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Neoproterozoic subducted materials in the generation of Mesozoic Luzong volcanic rocks: Evidence from apatite geochemistry and Hf–Nd isotopic decoupling

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ABSTRACT

A combined study including apatite geochemistry, zircon U-Pb, Lu-Hf isotopes and whole-rock geochemistry including Nd isotopes was carried out for the late Mesozoic volcanic rocks from the Luzong Basin, in the lower Yangtze River region, South China. Whole-rock geochemistry indicates the enrichments of large ion lithophile elements (LILE) and light rare earth elements (LREE) as well as depletions of Nb, Ta and Ti. The extremely low Cl contents in apatites strongly contrast with the rather high-K contents in whole rocks. Potential loss of Cl during syn- and post-magmatic processes having been ruled out, Cl-K decoupling is attributed to be a feature inherited from the primary magma, which indicates the involvement of highly dehydrated sediments and altered oceanic crust in the mantle source. A calculation based on apatite and whole-rock geochemistry further illustrates that the source was composed of four end-members in the perspective of Cl/K, Cl/Nb and F/K ratios. The Hf-Nd isotopes are decoupled for the basaltic trachytes from the lower volcanic sequences in the Luzong Basin, with rather low $\varepsilon_{Hf}(t)$ values (mean = -10.3) and inconsistent Hf–Nd model ages (Hf~1.8 Ga, Nd~1.3 Ga), which indicate the "zircon effect" that in turn requires the incorporation of continental detritus in the source via subduction. However, Hf and Nd isotopes are nearly coupled for the rocks from the upper volcanic sequences in the Luzong Basin. Late-Mesoproterozoic two-stage Hf and Nd model ages (ca. 1.2 Ga) of rocks from the upper volcanic sequences in the Luzong Basin are similar to those of the Neoproterozoic igneous rocks from the Jiangnan orogen, suggesting their relationship with the same subduction event. Based on the combined apatite geochemistry and Hf–Nd isotopes, this work suggests that the source of Luzong volcanic rocks might incorporate Neoproterozoic subducted slab fragments and detrital sediments that had been blocked in the deep lithospheric mantle below the Luzong area since the Neoproterozoic assembly between the Yangtze and Cathaysia blocks. The partial melting may be triggered by the back-arc lithospheric extension related to the subduction of Paleo-Pacific plate in the late Mesozoic.

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1. Introduction

Recycling of crustal materials at convergent margins has been a key issue in understanding the evolution of lithosphere (e.g., Stern, 2011). Subduction is a major process for the recycling of oceanic crust, through the incorporation of fluids released from the oceanic crust or the melting of itself. However, the identification of subduction-related magmatism in specific regions has been debated, since crustal delamination and contamination are both able to introduce crustal materials into the magma and usually predestine similar whole-rock elemental and isotopic features typical of arc volcanic rocks (e.g., LILE enrichment, Nb and Ta depletion). In spite of the difficulty in determining the role of crustal materials in the arc-like rocks with

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whole-rock geochemistry, the detailed behaviors of different elements (including LILE) during subduction, identified in recent studies (e.g., Busigney et al., 2003; Spandler et al., 2003; Straub and Layne, 2003a; Rapp et al., 2008), has revealed new fingerprints of arc magmas and the materials that are modified by early-stage subduction which enter into the deeper mantle. Especially, the decoupling of volatiles and LILEs observed by Straub and Layne (2003a), might provide new clues for the determination of subduction process. Moreover, the detrital sediments may also be involved in subduction along with the oceanic slab (Stern, 2011), which might lead to the "zircon effect" (Patchett et al., 1984). This, undoubtedly, will also help uncover potential evidences for subduction in a certain area.

There are a great volume of Mesozoic volcanic rocks in eastern China, and their relationships with the westward and northwestward subduction of Paleo-Pacific plate have been in debate for a long time (e.g., Li, 2000; Li and Li, 2007; Yuan, 2007; Zhang et al., 2008). It has been proposed that the lithosphere of the North China Craton (NCC) was thinned and the thinning accompanied by decratonization took place in the Mesozoic (Fan et al., 2000; Zhang et al., 2002; Yang et al.,

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2010). However, the lithospheric evolution of the South China Block (SCB) seems to be different at the time, since it was not as stable as the NCC, especially during Neoproterozoic time. Determination of the magma source for the Mesozoic volcanic rocks will provide clues for the study of lithospheric recycling and related geodynamic processes. The volcanic rock from the Luzong Basin, which is located in the SCB, adjacent to NCC, is a good target for addressing the puzzling problem. Furthermore, this study will provide insight into the Mesozoic geodynamic framework of eastern China, as the Luzong Basin is a representative one among the five volcanic basins in the lower reaches of the Yangtze River.

Based on the detailed study on apatite geochemistry and zircon U–Pb and Lu–Hf isotopes as well as whole-rock Sm–Nd isotopes from the Luzong volcanic rocks, we recognized a strong Cl–K decoupling in the source and bring out the contrasting Lu–Hf and Sm–Nd isotopic evolution from early to late stages of the volcanic rocks. Integrating with previous geochemical data, the early Neoproterozoic oceanic crust and sediments are proposed to have played an important role in the generation of the volcanic rocks.

2. Geological background

The NCC and SCB are separated by the Qinling orogenic belt in the west and Dabie-Sulu ultra high pressure (UHP) orogenic belt in the east, with the north-striking Tancheng-Lujiang (Tan-Lu) fault cutting through. The SCB is composed of two small blocks integrated by the Jiangnan orogen: the Yangtze Block to the northwest and the Cathaysia Block to the southeast, and both blocks are characterized by distinct geochemical and geochronological record (e.g., Wong et al.,

2011; Yao et al., 2011). The Luzong Basin occurs in the southwest Anhui Province, China, located in the northeastern Yangtze Block (Fig. 1). It is bounded by the Tan-Lu fault and the Dabie Orogen to the west and other secondary faults around (Fig. 1). The Luzong volcanic rocks, which are exposed in the Luzong Basin, could be subdivided into four formations: from bottom to top, the Longmenyuan Formation (*F*-1), the Zhuanqiao Formation (*F*-2), the Shuangmiao Formation (*F*-3) and the Fushan Formation (*F*-4). The rocks are generally rich in potassium, with compositions from trachy-basalt to trachyte. Available laser-ablation (LA)-ICP-MS zircon U–Pb dating shows that the volcanic rocks erupted within the duration from 135 Ma to 127 Ma (Zhou et al., 2008).

Contemporaneous volcanic rocks are widespread along the midlower reaches of the Yangtze River. As a typical case of the Cretaceous volcanic rocks in the area, the Luzong volcanic rocks have drawn much attention in the last three decades. Although their petrogenesis remains in debate, it has been widely accepted that these rocks formed in an extensional setting (Liu et al., 2002; Wang et al., 2006; Xie et al., 2007). In recent years, progressively more attention has been paid to the lithospheric thinning events in eastern China in the late Mesozoic (Yang et al., 2010; and other references). Available whole-rock geochemical data for the Luzong volcanic rocks show arclike geochemical features and enriched mantle like isotopic signatures (Wang et al., 2006). However, the mechanism for the enrichment remains unresolved. Liu et al. (2002) argued that the enrichment resulted from the metasomatism by LILE-rich fluids released from the subducted Paleo-Pacific slab in the Mesozoic. Xie et al. (2007) held that delamination, which involved the lower continental crust, was responsible for the isotopic enrichments. Zhang et al. (2011) showed



Fig. 1. Geological sketch map of Late Mesozoic Luzong volcanic rocks. The Longmenyuan Formation (*F*-1) is composed of basaltic trachytes and trachy-andesites; the Zhuanqiao Formation (*F*-2) is composed of trachy-andesites and basaltic trachytes; the Shuangmiao Formation (*F*-3) is composed of trachy-basalts and basaltic trachytes; the Fushan Formation (*F*-4) is composed of trachytes and phonolites.

that episodic magmatism in the Paleozoic and Mesozoic modified the composition of the ancient lower crust of the NCC and destroyed the structure of the lithosphere, substantially contributing to the decratonization of the NCC. The incorporation of subducted sediments in the mantle source has also been proposed as a possible mechanism (Wang et al., 2006), but the evidence is inadequate and the timing of the subduction event is unclear. Presently, Paleo-Pacific plate subduction is a model favored by those who viewed the late Mesozoic igneous activities in eastern China from the perspective of subduction-related petrogenesis (e.g., John et al., 1990; Faure and Natal'in, 1992; Lapierre et al., 1997; Li and Zhou, 1999; Li, 2000; Li and Li, 2007; Yang et al., 2011), whereas the Paleo-Pacific plate subduction model does not answer some of the key issues (e.g., most late Mesozoic igneous rocks in eastern China are located too far away from the subducting trench; Shao et al., 2000; Zhang et al., 2001; Yuan, 2007; Zhang et al., 2008).

3. Whole-rock and apatite samples

3.1. The Luzong volcanic rocks

Seven representative samples were collected from the Luzong Basin, including two trachy-andesites from F-2 (LZ-1, LZ-2), two trachy-basalts from F-3 (LZ-3, LZ-4), and three trachytes from F-4 (LZ-5, LZ-6 and LZ-7). The trachy-andesites and trachy-basalts show porphyritic textures, and are mainly composed of plagioclase (60%, An = 28%), clinopyroxene (18%), alkaline feldspar (15%), minor olivine (4%) and accessory biotite, magnetite and apatite. Clinopyroxene and most plagioclase occur as phenocrysts while alkaline feldspar and plagioclase microlite are the dominant components of the matrix. Trachytes generally show trachytic texture, and are mainly composed of alkaline feldspar (70%), plagioclase (10%, An = 10%), minor clinopyroxene (7%) and biotite (5%), accessory magnetite (3%) and apatite. Alkaline feldspar (sanidine) and biotite exist as phenocrysts while matrix mainly consists of alkaline feldspar. The sampling was conducted on fresh outcrops. However, careful microscope observations indicate post-magmatic alteration to different extent, with olivine and clinopyroxene being partly replaced by chlorite and calcite.

3.2. Apatite

Apatite is a ubiquitous accessory mineral, which exhibits euhedral crystal with grain size ranging from 50 μ m to 500 μ m in length and 30 μ m to 100 μ m in width. The apatite occurs either as single crystal in groundmass or as inclusions in plagioclase, clinopyroxene, magnetite and in other apatites in all samples from *F*-2 and *F*-3 (Fig. 2a–c). Apatites are much less common in the samples from *F*-4 due to the relatively low whole-rock P contents. The mineral also occurs in groundmass or as inclusions in plagioclase, biotite and magnetite (Fig. 2d). These observations imply that apatites from *F*-2 and *F*-3 crystallized earlier than plagioclase and clinopyroxene whereas those from *F*-4 formed earlier than plagioclase and biotites. BSE images reveal the broad homogeneity of these apatites. Although one of the CL images (Fig. 2e) exhibits weak zoning, later EMPA and laserablation ICP-MS studies show little compositional heterogeneity.

4. Analytical methods

Major elements in apatite were determined using a JEOL JXA-8100 electron microprobe at State Key Laboratory for Mineral Deposits Research (MiDeR), Nanjing University (NJU). All analyses were performed under an accelerating voltage of 15 kV, with a beam current of 20 nA and a beam diameter of 1 μ m. Standards used in this work include natural apatite for F, Cl, Ca and P, galena for S, hornblende for Na, Si, Mg, and Fe. OH was calculated by site difference (Cl+F+

OH = 1 pfu). Each apatite grain was analyzed with 2–10 spots, from core to rim.

All whole-rock samples were prepared by crushing in an agate shatterbox. Major elements and most trace elements were analyzed at ALS Chemex Company (Guangzhou, China), using an X-ray fluores-cence spectrometer (XRF) and an ICP-MS, respectively. The analytical precisions for major elements are less than 1%, and 5–10% for most of the trace elements.

Zircon grains were separated from sample LZ-1(F-2), LZ-5 and LZ-6 (F-4) for U–Pb dating and Lu–Hf isotopic analyses. Trace elements in apatite and zircon U-Th-Pb isotopes were determined using an Agilent 7500a ICP-MS coupled with a New Wave UP213 laser-ablation system at MiDeR, NJU. He carrier gas was used to transport the ablated sample from the standard laser-ablation cell, and then mixed with the Ar gas via a mixing chamber before entering into the ICP-MS torch. All of the spot analyses were carried out using a repetition rate of 5 Hz. The laser-ablation beams are 18 µm and 25 µm in diameter for zircon U–Th–Pb isotopes and apatite trace elements, respectively, NIST610 glass was used as external standard for trace elements with Ca contents (from EMPA) being internal standard. KL-2 glass was used as the reference material to monitor the analytical accuracy. GI-1 was used for external standard with a recommended ²⁰⁷Pb/²⁰⁷U age of 608.5 ± 1.5 Ma (Jackson et al., 2004). Zircon standard Mud Tank (intercept age of 732 ± 5 Ma; Black and Gulson, 1978) was used to monitor the analytical accuracy. U-Pb ages were calculated from the raw signal data using the online software package GLITTER (ver. 4.4) (www.mq.edu.au/GEMOC). The common Pb correction was carried out by using the EXCEL program of ComPbCorr# 151 (Andersen, 2002). The calculations of weighted average ages and plotting on Concordia were finished using the ISOPLOT program (ver. 2.06) of Ludwig (1999). The results are reported in 1σ error.

Dated zircon grains were further analyzed for Lu-Hf isotopes using a Geolas193 laser-ablation system attached to a Nu Plasma HR multicollector ICP-MS at the State Key Laboratory of Continental Dynamics, Northwest University, Xi'an. Instrumental conditions and data acquisition in detail are described in Yuan et al. (2008). Typical ablation time was about 30 s for 200 cycles, with a 5 Hz repetition rate and a beam diameter of 44 μ m (predominantly). He was used as the carrier gas, then combined with argon in a small mixing chamber prior to transport into the ICP torch. In case of the influence of isotopic zoning or with intersecting cracks/inclusions, only the flattest, most stable portions of the time-resolved signal were selected for integration. Isobaric interference of ¹⁷⁶Lu on ¹⁷⁶Hf was corrected by measuring the intensity of the interference-free ¹⁷⁵Lu isotope and using a recommended ¹⁷⁶Lu/¹⁷⁵Lu ratio of 0.02655 (Machado and Simonetti, 2001) to calculate ¹⁷⁶Lu/¹⁷⁷Hf ratios. Correction for isobaric interference of ¹⁷⁶Yb on ¹⁷⁶Hf was performed in 'real time' as advocated by Woodhead et al. (2004) and developed by Iizuka and Hirata (2005). This involves measuring the interference-free ¹⁷²Yb and ¹⁷³Yb during the analysis, calculating mean β_{Yb} value from ¹⁷²Yb and ¹⁷³Yb and using a recommended ¹⁷⁶Yb/¹⁷²Yb ratio of 0.5886 (Chu et al., 2002) and mean β_{Yb} value to calculate $^{176}\text{Yb}/^{177}\text{Hf}$ ratios (Wu et al., 2006). Zircon 91500 was used as the reference standard afterwards, with a recommended 176 Hf/ 177 Hf ratio of 0.282306 \pm 10 (Woodhead et al., 2004). The analytical error for single Lu-Hf isotope analysis results is in 1σ . We have adopted a decay constant for 176 Lu of 1.865×10^{-11} year⁻¹ (Scherer et al., 2001). Initial ¹⁷⁶Hf/¹⁷⁷Hf ratios $\varepsilon_{\rm Hf}(t)$ were calculated with reference to the chondritic reservoir (CHUR) of Blichert-Toft and Albarede (1997) at the time of zircon growth from the magma. Two-stage Hf model age (T_{DM2}) is calculated relative to the depleted mantle with present-day ${}^{176}\text{Hf}/{}^{177}\text{Hf} =$ 0.28325 and ${}^{176}Lu/{}^{177}Hf = 0.0384$ (Nowell et al., 1998; Griffin et al., 2000), bulk crust with present-day $^{176}Lu/^{177}Hf = 0.015$ (Griffin et al., 2002).

The Nd isotopic compositions were determined using the ID-TIMS (Finnigan MAT Triton TI) at MiDeR, NJU. Chemical separation procedures



Fig. 2. (a) and (b): Photomicrographs of apatites from the Luzong volcanic rocks (crossed nicols); (c) and (d): Back scattered electron images; (e) and (f): Cathodoluminescence (CL) images. Note that apatites are quite homogeneous in BSE images and slightly exhibit zonation in CL images. Ap = apatite, Pl = plagioclase, Bt = biotite.

follow Pu et al. (2005), with relative standard deviation (RSD) lower than 5×10^{-6} . Mass fractionation was corrected according to 146 Nd/ 144 Nd = 0.7219. The long-term 143 Nd/ 144 Nd analyzing value of the international standard JNdi-1 is 0.51211±0.000016 (2 σ , n = 67), consistent with the value (0.512115±0.000007) of Tanaka et al. (2000). The $\varepsilon_{Nd}(t)$ values were calculated based on the Nd isotopic compositions of 143 Nd/ 144 Nd (CHUR) = 0.512638 and 147 Sm/ 144 Nd (CHUR) = 0.1967.

5. Results

5.1. Whole-rock major and trace elements

A TAS classification was applied for our samples, as shown in Fig. 3. Samples from all three formations display enrichments of LILEs (K, Rb, Sr, Ba, Pb, U, etc.), which is typical of arc volcanic rocks (Fig. 4). The LREEs are enriched relative to HREEs, while the HFSEs (Nb, Ta, Ti) are obviously depleted. However, it should be noted that the abundances of incompatible elements of the Luzong volcanic rocks are significantly higher than those of typical arc volcanic rocks but close to that of global subducting sediment (GLOSS) calculated by Plank and Langmuir (1998). Detailed data are available in Background Dataset 1.



Fig. 3. TAS diagram for the Luzong volcanic rocks. Data from Liu et al. (2002), Wang et al. (2006), Xie et al. (2007) and this study.



Cs Rb Ba Th U Nb Ta K La Ce As Pb Pr Sr P Nd Sm Zr Hf Eu Ti Gd Tb Dy Y Ho Er Tm Yb Lu

Fig. 4. Primitive mantle normalized spider diagram for the Luzong volcanic rocks, in comparison with GLOSS and arc andesite MI. GLOSS is from Plank and Langmuir (1998); arc andesite MI compositions are from Straub and Layne (2003a); OIB composition is from Sun and McDonough (1989). Primitive mantle composition is from Lyubetskaya and Korenaga (2007).

5.2. Apatite composition

5.2.1. Major elements

Apatites from the Luzong volcanic rocks are generally fluorapatites, while these apatites are also high in hydroxyl, especially for samples from *F*-4. Detailed data are available in Background Dataset 2. It should be noted that nearly all these apatites contain extremely low chlorine (<1000 ppm). Na₂O is hardly detected in apatites from *F*-2 and *F*-3, while apatites from *F*-4 have relatively higher Na₂O (1000– 2000 ppm) and SiO₂ (500–1000 ppm). Additionally, apatites from *F*-4 have relatively low MgO and FeO, indicating the melt in equilibrium with the apatites is more evolved.

5.2.2. Trace elements

As a mineral phase that crystallizes at early stage, apatite could be used to probe into the melt composition in the magma chamber. Profile composition indicates that most major and trace elements exhibit neglectable variation from core to rim, which agrees well with previous BSE observations. Detailed data are available in Background Dataset 3. Generally, apatites from the three formations of the Luzong Basin have similar REE patterns (Fig. 5). Eu(II) and Eu(III) have different partitioning coefficients between apatite and melt due to their different ion radii (Watson and Green, 1981). Therefore, δ Eu in apatite is co-controlled by the δ Eu and the Eu(II)/Eu(III) ratio in the



Fig. 5. Chondrite-normalized REE pattern diagram for apatites from the Luzong volcanic rocks. Chondrite value is from Sun and McDonough (1989).

melt when apatite crystallizes. Whole-rock δEu changes insignificantly from LZ–1 to LZ-7. In addition, most apatite grains occur as inclusions in plagioclases, suggesting their earlier crystallization. Therefore, the constant δEu values (most analysis ranging from 0.40 to 0.50) in all analyzed grains require that the Eu(II)/Eu(III) ratio in the melts in equilibrium with the apatites vary little from LZ1 to LZ-7. The \sum REE contents in apatites from *F*-4 are remarkably higher than those in apatites from *F*-2 and *F*-3, consistent with the relatively low CaO, MgO and FeO contents in the apatites from *F*-4. This might be caused by relatively late crystallization of apatites from *F*-4 because REE partition coefficients between apatite and melt increase with increasing degree of polymerization of the melt (Prowatke and Klemme, 2006).

Other trace elements provide further information about the source and magma evolution. Firstly, Sr drops dramatically in apatites from *F*-4, consistent with the observation in whole-rock composition. The loss of Sr has often been interpreted as a result of fractional crystallization of plagioclase due to the large partitioning coefficient of Sr in plagioclase (Drake and Weill, 1975). However, it is not the case for the volcanic samples from F-4. The apatite grain LZ-6-06 from F-4 is included in plagioclase, suggesting that its formation might have not been affected by the fractional crystallization of plagioclase, but this grain still has fairly low Sr content (382-406 ppm) compared with apatites (Sr>1000 ppm) from F-2 and F-3. Therefore, low Sr contents in apatites and whole rocks of F-4 should be a feature inherited from the magma source, rather than the results of fractional crystallization. Secondly, Zr, U and Th contents increase systematically from F-2 and F-3 to F-4 in whole rocks while such increase is neglectable in apatites. A possible explanation may refer to the crystallization of zircon which is the major reservoir of Zr, U and Th. The low Zr, U and Th contents of apatites from F-4 suggest that zircons might crystallize earlier than apatites in the case of F-4. This is also supported by its highest zircon saturation temperature among the three formations (Wang et al., 2006).

5.3. Zircon dating and Lu-Hf isotopes

Zircons from all samples are clear and transparent, typically 50– 150 µm long and 30–100 µm wide. All zircon grains display prismatic and pyramidal crystal faces, among which, most grains exhibit weakly oscillatory zonation (Fig. 6). Contents of Th (89–5106 ppm) and



Fig. 6. Cathodoluminescence (CL) images of representative zircons from the Luzong volcanic rocks. Small red circles are spots for U–Pb isotopes using LA–ICPMS, and big white dashed circles are spots for Hf isotopes using LA–MC-ICPMS.

U (34–1815 ppm) are high and varying with Th/U ratios ranging from 0.42 to 20.89. Detailed data for zircon U–Pb dating are available in Background Dataset 4.

All of the zircons separated from LZ-1 are concordant in Fig. 7a. The analyzed zircons yield a weighted mean ${}^{206}\text{Pb}/{}^{238}\text{U}$ age of 132 ± 1.5 Ma (MSWD = 0.37, n = 14) for *F*-2. 25 of the zircon grains separated from LZ-5 and LZ-6 are concordant and these ages were adopted when calculating the ages (Fig. 7b). The main population of zircons yield a weighted mean ${}^{206}\text{Pb}/{}^{238}\text{U}$ age of 129 ± 1.2 Ma (MSWD = 1.03, n = 25) for *F*-4.

Zircons from *F*-2 and *F*-4 have been further studied for *in situ* Hf isotopes and the detailed data are available in Background Dataset 5. Zircons $\varepsilon_{\text{Hf}}(t)$ values of *F*-2 show a small variation and yield a mean value of -10.3. However, the values of zircons from *F*-4 (LZ–5 and LZ-6) have relatively large variations (from -7.9 to +6.1), which might result from the incorporation of zircons from earlier erupted rocks (e.g. *F*-2, *F*-3) during magma upwelling. Apart from three zircons from *F*-4 with rather low $\varepsilon_{\text{Hf}}(t)$ (< -5), the other grains have $\varepsilon_{\text{Hf}}(t)$ values ranging from -2.6 to +6.1, constituting a normal distribution. We thus eliminated the low- $\varepsilon_{\text{Hf}}(t)$ zircon grains (LZ-5-06, LZ-5-08 and LZ-5-11) when calculating mean $\varepsilon_{\text{Hf}}(t)$ for *F*-4. Contrasting differences in both $\varepsilon_{\text{Hf}}(t)$ values and zircon two-stage Hf model ages exist between *F*-2 and *F*-4 (Fig. 8). $\varepsilon_{\text{Hf}}(t)$ rises notably from *F*-2 (mean = -10.3) to *F*-4 (mean = +1.2). Zircon two-stage Hf model age peak of *F*-2 occurs at around 1.8 Ga while *F*-4 around 1.2 Ga.

Nd isotopic data of the Luzong volcanic rocks are listed in Background Dataset 6. Basaltic trachytes (LZ-1 and LZ-2) from *F*-2 have $\varepsilon_{Nd}(t)$ ranging from -3.89 to -4.28, similar to that of trachytes (LZ-5, LZ-6 and LZ-7) from *F*-4 (-3.03 to -4.17). Trachy-basalt (LZ-3) from *F*-3 has a lower $\varepsilon_{Nd}(t)$ (-7.99). Two-stage model ages for Nd vary insignificantly from *F*-2 to *F*-4 (1.3-1.2 Ga), except that LZ-3 has a relatively old age (ca. 1.6 Ga).

6. Discussion

Apatite can be a secondary mineral and crystallize at postmagmatic alteration stage. However, it is not the case for the apatites from the Luzong volcanic rocks because most apatite grains analyzed occur as inclusions in plagioclase, and those grains in groundmass display little compositional differences compared with those occurring as inclusions. The zircon grains are also believed to be primary



Fig. 7. Zircon U–Pb concordia diagram for volcanic rocks from the Zhuanqiao (a) and Fushan Formation (b).



Fig. 8. Plot of zircon $\varepsilon_{\text{Hf}}(t)$ vs. ages (a) and histogram of zircon two-stage Hf modal ages (T_{DM2}) (b) for the Zhuanqiao and Fushan Formation samples (LZ-1, LZ-5 and LZ-6).

magmatic because of their euhedral form, weak oscillatory zonation, high Th/U ratios, and their U–Pb ages which are consistent with the previous dating results (Zhou et al., 2008, and the references therein concerning K-Ar and Rb–Sr dating).

Before we address the petrogenesis of the volcanic rocks, the role of crustal contamination should be evaluated, because large-scale crustal contamination can produce a melt similar to what stems from an enriched mantle or arc volcanic rocks (e.g., Dungan et al., 1986; Wolff et al., 2005). Based on the available geochemical data, crustal contamination may be insignificant for the generation of the Luzong volcanic rocks. Firstly, Sr–Nd isotopic studies from previous authors (e.g., Liu et al., 2002; Wang et al., 2006 and Xie et al., 2007) found no co-variation in $\varepsilon_{Nd}(t)$ vs. SiO₂ and (87 Sr/ 86 Sr)_i ratios vs. 1/Sr. Secondly, the negative correlation between La/Nb ratios and SiO₂ contents in whole rock (Fig. 9) also precludes significant crustal contamination.

6.1. Low Cl/K ratio in the Luzong volcanic rocks

6.1.1. Extremely low Cl apatites

According to the available geochemical data, most apatites from intermediate igneous rocks, especially andesites originated from subduction-related settings, have relatively high chlorine contents



Fig. 9. Plot of La/Nb vs. SiO₂. Data from Liu et al. (2002), Wang et al. (2006), Xie et al. (2007) and this study.

(Boyce and Hervig, 2009), while the apatites from granites, especially S-type granites, contain much less chlorine (Sha and Chappell, 1999). The overall high F and low Cl contents in apatites from S-type granites are interpreted to mainly reflect the halogen abundances in the sedimentary source rocks, since they had experienced a relative enrichment of F and preferential loss of Cl during prior weathering process, because of the high solubility of Cl in aqueous solutions (Brehler and Fuge, 1974). Apatites from the Luzong volcanic rocks are characterized by low Cl contents, even lower than apatites from S-type granites (Fig. 10).

Halogen degassing might result in low-Cl contents in apatites. But halogen degassing takes place only when magma gets very near the surface (e.g., <200 MPa; Aiuppa et al., 2004; Stelling et al., 2008). As for the Luzong apatites, neither major nor trace element compositions exhibit remarkable variations from core to rim in these large-size grains ($50-500 \mu m$). Such compositional homogeneity requires stable crystallizing conditions for the apatites, including melt composition, temperature and pressure. Moreover, our apatites, as estimated above, crystallized earlier than plagioclase and even clinopyroxene.



Fig. 10. Logarithmic plot of halogen composition of apatites from the Luzong volcanic rocks, in comparison with apatites from different types of igneous rocks. Data sources, apatites from mafic igneous rocks: Brearly and Jones (1998), Min et al. (2003), McCoy et al. (1996), Semenenko et al. (2004), Kurat et al. (2002), Folco et al. (2004), Boudreau and McCallum (1989), Huntington (1979), Nash (1976), Willmore et al. (2000), Boudreau et al. (1995), Warner et al. (1998), Fodor (2001), Meurer and Natland (2001), Spengler and Garcia (1988); apatites from 1-type and S-type granite: Sha and Chappell (1999); apatites from mantle: O'Reilly and Griffin (2000).

Hence, a deep magma chamber with stable P–T condition may be appropriate for the generation of these apatites, which implies that the occurrence of halogen degassing is unlikely to have occurred.

Brenan (1993) noted that the rapid cooling of eruptive and hypabyssal magmatic systems prevented the diffusion of halogen from the magma, and thus the apatites trapped as inclusions in other minerals might preserve the initial halogen chemistry of magma source (Webster et al., 2009). Piccoli and Candela (2002) pointed out that volcanic apatites rarely exhibit chemical zoning, well consistent with the observation of this work. Thus, it could be concluded that apatites from the Luzong volcanic rocks are free from post-magmatic diffusion of halogens and may preserve the initial information of the halogen composition of the parental melt. Therefore, these low-Cl apatites reflect the low Cl concentration in the parental magma, which is a feature inherited from the source.

Fluorine and chlorine are both incompatible elements, their contents vary with the fractionation of melt. Therefore parameters that index fluorine and chlorine to other highly incompatible elements have been introduced to study halogen compositions of the source. K is commonly used (e.g., Aoki et al., 1981; Jambon et al., 1995; Douce and Roden, 2006) because F/K and Cl/K ratios vary within narrow ranges among continental basalts, MORB and OIB samples that are not contaminated by seawater. Arc volcanic rocks exhibit significantly elevated Cl/K ratios due to the large incorporation of Cl released from the altered oceanic crust and sediments during early-stage subduction. In fact, in addition to F/K and Cl/K ratios, other ratios (e.g., F/P, Cl/Nb and etc.) were also frequently adopted, since these element pairs are similarly incompatible (Schilling et al., 1980).

The Luzong volcanic rocks are rather enriched in K, strongly contrasting with the extremely low Cl contents in apatites. In fact, the Luzong volcanic rocks generally have much higher K contents than those referred in Fig. 10 while the Luzong apatites have the lowest Cl contents. It is unrealistic to expect a sort of melt with normal Cl/K ratio to produce such low-Cl apatites and high-K whole rocks simultaneously, when potential loss of Cl during syn- and post-magmatic processes has been eliminated. An alternative explanation is the Cl–K decoupling in the source.

A low Cl/K ratio source is required to be responsible for the low-Cl apatites and high-K contents in whole rocks. The incorporation of lower continental crust in the mantle source, as suggested by Xie et al. (2007), is unable to produce a magma with low Cl/K ratio because Cl/K ratio of lower continental crust (~0.05; Rudnick and Gao, 2004) is believed to fall in the variation range of uncontaminated MORB (i.e. 0.01-0.07; Michael and Cornell, 1998). Moreover, crustal delamination would more likely produce Cl-Nb decoupling, instead of Cl-K decoupling, because Nb might be trapped in rutile in the residue (Hermann and Rubatto, 2009). Since Nb is also depleted in the Luzong volcanic rocks, Cl-Nb decoupling may be insignificant. Contamination of middle and/or upper continental crust is able to decrease Cl/K ratio in the magma (Rowe and Lassiter, 2009). However, large crustal contamination is unlikely as discussed above. Moreover, it remains rather questionable whether crustal contamination alone can produce such low-Cl apatites because apatites from S-type granite have even higher Cl contents. Meanwhile, apatites from continental basalts, as far as we are aware of, have much higher Cl contents than the Luzong apatites.

6.1.2. A possible mechanism responsible for the low Cl/K ratio

The low Cl/K ratios of the Luzong volcanic rocks might be accomplished in two steps.

As one of the most soluble elements in aqueous solutions (Brehler and Fuge, 1974), Cl tends to be lost during weathering. In contrast, K and F are much more conservative due to their relatively low migration rate, resulting in the low Cl/K ratios of continental detritus. And this is why Cl is depleted in upper continental crust, because Cl is transported to the oceans and accumulates in seawater (Pyle and Mather, 2009).

When the detrital sediments were transported into subduction trench, the Cl concentration might be elevated to a certain extent through the interaction with seawater which contains as much as 20,000 ppm Cl. But the following dehydration process during the early-state subduction is able to completely remove the Cl buried in the subducted detrital and pelagic sediments (Straub and Layne, 2003b).

Weathering and dehydration are both able to remove Cl in rocks but the latter process, subduction-related dehydration, might be more efficient in Cl-extraction. Much work has been done to quantitatively assess the amount of halogens and LILEs that survive the early-stage subduction and enter into the deeper mantle. Straub and Layne (2003b) estimated that nearly all Cl recycled into the mantle wedge during shallow subduction. Lassiter et al. (2002) argued that Cl retained in deep recycled crust was less than 50 ppm after the extraction during subduction. Scambelluri et al. (2004) calculated that about 45 ppm Cl was hosted in the residual high pressure minerals of subducting oceanic slab. However, F extraction is not as efficient as Cl during early-stage subduction. For example, only 4-5% F was recycled at the Izu arc, leading to the strong fractionation of F from Cl and H₂O (Straub and Layne, 2003b). F contents in sediments vary in a large range, depending on the rock type of sediments. Pelagic clay contains as much as 1300 ppm F (Li and Schoomaker, 2003) while continentderived sediments might contain about 600 ppm F, similar to the upper crust (Straub and Layne, 2003b). As for LILEs, geochemical studies of meta-sedimentary rocks in the western Alps (Busigney et al., 2003) suggested that LILEs behaved rather conservatively and were not significantly lost during subduction-related devolatilization (Busigney et al., 2003; Spandler et al., 2003). Decoupling between volatiles and LILEs was observed in halogen-rich andesite melt inclusions from the Izu arc volcanic front (Straub and Layne, 2003a). A continuous series of hydrous phases (e.g., amphibole, lawsonite, muscovite, phengite, paragonite, epidote-group minerals, phase egg, K-hollandite) would be stable in subducted continental sediments down to a depth with 15 GPa in pressure (Schmidt, 1996; Ono, 1998; Hermann and Green, 2001; Hermann, 2002; Spandler et al., 2003; Rapp et al., 2008). LILEs released from a decomposed hydrous phase could be re-trapped into another hydrous phase that is stable in higher metamorphic grades. Thus, the whole-rock budget of LILEs in deeply subducted sediments may stay broadly unmodified when passing through the sub-arc regime (Rapp et al., 2008).

There is increasing evidence to suggest that dehydration processes which take place during early subduction are able to fractionate Cl from F and LILEs. Dehydrated residue might be rather low in Cl but still rich in LILEs, as is the key to the Cl–K decoupling in the source.

6.1.3. A four-end-member source model—implication from calculated Cl/K, Cl/Nb and F/K ratios

Since apatites from the Luzong volcanic rocks have rarely undergone obvious subsequent re-equilibration with late-stage fluids or melts, they provide an ideal opportunity to calculate the halogen compositions of the melt. Boyce and Hervig (2009) gave a detailed study on the halogen composition of the volcanic rocks from Irazú volcano formed in 1963 and 1723 eruptions. Samples from the two eruptions have weak whole-rock compositional differences comparing with those from *F*-2 and *F*-3. Therefore, we choose their partitioning coefficients for fluorine ($D_F = 15$) and chlorine ($D_{CI} = 5$) between apatite and silicate melt. As for *F*-4, calculation of halogen compositions for the melt was given up because of the relatively larger compositional variations in whole rock.

The influence of fractional crystallization should be evaluated before using whole-rock K content as that of the melt in equilibrium with apatites. Prowatke and Klemme (2006) observed that REE partitioning coefficients between apatite and melt increase with increasing degree of polymerization of the melt. Their study outlined the Sm partitioning coefficient array with increasing SiO₂ in the melt. Accordingly, a modeling has been carried out to estimate the extent of fractional crystallization after the crystallization of apatite, by comparing the whole-rock SiO₂ contents with the assessed SiO₂ content in the melt in equilibrium with apatite (Fig. 11). The cross range estimates that SiO₂ in the melt when apatite crystallizes is 54% to 57%, which is slightly higher than the whole-rock SiO₂ contents of samples from *F-2* and *F-3*. This might be caused by the involvement of earlier crystallized, low-SiO₂ phases (e.g., olivine) into the magma after apatites had crystallized. Anyway, fractional crystallization, which might lead to underestimated ratios like Cl/K in our calculation, is insignificant.

There might still be other unknown factors that may affect the accuracy of this calculation. But these do not hamper our discussion of the Cl–K decoupling in the source. As an attempt, this calculation helps quantify the Cl–K decoupling and helps better illustrate source compositions.

The calculated halogen and related element ratios are listed in Table 1. Cl/Nb ratios (3.49-13.88) are roughly between that of N-MORB (~11; compiled from Sun and McDonough, 1989) and primitive mantle (~3; Lyubetskaya and Korenaga, 2007) and F/P values (0.35-1.07) between that of N-MORB (~0.4; compiled from Sun and McDonough, 1989) and crust (~1; Rudnick and Gao, 2004). Obviously, F/K ratios of our samples generally fall on the array defined by MORBs, OIBs, arc volcanic rocks and continental basalts (Fig. 12). As to the Cl/K ratios, the story is totally different. Extremely low Cl/K ratios (<0.004), even lower than in uncontaminated MORBs (i.e. 0.01–0.07; Michael and Cornell, 1998), make the volcanic rocks of *F*-2 and *F*-3 strongly deviate from the array (Fig. 12), indicating that chlorine decouples with potassium remarkably.

Four possible end-members, including dehydrated altered and unaltered oceanic crust, dehydrated sediments and depleted mantle, are introduced to model the source. Conservative elements, like Nb and Ta, are expected to be free of modification through subduction due to their immobility in fluid. Hence, we refer to GLOSS (Plank and Langmuir, 1998) for LILEs and Nb concentrations of dehydrated sediments.

As summarized in Table 2, sediments and altered oceanic crust that survive the subduction-related dehydration do display extremely low Cl/K ratios, as is associated with the decoupling between volatiles and LILEs at arc front which is marked by high Cl/K ratio. F/K, Cl/Nb and Cl/K ratios of the Luzong volcanic rocks generally plot in the ranges



Fig. 11. Plot of Sm vs. SiO₂ concentrations in the melt. Black line represents the evolving trend of Sm concentration in the melt estimated from whole-rock data. Grey belt represents estimated Sm concentration in the melt which is in equilibrium with apatites from the Luzong volcanic rocks (see Prowatke and Klemme (2006) for detailed relationship between Sm partitioning coefficient and SiO₂ in the melt). The cross range of the black line and the grey belt corresponds to the estimated variation range of SiO₂ content in the melt during the apatite crystallization.

Table 1

Calculated halogen concentrations (ppm) in the melt.

	LZ-1	LZ-2	LZ-3	LZ-4
F ^a Cl ^a K ^b Nb ^b P ^b Cl/K	$\begin{array}{c} 1640 \ (136) \\ 76 \ (22) \\ 41,300 \\ 21.8 \\ 1530 \\ 1.84 \times 10^{-3} \end{array}$	1547 (183) 85 (28) 32,900 15.5 1820 2.58×10 ⁻³	$\begin{array}{c} 1518 \ (142) \\ 84 \ (24) \\ 27,100 \\ 9.5 \\ 2070 \\ 3.10 \times 10^{-3} \end{array}$	$\begin{array}{c} 1108 \ (341) \\ 111 \ (48) \\ 35,400 \\ 8.0 \\ 3190 \\ 3.14 \times 10^{-3} \end{array}$
Cl/Nb F/P F/K	3.49 1.07 0.040	5.48 0.85 0.047	8.84 0.73 0.056	13.88 0.35 0.031

 1σ of calculated melt F and Cl contents reported in terms of last significant digits in the parentheses.

^a Based on F (Cl)_{melt} = F (Cl) average of EMPA data/ $D_{F (Cl)}$.

^b Comes from whole-rock data.



Fig. 12. Plots of halogens vs. K, comparing the Luzong volcanic rocks with MORB, OIB, arc volcanic rocks and continental basalts. F generally develops a good coupling with K in different igneous rocks including the Luzong volcanic rocks. In the case of Cl/K ratio, both arc and the Luzong volcanic rocks deviate from the Cl–K coupling array but stretch to opposite sides. Data sources, MORB: Sigvaldason and Óskarsson (1986); Michael and Cornell (1998); Moune et al. (2007); PETBD database: http://beta.www.petdb.org/. OIB: Aoki et al. (1981); Hansteen and Gurenko (1998); Davis et al. (2003). Arc volcanic rocks: Straub and Layne (2003a). Continental basalts: Aoki et al. (1981); McHone (2002).

Table 1	2
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	Luzong volcanic rocks	Dehydrated unaltered oceanic crust ^a	Dehydrated altered oceanic $\mbox{crust}^{\rm b}$	Dehydrated sediments ^c	Depleted mantle ^d
F ppm	1108-1640	100	/	611-1300	11
Cl ppm	76–111	45	45	45	0.51
K ppm	$(2.71-4.13) \times 10^4$	1250	2.68×10^{4}	1.69×10^{4}	60
Nb ppm	8.0-21.8	2.20	3.56	8.94	0.21
F/K	0.03-0.06	0.08	/	0.04-0.08	0.183
Cl/K	$(1.84 - 3.14) \times 10^{-3}$	36×10^{-3}	1.68×10^{-3}	2.65×10^{-3}	8.50×10^{-3}
Cl/Nb	3.49-13.88	20.45	12.64	5.03	2.43

45 ppm is estimated as an average Cl concentration of dehydrated altered/unaltered crust plus sediments as a whole.

Slash represents data not available.

Note that GLOSS represents an average composition of subducted sediments. More continent-derived sediments involved in the source might still lower the Cl/K ratio of dehydrated sediments. Moreover, we use the original abundance of F in sediments as that of dehydrated sediments so that the F/K ratio of completely dehydrated sediments estimated here should be regarded as an upper limit.

^a Compiled from Taylor and McLennan (1985) and Michael and Cornell (1998).

^b K and Nb concentrations estimated according to ODP Sites 801 and 1149 (Kelley et al., 2003).

^c See text for data sources.

^d Depleted mantle composition according to Salters and Stracke (2004).

defined by dehydrated altered and unaltered oceanic crust, dehydrated sediments and depleted mantle. Therefore, it is possible that the halogen compositions of F-2 and F-3 were produced by mixing some or all of the four end-members, but at least one of the two low-Cl/K end-members (dehydrated sediments and dehydrated altered crust) is required to be present in the source. Because K abundances of dehydrated sediments and dehydrated altered oceanic crust are higher than unaltered crust and depleted mantle by more than a magnitude, even a small input of the low-Cl/K end-members would strongly affect the Cl/K ratio in the source. For example, an ideal source model consisting of 70% from depleted mantle, 10% from dehydrated sediments and 10% from dehydrated altered oceanic crust as well as 10% from dehydrated unaltered oceanic crust (all in wt.%) renders a Cl/K ratio as low as 0.003, close to a source model composed of pure dehydrated sediments or dehydrated altered oceanic crust. So the mixing proportion of the four end-members might not be significant as long as either of the two low-Cl/K end-members is present in the source. Cl/Nb and F/K ratios of the four end-members outline relatively narrow variation ranges. Mixture of them in any proportion would produce the final Cl/Nb and F/K ratios similar to those of F-2 and F-3.

We did not calculate the halogen contents in the melt of *F-4*, but the similar low-Cl apatites and the even higher K contents in whole rocks seem to promise the similar low Cl/K ratio of the magma, which requires the presence of low-Cl/K end-members in the source of *F-4*.

6.2. "Zircon effect": Hf isotopic evidence

6.2.1. Hf-Nd decoupling—evidence for detritus-rich sediments in the source

Zircon $\varepsilon_{\rm Hf}(t)$ values dramatically rise from *F*-2 (mean = -10.3) to *F*-4 (mean = +1.2) while whole-rock $\varepsilon_{Nd}(t)$ values vary little from F-2 (-3.89 to -4.28) to F-4 (-3.03 to -4.17). It seems that there exists a component in the source which controls the Lu–Hf isotopic system but has little effect on the Sm-Nd isotopic system. The Nd and Hf isotopes are nearly coupled ($\varepsilon_{Nd}(t)$ slightly lower) in F-4 but obviously decoupled in F-2. Similarly, the two-stage Nd model ages are fairly similar from F-2 to F-4 (1.3-1.2 Ga), close to the two-stage Hf model ages of F-4 (ca. 1.2 Ga), while totally different from the twostage Hf model ages of F-2 (ca. 1.8 Ga). The Hf-Nd decoupling of F-2 results from the rather low $\varepsilon_{\rm Hf}(t)$. Large Hf isotopic deviation might be a result of involvement of detritus-rich sediments in the source, which contain considerable amounts of detrital zircons, i.e., the sediments incorporated in the source should have a large proportion of continental detritus. Early studies questioned the possibility of presence of zircon in the sediments that subducted into the mantle. Patchett et al. (1984) considered the mixing of turbidites and pelagic sediments to be unlikely, because they might be deposited far from one another in the ocean. Thus they concluded that detrital zircons could not be transported to the abyssal plains and would not be involved in subduction-related recycling. However, Plank and Langmuir (1998) argued that Patchett's conclusion was based on the distribution of surface sediments, where there were indeed great distances between different sedimentary provinces. Plate movement produces total sedimentary columns that have Lu-Hf parameters similar to terrigenous sediments (Plank and Langmuir, 1998). Recently, "zircon effect" has been clearly observed in chemical and Hf isotopic compositions of Lesser Antilles forearc sediments (Carpentier et al., 2009). In fact, models that neglect the direct incorporation of zircon-rich detritus in the source cannot render the striking Hf–Nd decoupling in the volcanic rocks from F-2. Delamination of lower crust would introduce crustal material that is expected to change Hf and Nd isotopes simultaneously in the melt. Especially, garnet in the lower crust, which is enriched in Lu, might even potentially elevate ¹⁷⁶Hf/¹⁷⁷Hf relative to ¹⁴³Nd/¹⁴⁴Nd in the melt (Patchett et al., 1981; Schmitz et al., 2004), rather than the lowered ¹⁷⁶Hf/¹⁷⁷Hf observed in F-2. The melting of fluid-metasomatized mantle, as suggested by Liu et al. (2002), should produce magma that displays no appreciable Hf isotopic deviation from depleted mantle because Hf behaves as an immobile element in fluids (Barry et al., 2006).

6.2.2. The consumption of sediments in the source from F-2 to F-4

The variation of $\varepsilon_{\rm Hf}(t)$ values from *F*-2 to *F*-4 reflects the varied amount of detritus-rich sediments incorporated into the source. $\varepsilon_{\rm Hf}(t)$ values rise significantly from *F*-2 to *F*-4, indicating that the detritusrich sediments were gradually consumed relative to other endmembers in the source. This speculation is supported by a comparative study of LILEs of the three formations. We index incompatible elements to Nb, which is both conservative and highly incompatible, and plot these ratios against Nb, to distinguish between source variation and fractional crystallization during magma ascending.

Fig. 13 shows that Rb/Nb, Ba/Nb, K/Nb, Sr/Nb and La/Nb form remarkable decreasing trends with increasing Nb. Assuming that trace element concentrations satisfy the fractional crystallization equation:

$$\frac{C'_{\alpha}}{C_{\alpha}} = F^{D_{\alpha} - 1} \tag{1}$$

$$\frac{C'_{\beta}}{C_{\beta}} = F^{D_{\beta}-1} \tag{2}$$

whereby C_{α} and C_{β} = concentrations of element α and β in the parental magma respectively, C'_{α} and C'_{β} = concentrations of element



Fig. 13. Plots of Rb/Nb vs. Nb, Ba/Nb vs. Nb, K/Nb vs. Nb, Sr/Nb vs. Nb and La/Nb vs. Nb. See the negative correlation in all plots. LILE/Nb and La/Nb decrease with the increase of Nb and reaches the low-most point when it comes to the Fushan Formation. Data from Liu et al. (2002), Wang et al. (2006), Xie et al. (2007) and this study.

 α and β in the upwelling melt respectively, D_{α} and D_{β} = bulk partitioning coefficients of element α and β respectively, F = crystallinity. Accordingly:

$$\frac{C'_{\alpha}}{C'_{\beta}} = \frac{C_{\alpha}}{C_{\beta}} \bullet F^{D_{\alpha} - D_{\beta}}$$
(3)

where F could be substituted as:

$$F = \left(\frac{C'_{\beta}}{C_{\beta}}\right)^{\frac{1}{D_{\beta}-1}} \tag{4}$$

Thus, Eq. (3) could be written as:

$$\frac{C'_{\alpha}}{C'_{\beta}} = \left(\frac{C_{\alpha}}{C_{\beta}}\right) \cdot \left(\frac{C'_{\beta}}{C_{\beta}}\right)^{\frac{D_{\alpha} - D_{\beta}}{D_{\beta} - 1}}$$
(5)

The partial derivative of Eq. (5) with respect to C_{β} :

$$\frac{\partial \left(\frac{C'_{\alpha}}{C'_{\beta}}\right)}{\partial C'_{\beta}} = \left[\left(\frac{C_{\alpha}}{C_{\beta}}\right) \cdot \left(\frac{D_{\alpha} - D_{\beta}}{D_{\beta} - 1}\right) \cdot C^{\frac{D_{\beta} - D_{\alpha}}{D_{\beta} - 1}}_{\beta} \right] \cdot C'_{\alpha}^{\frac{1 + D_{\alpha} - 2D_{\beta}}{D_{\beta} - 1}}$$
(6)

A more incompatible element corresponds to a smaller D. Suppose that element α is more incompatible than element β , and we find the

right side of Eq. (6)>0. With the left side of Eq. (6) symbolizing the slope of the trend plotted in α/β vs. β , we expect a positive correlation between α/β and β . On the contrary, when element α is less incompatible than element β , a negative correlation should be expected.

Generally, Rb and Ba are more incompatible than Nb in magmatic system while K, Sr less incompatible. Therefore, a positive correlation between Rb/Nb vs. Nb and Ba/Nb vs. Nb is expected and a negative one for K/Nb vs. Nb and Sr/Nb vs. Nb. However, Fig. 13 shows a pronounced negative correlations between LILE/Nb ratio vs. Nb for all Rb, Ba, K and Sr. Negative correlation between Rb/Nb vs. Nb and Ba Nb vs. Nb is hard to understand for a magmatic evolution that is governed by fractional crystallization. In fact, the above modeling is based on the assumption that the source is broadly homogeneous and stays constant in composition throughout the time. Ideally, a dualend-member source is required to produce batches of magma that render the elemental evolution observed in Fig. 13: one is characterized by relative LILE (and probably LREE) enrichment while the other is characterized by relative Nb enrichment. Higher Nb content in whole rock requires more Nb-rich end-member added to the source, thus yielding lower LILE/Nb ratios as observed. The younger F-4 trachytes has the lowest Rb/Nb, Ba/Nb, K/Nb and Sr/Nb ratios, suggesting that the LILE-rich end-member accounts for a remarkably smaller proportion in the source of F-4. Similarly, La/Nb ratio drops significantly from F-2 and F-3 (~5) to F-4 (~1.5), indicating less addition of supracrustal materials into the magma source for F-4. As suggested above, there are four possible end-members in the source of the Luzong volcanic rocks: dehydrated sediments, dehydrated

altered oceanic crust, dehydrated unaltered oceanic crust and depleted mantle. Dehydrated sediments and/or altered oceanic crust act as the LILE-rich end-member(s) while unaltered oceanic crust and/or depleted mantle act as the relative Nb-rich end-member(s). Sediments are gradually consumed relative to the other three end-members from F-2 to F-4, making altered oceanic crust the dominant contributor for the excess LILEs of F-4. Exactly, the source of F-4 might be mostly composed of altered/unaltered oceanic crust, depleted mantle and the residual products (crust- and mantle-derived) of the previous partial melting batches.

6.3. Timing of the subduction event

Though the Lu–Hf isotopes obtained from zircons from the Luzong volcanic rocks represent mixing isotopic compositions of four endmembers, these four end-members played quite different roles in producing the Hf isotopic compositions in the source. Depleted mantle contributed little to the Hf isotopic compositions of the source for two reasons below:

- (1) The Luzong volcanic rocks are characteristic of rather low Cr and Ni contents (less than 10 ppm, generally), both of which are quite abundant in depleted mantle (Cr~2500 ppm, Ni~1960 ppm; Salters and Stracke, 2004) but relatively low in oceanic crust (Cr~270 ppm, Ni~135 ppm; Taylor et al. 1985) and GLOSS (Cr~78.9 ppm, Ni~70.5 ppm; Plank and Langmuir, 1998). Though Cr and Ni are gradually consumed during fractional crystallization, it will still be puzzling to observe the whole-rock Ni and Cr concentrations as low as low-Ca granite (Cr~4.1 ppm, Ni~4.5 ppm; Turekian and Wedepohl, 1961) while whole-rock compositions stay basaltic or trachytic. One possible explanation is that the materials derived from depleted mantle had very limited contribution to the source of the Luzong volcanic rocks.
- (2) Abundances of trace elements, including Hf, are extremely low in depleted mantle, less than those of oceanic crust and sediments by a magnitude or more. For example, an ideal source consists of 70% from depleted mantle and 20% from oceanic crust as well as 10% from detrital sediments (all in wt.%) receives less than 20% Hf from depleted mantle. Therefore, low Hf content makes depleted mantle rather inefficient in controlling the source Hf isotopic compositions in the presence of oceanic crust and sediments.

and the subducted oceanic crust. "Zircon effect" is remarkable in *F*-2, making the calculated zircon Hf T_{DM2} of *F*-2 significantly older than the actual age of the oceanic crust. Whereas, detrital sediments gradually faded away from *F*-2 to *F*-4 and two-stage Hf model ages of *F*-4 (ca. 1.2 Ga) are fairly consistent with the two-stage Nd model ages (1.3–1.2 Ga). Approximately coupled Hf–Nd model ages indicate that the source of *F*-4 receives the least contribution from detritus-rich sediments. Thus, the two-stage Hf and Nd model ages of *F*-4 provide a relatively close proxy for the formation age of the oceanic crust. It goes, therefore, the subducted oceanic crust might be around 1.2 Ga old, and the subduction process should take place at ca. 1.0 Ga, which is absolutely older than the formation time of the Luzong volcanic rocks.

are actually controlled by two end-members: detritus-rich sediments

The subducted Paleo-Pacific plate has been favored by quite a lot of researchers as the material source (or an end-member) when they explained the large-scale Mesozoic magmatic events that took place in eastern China, including the Luzong area (e.g., John et al., 1990; Faure and Natal'in, 1992; Lapierre et al., 1997; Li and Zhou, 1999; Li, 2000; Liu et al., 2002; Li and Li, 2007). However, modern oceanic crust is hardly older than 200 Ma, and thus the Paleo-Pacific oceanic crust that subducted in the Mesozoic is unlikely to be much older than 400 Ma, suggesting that the Mesozoic Pacific plate subduction model is an impossible material source for the generation of the Luzong volcanic rocks.

In fact, the Neoproterozoic magmatism and the related subduction processes are very significant in South China, especially in the periphery of the Yangtze Block. According to the tectonic location of the Luzong volcanic rocks, two possible early Neoproterozoic subduction zones can be responsible for the oceanic crust and sediments in the formation of these volcanic rocks: one in the northern margin of the Yangtze Block, and the other in the southeastern margin of the Yangtze Block. The former area consists of a great amount of Neoproterozoic granitic and basaltic rocks that were finally metamorphosed to form the UHP metamorphic rocks in the Sulu-Dabie Orogen during the Triassic continent-continent collision (Li et al., 1993) between the South and North China blocks, while the latter shows a clear arc-continent collision history that led to the formation of the Jiangnan orogen at ca. 860–800 Ma (Wang et al., 2007). We think the subduction in the northern margin of the Yangtze Block may not be responsible here, because the Luzong area could not have been so close to the north margin of the Yangtze Block at that time, or it will be difficult to imagine that the subducted oceanic crust and related sediments could be preserved so well in the lithospheric mantle and survived later Triassic intense tectonic events that included continental deep subduction and significant



Since altered and unaltered oceanic crust have similar Lu–Hf isotopes, zircon two-stage Hf model ages of the Luzong volcanic rocks

Fig. 14. Cartoons illustrating the genesis of the late Mesozoic Luzong volcanic rocks. Slab fragments of the ancient ocean between the Yangtze and Cathaysia Block that closed up at ~860 Ma, were retained in the upper mantle till back-arc extention gave rise to their partial melting in the late Mesozoic.

exhumation. Moreover, there is strong rift magmatism along the north margin of the Yangtze Block in the Neoproterozoic (Zheng et al., 2004), which is hardly seen in or near the Luzong area. Strong mantle convection, as the trigger mechanism of rift magmatism, might have also seriously endangered the preservation of the slab fragments and sediments blocked in the nearby mantle.

On the other hand, however, the Jiangnan Orogen has remained stable since its formation, with few weak interferences, e.g., Paleozoic and earlier Mesozoic activities. The early subduction process in the area is supported by the Neoproterozoic ophiolites (Zhou et al., 1989; Chen, et al., 1991; Li and McCulloch, 1996) and arc volcanic rocks in the eastern Jiangnan orogen (Li et al., 2009). Moreover, it has been proposed that the juvenile crust growth took place in late Mesoproterozoic to early Neoproterozoic (Zheng et al., 2008), with the model Hf–Nd model ages ranges from 1.3 to 1.2 Ga, similar to those of the Luzong volcanic rocks. Therefore, the early Neoproterozoic subduction to the eastern margin of the Yangtze Block might bring some materials (including the oceanic crust and overlain sediments) to the deeper part which became the source for the later formation of Luzong volcanic rocks.

The scenario could now be described as follows: ~200 km lateral subduction brought the slab and sediments through the sub-arc regime and probably down to the keel of the lithospheric mantle beneath the Yangtze Block. The early-stage subduction had efficiently extracted Cl and part of the water buried in the slab and the sediments above. The slab then broke down into fragments in the mantle during deep subduction. Given their slightly higher density, a few of the small-size slab fragments might still be blocked in the rigid lithospheric mantle and stopped subducting. These slab fragments were retained in the upper mantle for a long time until the late Mesozoic, when the west and northwest subduction of the Paleo-Pacific plate produced back-arc extentional settings (Xu et al., 2010). Lithospheric extension triggered the decompression melting of the ancient slab fragments and sediments retained in the upper mantle as well as depleted mantle beneath the Luzong area in the late Mesozoic, and resulted in the formation of the Luzong Basin (Fig. 14).

7. Conclusions

We conclude that the late Mesozoic Luzong volcanic rocks originated from partial melting of Neoproterozoic oceanic crust and overlain sediments as well as depleted lithospheric mantle. Contrasting low-Cl apatites and high-K whole rocks indicate the strong Cl–K decoupling of the primitive magma, which requires the existence of highly dehydrated (detritus-rich) sediments and dehydrated altered oceanic crust in the source. Presence of detritus-rich sediments in the source is supported by the observed "zircon effect". Coupled Hf–Nd model ages of the younger Fushan Formation imply that the subduction responsible for the oceanic crust and sediments might take place in the early Neoproterozoic era.

Apatite, a ubiquitous mineral in igneous rocks, could provide a precious tool to learn the halogen composition of melt, especially for old rocks when melt inclusions are unavailable. More attention should be paid to the Cl-poor apatite from K-rich volcanic rocks, which may imply the strong Cl–K decoupling in the primitive magma. Extremely low Cl/K ratio might prove to be a powerful indicator of highly dehydrated subducted materials preserved in the deep lithospheric mantle for intraplate magmatism.

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