50	-1.70	-	2.2	-	2.97	208	-	gl
25	1550	800	13	695	-14.89	-1.09	-	3.7	-	4.47	208	-	gl
26	1550	800	13	545	-15.10	-1.30	-	6.2	-	3.90	208	-	gl, vp

tr: phase present in trace amounts. gl: glass; vp: vapour; amph: amphibole; af: alkali feldspar; qz: quartz; aeg: aegerine, nacl: sodium chloride.
^a $f\text{H}_2\text{O}$ calculated
^b ΔNNO : $\log f\text{O}_2$ relative to the value of the Ni-NiO buffer.
^c Initial mole fraction of H_2O in the fluid phase for CO_2 -bearing capsules.
^d Weight percent of H_2O loaded to the charge.
^e Weight percent of CO_2 loaded to the charge.
^f Amount of dissolved H_2O in the glass.

The crystalline phases identified are sodic amphibole, aegirine, alkali feldspar and quartz. Fluorite was formally detected in one charge (#4) and, although no effort was made to systematically look for this mineral, we believe that its stability field is restricted to below 750 °C at all water contents, since no bright mineral in SEM images was seen in any of the charges run at or above 750 °C. Glass was present in all charges, but could not be probed in charges #3 and #4, owing to their high crystallinity. All CO₂-bearing or H₂O saturated charges contained large bubbles (> 20 µm) indicative of fluid saturation. Charges that were supposed not to be saturated in fluid also contained small bubbles (< 10 µm), whose proportion is 1–3% by volume, regardless of charge crystallinity. The lack of variation with crystallinity of this population of bubbles suggests that they probably represent remnants of air bubbles originally trapped in the glass powder. Aegirine was positively identified only at 650 °C in the two driest charges (#3 and #4). No oxide was detected.

4.2. Phase relationships

The phase relationships are shown on [Fig. 2](#). At H₂O saturation, amphibole is the liquidus phase, crystallising slightly above 650 °C. It has not been found at temperatures higher than 700 °C and its stability field is drawn flat, although crystallisation of amphibole under very dry conditions (< 1 wt.% H₂O in melt) at temperatures higher than 700 °C cannot be excluded. Similarly, aegirine stability is restricted to below 700 °C but the stability field shown in [Fig. 2](#) is only approximate. In particular, the extension or termination of this phase boundary in the high temperature and dry ranges are not constrained by the current data set. Below 700 °C, quartz and alkali feldspar co-crystallise next to amphibole. Above 700 °C, they both precede amphibole, quartz appearing slightly earlier than the alkali feldspar. At 800 °C, quartz and alkali feldspar crystallise at melt H₂O contents of 2 and 1 wt.%, respectively. Extrapolation of the quartz boundary up-temperature suggests that the dry liquidus of the Eburru pantellerite is at, or above, 850 °C at 150 MPa. Our data indicate that crystal-poor (i.e. near liquidus) pantelleritic magmas saturated in tectosilicates and amphibole require conditions of both low temperature (650 °C) and near H₂O saturation (> 5 wt.% H₂O).

The phase relationships in the Eburru pantellerite can be compared to those of Olkaria comendite SMN 29 studied by [Scaillet and Macdonald \(2001\)](#) at 150 MPa and under slightly more oxidizing conditions (NNO-1.6). The boundaries of the main phases in both rhyolites are shown superimposed in [Fig. 3](#). It is apparent that whilst the amphibole stability field occupies approximately the same position in both compositions, the liquidus curves of the tectosilicates are depressed by 30–40 °C in the Eburru pantellerite compared to the Olkaria comendite. This downward displacement can be due to the combined effects of the more strongly peralkaline character of, and the more reducing conditions applied to, the Eburru rock. However, given the rather modest difference in fO_2 between the two studies (c. 0.5 log unit), we suggest that the first effect is dominant. The fact that liquidus temperatures are lower in pantellerite than in comendite magma is in agreement with the demonstration that extensive crystallisation of comendite can produce pantellerite ([Scaillet and Macdonald, 2003](#)). The small difference in melt H₂O content at saturation is not considered to be representative, given the error associated with the method of H₂O determination in the experimental glasses.

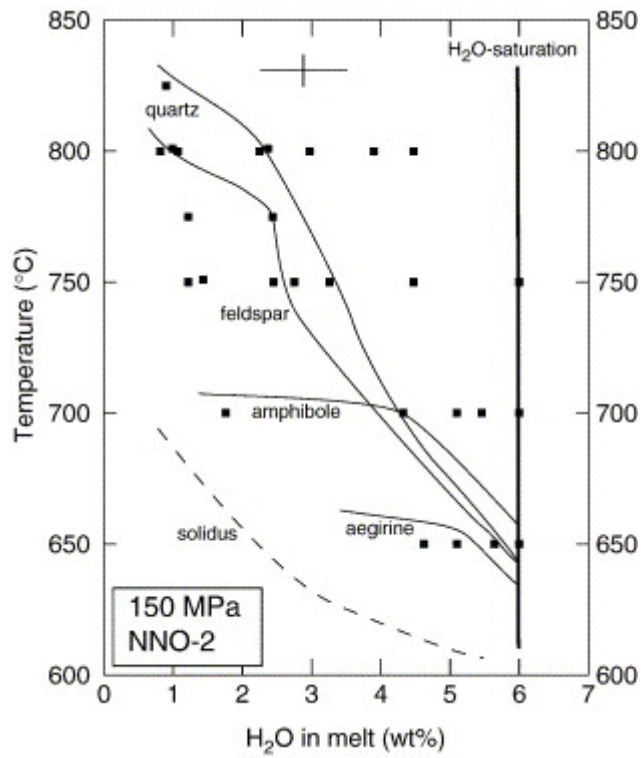


Fig. 2. Phase relationships of Eburru pantellerite at 150 MPa and NNO-2.

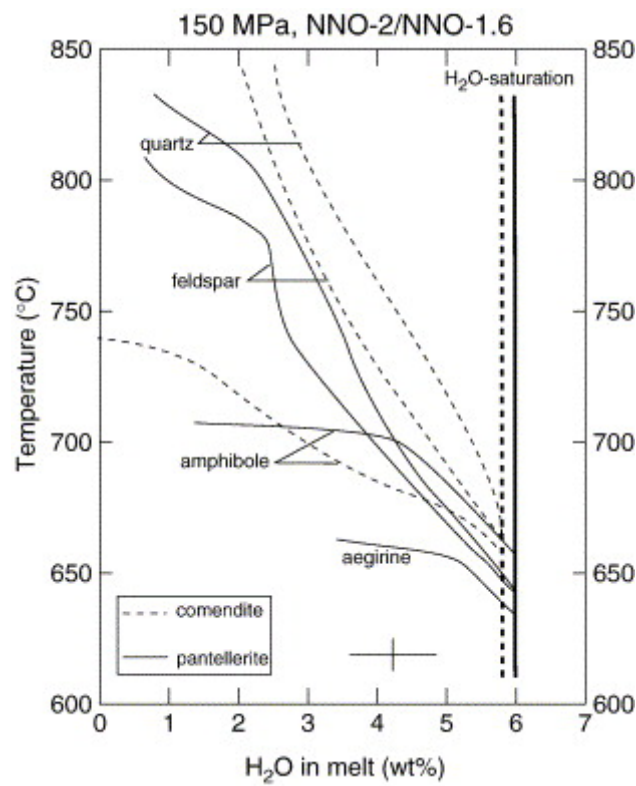


Fig. 3. Comparison between the phase relationships of Eburru pantellerite (continuous lines) and those of Olkaria comendite (dashed lines), the latter taken from [Scaillet and Macdonald \(2001\)](#), rock composition SMN 49. Note the downward displacement of tectosilicate curves in Eburru pantellerite relative to Olkaria comendite

4.3. Phase abundances

Mass balance calculations indicate that quartz and alkali feldspar are the dominant crystallising phases under all conditions except at near H₂O saturation and low temperature, at around 650 °C, where amphibole is the first phase to precipitate, reaching mass proportions of 6–7 wt.%. The most highly crystallised charge (#9, [Table 2](#)) has 44 wt.% melt and contains 7.6 wt.% amphibole, in addition to quartz (21.3 wt.%) and alkali feldspar (27.2 wt.%). The fact that the proportion of amphibole does not increase significantly with crystallinity suggests that it probably does not crystallise in much higher abundance at lower melt fractions. Such a lack of increase might be due to a limitation of the bulk iron content available in the residual melt (and perhaps also to that of Na₂O in excess of Al₂O₃), since amphibole crystallising in our experiments has FeO_{tot} in excess of 30 wt.% ([Table 3](#)). In contrast, the non-increase in amphibole proportion cannot be attributed to a decrease in melt water content since as H₂O_{melt} decreases, the water structurally bound in amphibole is exchanged with fluorine (i.e. the amphibole gets more F-rich, see next section) which helps stabilising sodic amphibole even in H₂O-poor magmas. Such a behaviour has been also noted for Olkaria comendites ([Scaillet and Macdonald, 2001](#) and [Scaillet and Macdonald, 2003](#)). Thus, contrarily to arc magmas in which amphibole occurrence is generally taken as evidence of water-rich conditions, such a correlation is not always true for peralkaline rhyolites. Charges that crystallised aegirine could not be probed for their residual glasses, so the proportion of sodic pyroxene remains unknown. [Fig. 4](#) shows the variation in the proportions of quartz and alkali feldspar with melt fraction. Both proportions increase almost linearly, although with different slopes, with decreasing melt fraction. Extrapolation of the quartz trend to the zero melt fraction intercept gives a quartz proportion at the solidus of 37 wt.%, which is similar to granite. Similar extrapolation of alkali feldspar yields a proportion of 57 wt.%, the remaining 6 wt.% (= 100 – 37 – 57) representing minor phases such as amphibole and aegirine. Amphibole precipitation produces a significant decrease of feldspar abundance relative to the trend drawn by amphibole-free charges ([Fig. 4](#)), which suggests that amphibole crystallises at the expense of feldspar-forming complexes in the melt.

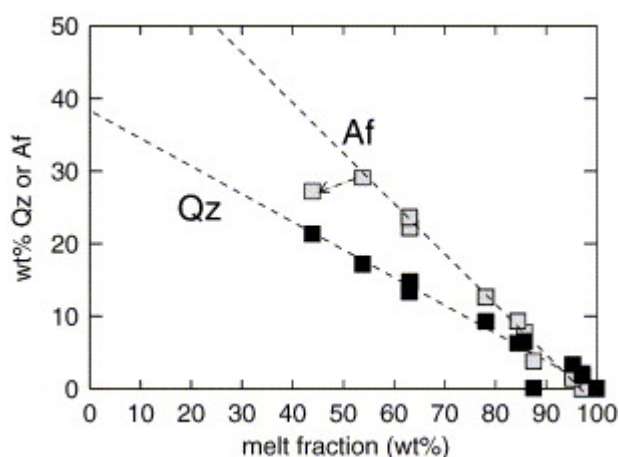


Fig. 4. Variations in quartz and alkali feldspar weight proportions with crystallisation. The dashed arrow shows the effect of amphibole crystallisation on the proportions of alkali feldspar.

Table 3. : Representative compositions of amphibole and aegirine in Eburru experiments

	Ebu-1 amph		Ebu-2 amph		Ebu-3 amph	Ebu-9 amph		Ebu-3 aeg
	<i>N</i> = 4	S.D.	<i>N</i> = 3	S.D.	<i>N</i> = 1	<i>N</i> = 4	S.D.	<i>N</i> = 1
SiO ₂	48.79	0.92	47.59	0.44	50.37	49.28	0.41	50.93
TiO ₂	1.05	0.17	1.24	0.12	0.79	1.72	0.22	1.08
Al ₂ O ₃	0.85	0.29	0.59	0.09	1.36	0.56	0.23	0.46
FeO	33.44	2.18	33.17	0.60	29.45	32.66	0.87	27.86
MgO	0.13	0.03	0.11	0.04	0.16	0.14	0.06	0.05
MnO	1.03	0.09	0.87	0.07	0.87	0.78	0.23	0.18
CaO	0.93	0.16	0.50	0.13	0.45	0.93	0.09	0.84
Na ₂ O	7.55	0.34	7.63	0.48	9.32	7.89	0.11	12.40
K ₂ O	1.61	0.04	1.75	0.15	1.54	1.64	0.03	0.10
ZrO ₂	0.10	0.14	0.24	0.18	0.40	0.17	0.13	0.00
Cl	0.03	0.02	0.02	0.01	0.03	0.02	0.03	0.01
F	1.60	0.50	1.88	0.41	1.92	2.53	0.42	0.21
Total	96.34		94.56		95.45	97.08	0.66	94.12
Si	7.93		7.89		8.07	7.87		2.02
Ti	0.13		0.15		0.1	0.21		0.02
Al	0.16		0.12		0.26	0.11		0.04
Fe ²⁺	4.54		4.6		3.95	4.36		0.14
Fe ³⁺	–		–		–	–		0.78
Mg	0.03		0.03		0.04	0.03		0.00
Mn	0.14		0.12		0.12	0.11		0.01
Ca	0.16		0.09		0.08	0.16		0.04
Na	2.38		2.45		2.9	2.44		0.95
K	0.33		0.37		0.31	0.33		

Structural formulas are calculated based on 23 O for amphibole and 6 O for pyroxene. For amphibole, no attempt has been made to determine its Fe²⁺/Fe³⁺ ratio and the Fe is given as Fe²⁺. For clinopyroxene, Fe³⁺ is calculated assuming ideal pyroxene stoichiometry and using charge balance constraints. Totals given are corrected for F. The calculations do not incorporate Zr since its concentration is not known with great accuracy given the applied analytical conditions and the overall low concentrations of this element in experimental minerals. The pyroxene was assumed free of both F and K₂O, whose presence in the analysis is inferred to reflect glass contamination.

4.4. Phase compositions

Alkali feldspar, amphibole, aegirine and glass were analysed (Table 3 and Table 4). Alkali feldspar was rather difficult to analyse, due to its small size and poor contrast with residual glass on BSE images, but the few acceptable analyses obtained reveal a constant composition at \sim Or₄₀Ab₆₀, with almost no detectable calcium. This composition is closely comparable to the phenocryst compositions in the natural rocks of similar AI (Ren et al., 2006-this issue).

Table 4. : Compositions of experimental glasses

Sample	SiO ₂	TiO ₂	Al ₂ O	FeO	MgO	MnO	CaO	Na ₂ O	K ₂ O	ZrO ₂	Cl	F	Original total
Ebu-1 (n = 6)	74.58	0.24	8.53	5.16	0.01	0.16	0.07	5.28	4.38	0.51	0.18	0.91	89.29
	0.36	0.12	0.08	0.35	0.01	0.08	0.06	0.07	0.10	0.10	0.04	0.08	
Ebu-2 (n = 6)	73.98	0.21	8.70	5.26	0.01	0.22	0.09	5.32	4.32	0.45	0.32	1.12	89.54
	0.15	0.07	0.17	0.12	0.01	0.07	0.07	0.14	0.16	0.19	0.06	0.57	
Ebu-4	71.63	0.32	9.11	6.15	0.00	0.34	0.01	5.49	4.45	0.91	0.42	1.19	86.24
Ebu-5 (n = 6)	72.00	0.31	8.09	6.96	0.01	0.17	0.25	5.79	4.34	0.48	0.28	1.32	91.39
	0.50	0.07	0.13	0.16	0.02	0.13	0.05	0.40	0.09	0.19	0.03	0.58	
Ebu-6 (n = 7)	72.08	0.27	8.11	6.81	0.01	0.22	0.25	5.81	4.44	0.43	0.34	1.23	92.15
	0.44	0.07	0.06	0.32	0.01	0.11	0.05	0.30	0.18	0.20	0.04	0.64	
Ebu-7 (n = 7)	72.02	0.28	8.02	6.92	0.02	0.20	0.22	5.86	4.31	0.45	0.37	1.34	92.93
	0.48	0.03	0.13	0.22	0.01	0.05	0.08	0.37	0.11	0.12	0.04	0.53	
Ebu-8 (n = 6)	71.49	0.33	8.02	6.87	0.00	0.23	0.21	6.01	4.53	0.44	0.38	1.48	92.52
	0.61	0.06	0.10	0.25	0.01	0.04	0.06	0.26	0.12	0.23	0.04	0.44	
Ebu-9 (n = 5)	63.56	0.49	6.68	9.70	0.01	0.35	0.25	9.82	4.77	0.91	0.87	2.59	96.06

Sample	SiO ₂	TiO ₂	Al ₂ O	FeO	MgO	MnO	CaO	Na ₂ O	K ₂ O	ZrO ₂	Cl	F	Original total
	3.73	0.16	1.98	1.81	0.01	0.08	0.09	0.97	0.53	0.16	0.24	0.87	
Ebu-10 (n = 4)	64.25	0.59	5.22	12.27	0.02	0.42	0.36	8.79	4.48	0.86	0.82	1.92	96.22
	0.43	0.06	0.60	1.12	0.02	0.12	0.08	0.17	0.46	0.30	0.05	0.39	
Ebu-11 (n = 6)	65.82	0.51	6.26	10.56	0.03	0.40	0.31	7.96	4.74	0.66	0.68	2.07	95.92
	0.34	0.11	0.38	0.46	0.02	0.10	0.04	0.10	0.18	0.37	0.07	0.29	
Ebu-12 (n = 5)	71.10	0.25	8.17	7.33	0.01	0.27	0.16	6.26	4.66	0.40	0.43	0.96	94.97
	0.21	0.06	0.14	0.11	0.01	0.11	0.03	0.15	0.08	0.17	0.04	0.40	
Ebu-13 (n = 6)	68.55	0.39	7.24	8.63	0.02	0.27	0.27	7.40	4.77	0.51	0.58	1.39	95.53
	0.40	0.06	0.31	0.31	0.01	0.07	0.06	0.17	0.13	0.14	0.03	0.47	
Ebu-14 (n = 6)	72.07	0.30	8.18	6.70	0.00	0.19	0.21	6.43	4.44	0.43	0.33	0.71	96.20
	0.45	0.06	0.08	0.18	0.00	0.10	0.03	0.31	0.06	0.19	0.05	0.46	
Ebu-15b (n = 7)	69.31	0.39	7.55	7.97	0.02	0.30	0.19	6.73	4.83	0.57	0.47	1.67	97.77
	0.63	0.06	0.14	0.24	0.02	0.09	0.07	0.15	0.09	0.19	0.06	0.50	
Ebu-15 (n = 7)	66.18	0.45	5.94	10.63	0.02	0.33	0.40	7.67	4.49	0.81	0.72	2.35	97.35
	0.30	0.11	0.10	0.14	0.02	0.09	0.06	0.35	0.13	0.34	0.04	0.67	
Ebu-16 (n = 4)	69.72	0.35	7.66	8.38	0.02	0.26	0.28	6.41	4.59	0.63	0.53	1.16	96.07
	0.42	0.04	0.14	0.26	0.01	0.06	0.04	0.12	0.12	0.15	0.04	0.32	
Ebu-17 (n = 8)	71.25	0.27	8.35	7.17	0.01	0.19	0.26	5.87	4.66	0.33	0.45	1.17	95.98
	0.29	0.06	0.27	0.25	0.02	0.10	0.05	0.10	0.19	0.15	0.05	0.60	
Ebu-18	72.3	0.23	8.10	6.95	0.03	0.20	0.25	5.73	4.5	0.49	0.3	0.7	94.81

Sample	SiO ₂	TiO ₂	Al ₂ O	FeO	MgO	MnO	CaO	Na ₂ O	K ₂ O	ZrO ₂	Cl	F	Original total
(n = 6)	4								2		7	6	
	0.33	0.08	0.13	0.18	0.02	0.13	0.03	0.13	0.09	0.29	0.04	0.79	
Ebu-19 (n = 7)	71.88	0.29	7.97	6.66	0.02	0.23	0.24	6.07	4.41	0.68	0.40	1.14	94.68
	0.37	0.06	0.15	0.22	0.02	0.08	0.04	0.35	0.09	0.13	0.05	0.78	
Ebu-20 (n = 6)	72.19	0.27	8.04	6.99	0.02	0.22	0.23	6.10	4.35	0.53	0.36	0.70	92.01
	0.70	0.06	0.15	0.32	0.01	0.10	0.04	0.38	0.16	0.25	0.04	0.62	
Ebu-21 (n = 6)	69.98	0.37	7.96	7.11	0.01	0.27	0.29	6.74	4.82	0.75	0.52	1.18	97.14
	0.38	0.08	0.17	0.26	0.02	0.09	0.06	0.13	0.15	0.26	0.05	0.44	
Ebu-22 (n = 6)	71.77	0.28	8.24	6.17	0.01	0.27	0.25	6.15	4.64	0.37	0.39	1.46	97.09
	1.82	0.06	0.31	0.36	0.01	0.10	0.08	0.19	0.23	0.32	0.04	0.47	
Ebu-23 (n = 6)	72.23	0.25	8.20	6.58	0.02	0.14	0.20	6.01	4.52	0.39	0.39	1.08	96.03
	0.17	0.05	0.11	0.21	0.02	0.06	0.07	0.10	0.08	0.27	0.05	0.63	
Ebu-24 (n = 7)	73.18	0.28	8.19	5.77	0.01	0.15	0.24	5.96	4.46	0.48	0.37	0.93	96.02
	0.31	0.05	0.05	0.35	0.01	0.10	0.05	0.26	0.18	0.15	0.03	0.60	
Ebu-25 (n = 5)	71.89	0.30	8.02	6.85	0.01	0.12	0.19	6.13	4.53	0.60	0.38	0.99	94.44
	0.74	0.07	0.05	0.24	0.03	0.06	0.04	0.37	0.05	0.29	0.05	0.39	
Ebu-26 (n = 6)	72.19	0.23	8.07	6.62	0.00	0.12	0.22	6.31	4.53	0.49	0.40	0.81	94.50
	1.14	0.05	0.09	0.21	0.01	0.04	0.04	0.25	0.12	0.27	0.07	0.65	

The amphibole is always sodic and can be classified as arfvedsonite (Leake et al., 1997). It is Fe-rich, with MgO contents below 0.2 wt.%, lower than MnO (0.7–1 wt.%). Although it occurs over a limited T -H₂O range, the analyses reveal some distinct trends. At 650 °C,

amphibole crystallising near H₂O saturation is richer in Ca than those near the solidus (compare #1, #2 and #3, [Table 3](#)). Such a trend of decreasing CaO content with melt H₂O content at fixed temperature has also been documented in amphibole crystallising in the Olkaria comendites ([Scaillet and Macdonald, 2003](#)). Similarly, the amphibole is F-rich, with F contents increasing with decreasing melt fraction, mirroring the increase in F of the coexisting melt. The mineral/melt partition coefficient for F ranges from 1.8 near the liquidus (#1 and #2) to 1 at higher melt fractions (#9), similar to that at Olkaria ([Scaillet and Macdonald, 2003](#)). Despite the elevated Cl content of the coexisting melt, amphibole has almost no Cl, as in Olkaria rocks and experiments ([Macdonald et al., 1987](#) and [Scaillet and Macdonald, 2003](#)), but contains detectable amounts of Zr. Its TiO₂ content, in the range 0.8–1.7 wt.%, shows no clear variation with changing melt fraction.

The few analyses of clinopyroxene show it to be sodic aegirine of nearly pure end-member composition ([Table 3](#)). The low total is presumably due to its high Fe³⁺ content, which amounts to 84% of total iron when calculated with stoichiometry and charge balance constraints ([Table 3](#)).

The variation in glass composition ([Table 4](#)) with melt fraction is shown on [Fig. 5](#). K₂O remains remarkably constant over the crystallisation interval investigated. SiO₂ and Al₂O₃ show a pronounced decrease with crystallisation, whilst FeO[□], Na₂O, CaO, MnO, TiO₂ and ZrO₂ contents increase regularly. The trend of decreasing SiO₂ and concomitant increase in Na₂O are similar to those documented for Olkaria comendites ([Macdonald et al., 1987](#) and [Scaillet and Macdonald, 2003](#)). The most peralkaline melt produced in our experiments has AI = 3.4, an Al₂O₃ content of 5.2 wt.%, and a FeO[□] content > 12 wt.%, closely resembling analyses of melt inclusions and interstitial glasses in peralkaline rhyolites from the Azores ([Mungall and Martin, 1996](#)). It has also an elevated ZrO₂ content, close to 1 wt.%. The monotonous decreases or increases in element abundances are affected only by amphibole crystallisation. In particular, amphibole crystallisation at near-liquidus conditions, that is when only a minor proportion of quartz and alkali feldspar has crystallised, is able to drive the residual liquids toward compositions less peralkaline than the bulk starting material ([Fig. 5](#)), producing melts with AI values approaching the comendite–pantellerite divide line which, following [Macdonald \(1974\)](#), is set at AI = 1.4. This is due to the role that amphibole exerts on both Na₂O and Al₂O₃ melt contents, the latter oxide being affected whether near-liquidus or crystal-rich conditions prevail, whilst the effect on Na₂O is significant only under crystal-poor conditions ([Fig. 5](#)). Crystallisation of sodic amphibole buffers melt peralkalinity, preventing it from reaching the Na contents observed when only tectosilicate phases precipitate, as shown for the Olkaria comendites ([Scaillet and Macdonald, 2003](#)). Amphibole crystallisation in peralkaline rhyolites counteracts to some extent the so-called orthoclase effect on melt peralkalinity proposed by [Bailey and Schairer \(1964\)](#) and [Bailey and Macdonald \(1969\)](#).

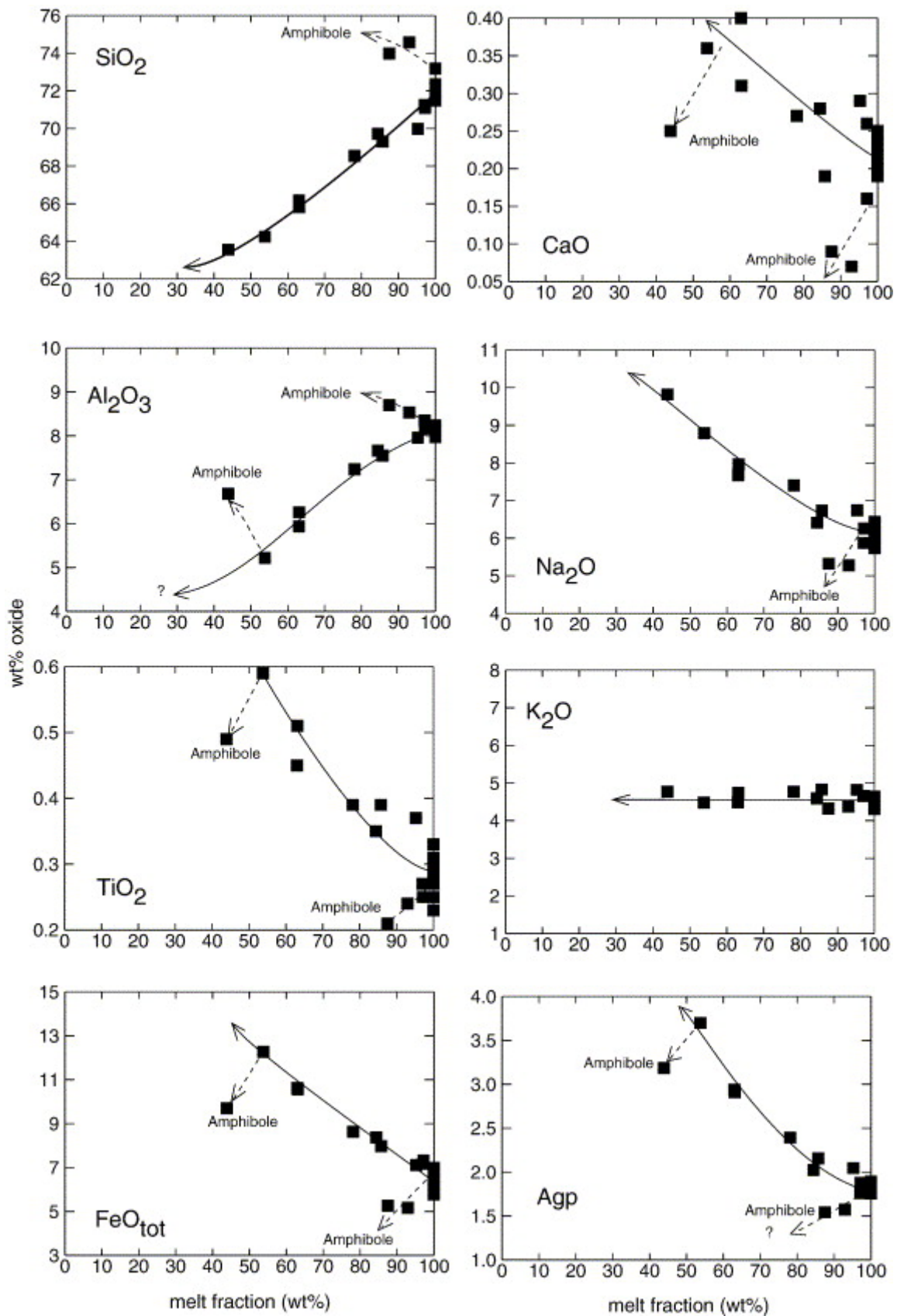


Fig. 5. Variation in the composition of the experimental glasses with crystallisation. The dashed lines show the effect of amphibole crystallisation on melt composition, either near the liquidus or under crystal-rich conditions.

Cl and F abundances increase with crystallisation (Fig. 6). Although some scatter is apparent, in the most crystal-rich charges analysed Cl displays contents (0.87 wt.%) similar to those predicted from incompatible behaviour (0.89 wt.%) (Fig. 6). F follows broadly the same pattern, except when amphibole crystallises, which somewhat dampens the increase in F. However, the effect remains limited (2.51 wt.% observed vs. 2.59 wt.% calculated in charge #9, Table 2). The scatter in F contents at near-liquidus conditions, as compared to Cl, may reflect some heterogeneity in F distribution in the starting glass

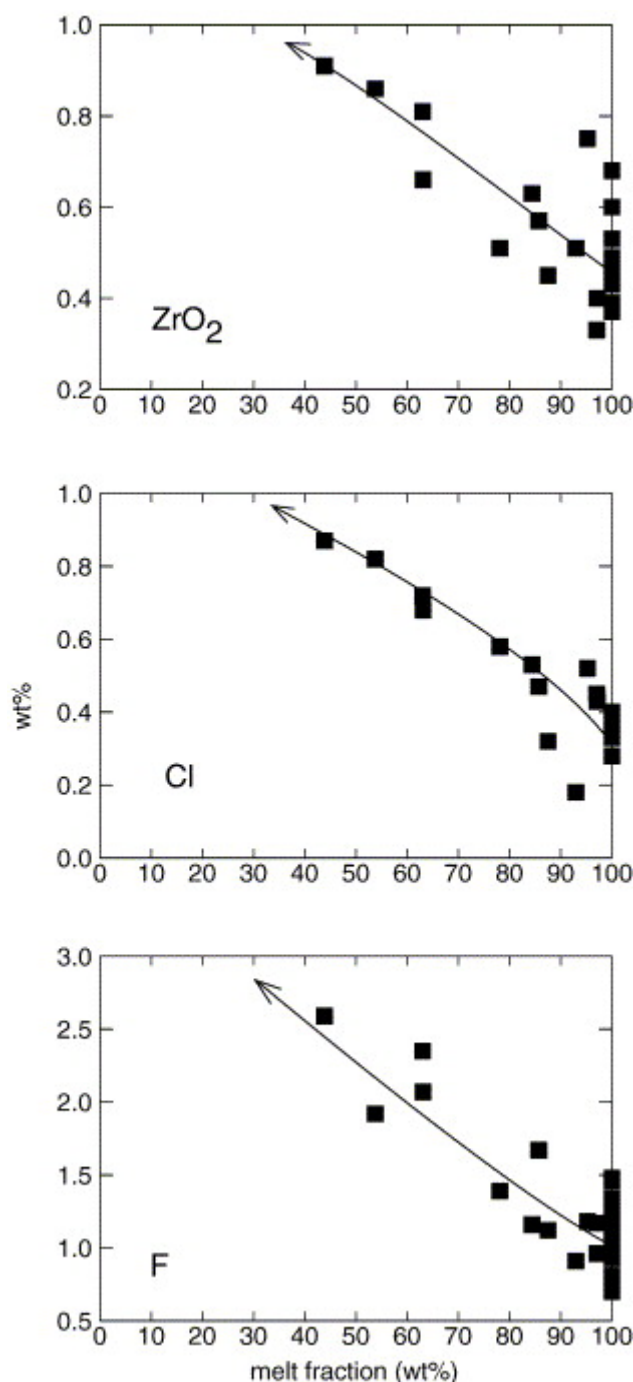


Fig. 6. Variation of Cl, F and ZrO₂ contents of the experimental glasses with crystallisation.

5. Discussion and conclusion

5.1. Equilibrium

The present study uses the same approach as our previous experimental study of the Olkaria comendites, for which conditions close to equilibrium are believed to have been attained ([Scaillet and Macdonald, 2001](#) and [Scaillet and Macdonald, 2003](#)). The use of dry glassy starting material, combined with the strongly peralkaline character of the rhyolite, and the long run durations applied in our experiments, in excess of 49 days for the lowest temperature run, are all factors that should promote attainment of crystal–melt equilibrium in our charges. Observed mineral textures are equant, and minerals are evenly distributed across the charges. From the chemical viewpoint, the trends obtained are regular and vary in a consistent fashion with experimental variables (T and H_2O). We thus conclude that, as for the Olkaria experiments, the results presented here closely approach crystal–melt equilibrium.

5.2. Liquid line of descent

Our experiments define the general liquid line of descent of pantelleritic magmas stored at shallow pressure, under low fO_2 conditions and for a variety of H_2O contents. On the Al_2O_3 – FeO^\square classification diagram of [Macdonald \(1974\)](#), the experimental glasses define a linear trend that extends the field defined by the rhyolites of the Eburru complex ([Fig. 7](#)). Also shown are the glasses obtained from the Olkaria experiments ([Scaillet and Macdonald, 2003](#)), whose most fractionated members were shown to closely approach the Eburru pantellerites. The present work shows clearly that fractionation of Eburru-type pantellerites yields melts that broadly follow the main trend defined by the Olkaria rocks (under reducing conditions). As for Olkaria, this trend of increasing FeO^\square and concomitant decrease in Al_2O_3 seems limitless as long as only tectosilicates fractionate. The Eburru trend is, however, displaced to lower Al_2O_3 compared to that of the Olkaria rocks, a feature that was already apparent in the data of [Scaillet and Macdonald \(2003\)](#). This small difference indicates that, although Olkaria-type comendites produce derivative liquids of clearly pantelleritic affinity, the comendite(s) parental to Eburru pantellerites must have been different from those erupted at the Olkaria complex. The small but perceptible difference in Al_2O_3 between the two groups of magmas possibly reflects a source signature or a pressure/ pH_2O -induced change in the feldspar/quartz ratio in the crystallising assemblage, affecting the rate of Al depletion in residual melts.

The Olkaria comendites are also precluded from being direct parents to the Eburru pantellerites by trace element characteristics. As well as having higher concentrations of most incompatible trace elements, they have higher LILE/HFSE ratios ([Macdonald et al., 1987](#) and [Clarke et al., 1990](#)).

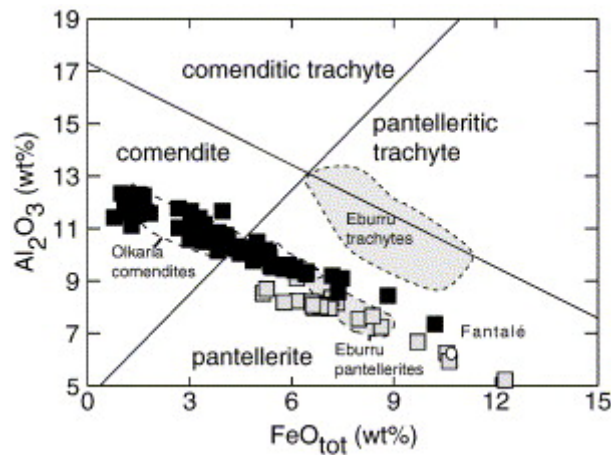


Fig. 7. Al_2O_3 - FeO^\square diagram showing the field of Eburru pantellerites and trachytes, together with the rhyolites of the Greater Olkaria Volcanic Complex. Superimposed are the experimental glasses obtained in this study (grey squares), by crystallising Eburru pantellerite and those from Olkaria rhyolite crystallisation (black squares) (Scaillet and Macdonald, 2001). Fields after Macdonald (1974). The Fantalé pantellerite is from the Fantalé Volcano, Ethiopia (Lacroix, 1930).

From a broader perspective, the facts that pantellerites have lower liquidus temperatures than associated comendites and that they can be demonstrably derived from near-metaluminous rhyolites (Scaillet and Macdonald, 2003) indicate that the most iron- and sodium-rich pantellerites represent the true near-minimum melt composition of alkaline oversaturated series. The fact that most pantellerites are crystal-poor may be due to the exceedingly low viscosity that such magmas can achieve (Dingwell et al., 1998), in particular under H_2O -rich conditions such as at Eburru, which allows efficient melt-crystal separation. Any source that has the potential to produce peralkaline rhyolites will, upon initial melting, yield melts characterised by extreme enrichment in Fe, possibly with FeO^\square contents > 10 – 12 wt.%, low SiO_2 contents (< 63 wt.%, i.e. trachyte-like) and Na_2O contents ≥ 10 wt.%. Owing to their elevated iron and low silica contents, these first (last) drops of liquid might conceivably be quite dense compared to those produced at an advanced stage of melting (or early stages of crystallisation), during which Fe concentrations and thus the density of the liquid decrease. This peculiarity may explain why pantellerites with extreme chemical characteristics have rarely been observed to erupt as lava flows (i.e. Fantalé, Fig. 7). Besides their low volume fraction, their relatively high density is likely to inhibit any crustal ascent, and they will remain trapped at depth. It would be instructive to study the compositions of the extrusive and intrusive rocks in individual volcanic-plutonic complexes, to try to determine if the intrusives are, in general, more Fe-rich than their erupted equivalents. In addition, the geochemical imprint of such hidden, denser liquids on the overall fractionation trends displayed by erupted oversaturated alkaline series remains to be explored quantitatively.

5.3. Pre-eruption conditions

Pre-eruption conditions can be assessed from the phase equilibrium results mainly because pantellerites tend to be sparsely phryic and phenocrysts do not display pronounced chemical zonation (i.e. alkali feldspar). In the present case, only amphibole is suitable to provide constraints on intensive parameters in the source reservoir. Unfortunately, as yet, there are no published chemical analyses of minerals of the Eburru pantellerites. The only such study of which we are aware is that of Ren et al. (2006-this issue), which provides first order constraints on the T - $f\text{O}_2$ conditions of magma evolution at Eburru by the use of the QUILF geothermometer and oxybarometer. The authors report amphibole (ferrichterite) in the

phenocryst assemblage. As stressed above, the occurrence of amphibole phenocrysts in a crystal-poor pantellerite implies a restricted set of T - H_2O conditions, basically below 700 °C and H_2O_{melt} higher than 4 wt.% (Fig. 2), if the pressure of magma storage is around 150 MPa. The temperature range retrieved from mineral equilibria in the Eburru pantellerites is 665–708 °C (Ren et al., 2006-this issue), which agrees well with the phase equilibrium results. The lower end of the temperature range would imply $H_2O_{\text{melt}} > 4$ wt.%. The constraints on temperature and melt H_2O content are very similar to those derived for the Olkaria comendites (Scaillet and Macdonald, 2001), and therefore seem to characterise the regional pattern of magma storage and perhaps production in the Kenya Rift Valley. The fact that many peralkaline rhyolites erupt explosively (e.g., Ayalew et al., 2002 and Peccerillo et al., 2003) is also in agreement with their elevated water contents under pre-eruption conditions. The present work confirms that reduced magmas are not necessarily dry but may have significant amounts of dissolved water, comparable to those found in arc settings. The origin of the difference in fO_2 between arc magmas and peralkaline rhyolites remains enigmatic, however. Perhaps it has to deal with the pronounced Fe-enrichment followed by transitional basalts upon crystallisation which, coupled to their lower H_2O contents relative to arc basalts, helps peralkaline rhyolites to keep the reduced signature of their parental magma. The high Fe contents of residual liquids all the way down to the rhyolite stage may buffer the fO_2 of alkaline magma series to low values.

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