ORIGINAL PAPER

Late Miocene to Pleistocene potassic volcanism in the Republic of Macedonia

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Received: 5 December 2007 / Accepted: 27 March 2008 / Published online: 8 May 2008 © Springer-Verlag 2008

Abstract The potassic (K) to ultrapotassic (UK) volcanic rocks cropping out in the Vardar Zone of Macedonia and southern Serbia span in age from Late Miocene (6.57 Ma)

Editorial handling: J.G. Raith

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ATOMKI, Hungarian Academy of Sciences, 4001 Debrecen, Hungary e-mail: pecskay@namafia.atomki.hu to Pleistocene (1.47 Ma). The main identified outcrops are in the Kumanovo, Sveti Nikole, Shtip and Demir Kapia areas; the southernmost occurrences of these volcanic rocks are located in the large Kozuf Massif (Voras Massif in Greece) at the Macedonia-Greek border. Three distinct groups may be distinguished. The first group has a shoshonitic affinity and occurs in the Kozuf Massif (LMg-K group); it includes shoshonites to rare rhyolites, with latites and trachytes being the most widespread products. The second group consists of potassic rocks (HMg-K group, K₂O/Na₂O between 1.0 and 1.8) occurring in both southern Serbia (Cer and Slavujevci) and Macedonia (Djuristhe, near Sveti Nikole). The third group, present only in Macedonia, consists of ultrapotassic rocks (UK group, K₂O/Na₂O >1.8, Mg# >71) classified as UK shoshonites, UK latites and UK phonotephrites; overall, they show a "Roman Province type" affinity (Group III of Foley, Venturelli, Green, Toscani, Earth Sci Rev 24:81-134, 1987). Geochemically, the studied rocks exhibit strong enrichment in LILE, Th and Pb, as well as relative depletion in Ta-Nb and Hf; such signatures are typical of magmas generated in convergent geotectonic settings. In the HMg-K and UK rocks, Sr and Nd isotopic ratios vary from 0.70768 to 0.71040, and 0.51243 to 0.512149, respectively. The rocks of the LMg-K group show relatively limited Sr and Nd isotope variations (0.7087-0.7093 and 0.51233-0.51229), which correlate with a decrease in MgO and increase in SiO₂ contents. The geochemical features of the LMg-K volcanic rocks indicate that their evolution was mainly driven by fractional crystallization coupled with contamination by feldspar-rich crustal materials. In contrast, the HMg-K and UK rocks have not been significantly modified by crustal contamination, and their geochemical features are considered to reflect lithospheric mantle heterogeneity acquired during the subduction of the

Western Vardar Ocean and the Apulian plate. The metasomatizing agent was apparently more enriched in Zr, Th, Ta and Ce than in fluid-mobile elements, such as Pb and Cs, suggesting that it was characterized by a high melt/fluid ratio. The potassic and ultrapotassic magmatic activity developed in response to the Pliocene–Pleistocene extension in the Vardar Zone, in turn related to the opposite propagation of extension in the Aegean and Pannonian basins (respectively SW and NE).

Introduction

Starting in the Jurassic–Cretaceous, the central and northeastern part of the Balkan Peninsula was affected by NEdirected subduction, with both SW (Hellenides-Dinarides) and NE (Balkans) migration of the conjugate orogenic fronts and intervening later extension (Doglioni et al. 1996). As a result of this complex geotectonic evolution, a calc-alkaline and shoshonitic volcanic arc developed in the Late Paleogene–Middle Miocene (Fig. 1; e.g., Pamić et al. 1998). Eruptive products form a NW–SE belt extending for over 1,600 km from southeastern Austria to northwest-



Fig. 1 Simplified tectonic map of the Eastern Mediterranean region (modified after Dumurdzanov et al. 2004) with the location of Eocene–Oligocene, Miocene–Pliocene and active volcanic belts. *SPFZ* Scutari-Peć Fault Zone, *KFZ* Kefalonia Fault Zone, *NAFZ* North Anatolian Fault Zone

ern Turkey, traversing the Balkan Peninsula (Harkovska et al. 1989; Kovács et al. 2007). The main phase of orogenic magmatism was locally followed by scattered potassic to ultrapotassic volcanism (Cvetković et al. 2004; 2007; Yanev 2003; Yanev et al. 2003; Agostini et al. 2007) that developed from Serbia to Turkey. From the Late Miocene to Pleistocene, in the region between southern Serbia and Macedonia, shoshonitic to ultrapotassic volcanism formed a discontinuous belt starting just south of the Scutari-Peć fault zone, regarded as the northern limit of the extensional area associated with the Hellenic subduction system (Boccaletti et al. 1974).

The ultrapotassic volcanism of southern Serbia and Macedonia has been the object of several recent studies mainly addressing its geochemical and petrological characteristics (Boev and Lepitkova 1991; Cvetković et al. 2004, 2007; Altherr et al. 2004; Prelević et al. 2001, 2005, 2007; Boev and Yanev 2001; Yanev et al. 2003). This paper describes the temporal and spatial evolution of the potassic and ultrapotassic rocks erupted since the Late Miocene to Pleistocene from the Scutari-Peć fault zone down to the southernmost part of Macedonia, to the volcanic Kozuf-Voras Massif at the border between Macedonia and Greece. New geochronological, geochemical and isotopic data, along with literature data, are used to discuss the petrogenesis of these rocks in the context of the geodynamic evolution of the area.

Geodynamic framework

Southern Serbia and Macedonia are located at the core of the Dinarides orogen, where the oceanic sliver of the Vardar unit is sandwiched between the continental basement rocks of the upper Western European plate (Serbo-Macedonian metamorphic rocks) and the lower Apulia plate (Pelagonian unit, e.g., Carminati et al. 2004). The subduction record goes back to at least the Paleocene (Carminati et al. 2004), but evidence from the Cretaceous supports an earlier onset. Collision took place in the Paleocene–Eocene (Pamić et al. 1998). Southern Serbia and Macedonia are still in the hanging wall of an active subduction zone, although the present subduction rate along the eastern coast of the Adriatic Sea seems to be very slow (e.g., Battaglia et al. 2004). Therefore, the calcalkaline-shoshonitic magmatic rocks occurring in the study area are compatible with the aforementioned long history of subduction-collision. However, widespread extension, with the formation of NNW-trending grabens, affected the area starting in the Paleogene (Dumurdzanov et al. 2004); southern Serbia and Macedonia form an area in which the southern margin of the eastward-propagating Pannonian extension and the northwestward margin of the southwestward-propagating Aegean extension overlap in time and

Fig. 2 Sketch map of Republic of Macedonia showing locations of the volcanic centers studied. In parentheses the K–Ar ages (Ma) of volcanic rocks (this work and literature data)



space. A number of Paleogene and Miocene to Pleistocene grabens, filled by coarse-grained siliciclastic sediments and volcanic rocks, accomplished the two independent and coexisting extensional settings. Due to its proximity, the



Fig. 3 Total alkali-silica classification diagram. T&B tephrite and basanite, *PTP* phonotephrite, *Sh* shoshonite, *L* latite, *Tr* trachyte, *R* rhyolite. The dashed line between alkaline and subalkaline rocks of Irvine and Baragar (1971) is also plotted. Literature data for this diagram and the following ones are from Altherr et al. (2004) and Cvetković et al. (2004)

study area was possibly more affected by Aegean tectonics, as also suggested by the southwestward migration of magmatism from the Paleogene to the Pliocene–Pleistocene in Macedonia and northern Greece.



Fig. 4 K_2O/Na_2O vs K_2O (wt%) diagram. Ultrapotassic rocks are defined following the chemical criteria of Foley et al. (1987). The line separating sodic and potassic rocks was drawn according to Le Maitre (1989). Symbols as in Fig. 3

Table 1 Major and trace element concentrations and Sr-Nd isotope ratios of selected samples

Sample Locality rock type group	KZ 3 M. Chuka Trachyte LMg-K	KZ 5 Mt Porta Latite LMg-K	KZ 6 Mt Porta Latite LMg-K	KZ 7 Mt Porta Shoshonite LMg-K	KZ 11 Sokol Trachyte LMg-K	KZ 12 Golubec Shoshonite LMg-K	KZ 13 Golubec Trachyte LMg-K	KZ 15 Vasov Grad Latite LMg-K	KZ 16 Pulevatz Latite LMg-K	KZ 17 Pulevatz Latite LMg-K
Major element	s									
SiO ₂ (wt%)	64.43	58.54	61.66	51.88	61.58	52.54	62.67	61.36	56.87	57.19
TiO ₂	0.37	0.68	0.57	1.16	0.64	0.92	0.54	0.62	0.66	0.66
Al_2O_3	18.15	18.36	17.63	19.25	17.74	18.06	17.64	17.35	17.44	17.15
$Fe_2O_3(t)$	3.06	5.23	4.56	7.84	4.43	8.85	4.50	4.87	5.38	5.31
FeO	-	_	_	_	-	_	-	-	_	-
MnO	0.04	0.10	0.09	0.11	0.06	0.16	0.10	0.09	0.12	0.07
MgO	1.43	2.85	2.40	4.69	2.38	3.85	1.90	2.87	3.17	3.04
CaO	3.59	5.22	4.82	7.62	3.83	7.55	4.33	5.40	5.56	5.79
Na ₂ O	4.32	3.53	3.57	2.89	3.29	3.18	3.46	3.53	3.58	3.45
K ₂ O	3.99	3.54	3.81	2.86	3.68	2.38	3.86	3.67	5.28	4.78
P_2O_5	0.10	0.30	0.26	0.49	0.26	0.49	0.25	0.28	0.43	0.43
LOI	0.79	1.30	1.23	1.33	1.94	1.99	1.33	0.57	1.13	2.71
Total	100.27	99.65	100.60	100.12	99.83	99.97	100.58	100.61	99.62	100.58
Mg#	57.3	59.5	58.6	60.9	60.7	53.1	54.8	61.3	61.3	60.7
Trace elements	5									
Li (ppm)	—	_	-	—	—	-	—	—	_	_
Be	3.8	4.3	4.3	3.5	3.5	6.4	4.1	4.3	7.5	6.1
Sc	13	18	15	20	17	23	14	17	19	18
V	46	102	86	179	97	162	81	97	141	127
Cr	24	15	11	19	12	60	13	28	27	24
Со	7	14	12	24	13	20	11	13	16	15
Ni	17	12	12	19	13	26	10	16	16	18
Cu	10	16	16	32	14	33	13	16	75	38
Ga	18.4	19	18	18	17	20	17	17	19	18
Rb	129	109	150	97	121	74	130	120	198	173
Sr	1284	1156	1169	1359	1192	1342	1154	1270	1143	1376
Y	11.5	21.1	18.3	22.6	17.5	30.4	15.9	18.5	28.1	22.7
Zr	81	94	76	160	76	37	45	77	295	33
Nb	9	12	11	13	11	14	11	11	20	18
Cs	11	9.0	12.7	7.2	7.6	6.5	9.4	9.0	5.0	12.9
Ba	2007	1533	1414	1530	1656	1099	1684	1591	1743	1735
La	58	73	71	60	63	49	57	69	85	79
Ce	102	130	132	118	119	109	112	131	159	156
Pr	10.6	15.3	14.3	14.4	13.6	16.0	12.3	14.5	18.1	17.8
Nd	35	53	49	54	49	67	44	52	65	64
Sm	4.7	7.8	6.8	8.5	6.8	12.0	6.4	7.6	10.5	10.3
Eu	0.26	1.51	1.13	1.57	0.98	2.32	1.09	0.96	1.69	1.56
Gd	3.6	5.6	4.7	6.2	5.1	8.0	4.7	5.6	8.0	7.1
Tb	0.39	0.69	0.61	0.79	0.63	1.07	0.56	0.69	1.04	0.87
Dy	1.91	3.6	3.10	4.1	3.2	5.6	2.84	3.3	5.2	4.4
Но	0.37	0.72	0.57	0.78	0.60	1.02	0.53	0.63	0.97	0.78
Er	1.02	1.93	1.66	2.12	1.56	2.71	1.47	1.61	2.49	1.98
Tm	0.15	0.30	0.25	0.30	0.21	0.39	0.22	0.25	0.35	0.26
Yb	0.95	1.82	1.65	1.93	1.49	2.51	1.30	1.59	2.30	1.63
Lu	0.15	0.27	0.25	0.28	0.22	0.34	0.19	0.22	0.32	0.21
Hf	2.50	2.90	2.40	4.4	2.60	1.80	1.70	2.30	8.2	1.32
Та	0.63	0.89	0.87	0.75	0.80	0.67	0.85	0.84	1.33	1.21
Tl	0.90	1.10	1.0	0.70	0.80	0.40	0.70	0.90	1.20	2.0
Pb	85	60	62	38	55	43	59	71	79	96
Th	24.9	28.8	31.0	17.4	25.7	8.9	25.5	30.0	54	41.0
U	7.8	8.0	8.0	3.7	7.5	3.3	7.4	8.6	16.0	8.7
°'Sr/ ⁸⁰ Sr ¹⁴³ Nd/ ¹⁴⁴ Nd	0.709162 0.512298		0.708868 0.512320	0.708706 0.512334				0.709280 0.512304	0.709318 0.512300	

- not determined

KZ 18 Staravina Trachyte LMg-K	KZ 22 Gradesnica Trachyte HMg-K	GU 5 Djurishte Latite HMg-K	MN 11 Djurishte Latite HMg-K	CER1 Cer Shoshonite HMg-K	SLAV Slavujevci Shoshonite HMg-K	EB 2 Jezevo Brdo Ph-tephrite HMg-K	GR 3 Gradishte Latite UK	K 01 K. Krasta Shoshonite UK	MN 13 M. Negor. Ph-tephrite UK	MN 9 Kishino Ph-tephrite UK
Maior eler	nents									
64 01	60.01	55 27	55.83	52 73	50.58	46 36	53 79	50.87	49 62	46 69
0.29	0.54	0.94	0.92	1.23	0.99	2.19	1.15	1.47	1.16	1.23
17.72	16.99	14.70	15.19	14.59	14.86	14.14	12.81	11.39	12.25	13.05
2.31	4.72	2.49	5.72	4.46	4.41	4.40	2.25	2.58	7.24	8.71
_	_	2.87	_	2.47	2.42	2.90	3.70	4.13	_	_
0.09	0.09	0.11	0.11	0.13	0.13	0.12	0.12	0.11	0.12	0.15
0.43	2.17	6.40	5.51	7.05	7.66	7.92	8.22	9.71	9.95	9.04
1.63	4.89	6.18	6.08	7.38	9.90	8.82	6.26	8.91	7.80	10.0
4.17	5.61	3.57	3.71	3.30	3.49	2.08	2.86	1.49	2.35	2.78
6.90	5.06	5.29	5.41	4.21	3.86	6.11	5.98	6.07	7.14	5.50
0.0	0.32	0.99	1.09	1.20	0.95	1.50	0.97	1.33	1.65	1.62
3.39	0.99	0.57	0.68	1.71	1.69	2.37	0.83	2.88	0.55	1.03
100.94	101.39	99.38	100.25	100.46	100.94	98.91	98.94	100.94	99.83	99.80
34.8	56.9	75.2	72.2	71.8	73.8	73.0	77.7	77.9	78.2	73.0
Trace elem	nents									
-	_	12.86	-	13.93	12.42	9.03	10.03	12.58	-	-
24.0	7.9	8.1	9.19	8.42	6.7	4.45	9.57	8.65	10.59	7.91
10	16	18	18	19	20	23	19	25	24	28
34	100	125	111	159	144	184	131	182	168	195
3	23	227	231	314	184	1/3	383	420	407	263
3	12	23	24 124	31	29	32	29	34 121	30	34
4	12	132	134	194	139	140	243	131	233	98
0	10	-		44	20	-	17	15	15	-
425	19	220	233	10.0	360	388	284	15	13	228
768	1200	13/1	1340	1069	1281	2104	1506	1041	1115	1311
23.4	24.0	24 56	24 52	27.97	28.2	26.12	28.17	22 32	25.36	29 79
465	188	321	312	330	20.2	415	20.17 497	458	319	342
44	20	24.0	25	26	22	30.0	22	25	20	25
84.1	19.7	9.9	9.3	15.5	12.9	10.1	6.3	14.7	24.9	14.9
316	1562	2500	2409	2312	1771	4163	2668	2268	2545	2263
149	84	67	68	48	59	116	99	54	37	56
233	160	130	130	96	119	243	223	126	81	120
20.8	17.3	14.9	15.0	11.8	14.5	28.9	29.4	17.1	10.7	15.4
60	60	54.6	55.4	45.6	54.5	109.3	119.1	70.2	43.8	62.6
7.7	9.3	8.8	8.7	8.0	9.0	15.7	16.5	10.6	8.5	11.3
1.21	1.47	1.54	1.50	1.95	2.18	2.32	2.93	1.63	1.37	2.28
5.5	6.7	6.7	6.5	6.1	6.6	10.5	9.6	7.0	6.6	8.5
0.68	0.81	0.84	0.83	0.85	0.87	1.09	1.10	0.82	0.89	1.09
3.5	4.4	4.3	4.3	4.6	4.8	5.2	5.3	4.3	4.7	5.7
0.69	0.80	0.86	0.81	0.95	0.96	0.92	0.99	0.75	0.86	1.03
2.07	2.14	2.20	2.17	2.50	2.57	2.21	2.46	1.86	2.15	2.50
0.35	0.33	0.34	0.34	0.38	0.39	0.31	0.33	0.28	0.33	0.38
2.41	2.09	2.08	2.04	2.29	2.34	1.92	2.26	1.69	1.94	2.22
0.36	0.30	0.30	0.32	0.33	0.36	0.27	0.32	0.23	0.29	0.33
13.2	5.5	8.0	8.0	8.4	7.1	10.6	12.8	11.8	8.3	8.7
1.95	1.42	1.35	1.39	1.45	1.40	1.39	1.10	1.23	1.04	1.47
3.90	2.10	0.59	0.27	1.31	0.59	1.46	1.82	2.31	2.07	1.86
128	81	46	55 21 0	30	31	26	54 (8.2	19	40	30 26 2
165.0	30 15-2	31.2 8.4	51.9 9.12	23.7	26.1	25.5	68.2 7.82	45.1	23.8	26.3
4/.0	13.3	0.4 0.71	0.13 0.709150	/.48	8.23	4./3	1.85	/.3	8.38 0.71	/.4/
	0.708938	0.71	0.708150	0.71	0.71	0.708093	0.710404	0.710090	0.71	
	0.512517	0.31	0.51	0.31	0.31	0.312337	0.312147	0.012224	0.51	

Analytical methods

K-Ar ages were determined on whole rock samples at ATOMKI, in Debrecen (Hungary). The groundmass was separated using standard techniques and purified by handpicking under a stereomicroscope. About 500 mg of sample were spiked with 38 Ar for the determination of 40 Ar. The mass spectrometer, operating in static mode, was cleaned with conventional getter material before introducing the gas sample. Interlaboratory standards HD-B1, GL-O, LP-6 and Asia 1/65 were used for calibration. Ages were calculated according to the decay constants of Steiger and Jäger (1977). All analytical uncertainties are reported as single standard deviations. Potassium concentrations were determined by flame photometry, adding 100 mg/l of Na and Li to ~100 mg dissolved sample. Major and trace element analyses were performed at the Pisa University's Dipartimento di Scienze della Terra. Major elements were determined by X-ray fluorescence on an ARL 9400 XP+ spectrometer using Li₂B₄O₇ glass disks. Estimated precision (relative standard deviation, RSD) is about 1% for SiO₂ and about 2% for the other major elements except those with low concentrations (approximately <0.50 wt%), for which the absolute standard deviation is about $\pm 0.01\%$. Loss on ignition (LOI) was determined by gravimetry at 1,000°C after preheating at 110°C. The concentrations of a set of 35 trace elements were determined by inductively coupled plasma-mass spectrometry (VG PQII Plus). Sample powders were dissolved in PFA vessels on a hot plate at about 120°C using HF + HNO₃. The sample solutions, spiked with Rh, Re and Bi as internal standards, were measured by external calibration using international reference materials of basaltic composition. Analytical precision (assessed through replicate analyses) is between 2 and 5% RSD, except for Gd, Tm, Be, Sc, Pb (6-8% RSD).

Sr and Nd isotope compositions were measured with a Finnigan MAT 262 multicollector mass spectrometer at

the Istituto di geoscienze e Georisorse, CNR, Pisa. Conventional ion exchange methods were used for Sr and Nd separations. Measured ⁸⁷Sr/⁸⁶Sr ratios were normalized to ⁸⁶Sr/⁸⁸Sr=0.1194, ¹⁴³Nd/¹⁴⁴Nd ratios to ¹⁴⁶Nd/¹⁴⁴Nd=0.7219. During the collection of isotopic data for this study, replicate measurements of NIST SRM 987 (SrCO₃) and La Jolla standards gave values of 0.710243± 13 (2SD, N=20) for ⁸⁷Sr/⁸⁶Sr and 0.511848±7 (2SD, N= 30) for ¹⁴³Nd/¹⁴⁴Nd.

Sample location

The studied volcanic rocks crop out in the Vardar Zone, south of the Scutari-Peć transverse fault zone considered to be the northern limit of Aegean extension (Kissel et al. 1995). Figure 2 reports the locations of the studied volcanic centres. The volume of emitted products is generally limited; they form small, thin, lava flows, which are in some cases connected with the feeder dykes, as in the Kureshnichka Krasta and Jezevo Brdo areas, where eroded remnants of vents (necks) are also present. The volcanic cover at Mlado Nagorichane is 10 to 30 m thick and has the greatest extension, covering an area of about 15 km². It consists of several superimposed lava flows separated by thin scoria levels. In addition, volcanics from Cer and Slavujevci in southern Serbia were sampled and analyzed. Noteworthy, the lavas of Jezevo Brdo and Kureshnichka Krasta crop out in the area of the NW-trending active faults of Skopje and Lakavica (Dumurdzanov et al. 2005), from which they were erupted.

The large Kozuf Massif (named Voras Massif in Greece) is the southernmost volcanic area sampled in this study. It comprises a cluster of domes, pyroclastics and subordinate lava flows; 22 samples from the Macedonian part of this massif were analyzed.

Fig. 5 CaO vs SiO₂ (a), and CaO vs Al_2O_3 (b) (wt%) diagrams. Boundary lines between Group I and Group III ultrapotassic rocks are slightly modified from Foley et al. (1987). Symbols as in Fig. 3



Results

Petrography and mineral chemistry

The Neogene volcanic rocks from the Vardar Zone constitute a complex association with a potassic (K) to ultrapotassic (UK, *sensu* Foley et al. 1987) alkaline and transitional signature (Figs. 3 and 4). For their classification we mainly followed the IUGS criteria, as reported in Le Maitre (2002).

The rocks of this study were divided into three groups: (1) ultrapotassic rocks (shoshonite, phonotephrite, latite); 2) high-Mg potassic rocks (HMg-K; shoshonite, phonoteph-

rite, latite with MgO>5 wt%); 3) low-Mg potassic rocks from Kozuf-Voras Massif (LMg-K; shoshonite, latite, trachyte, rhyolite with MgO<5 wt%).

A synopsis of petrographical and mineralogical features of the studied rocks are reported in the following paragraphs.

LMg-K Group These rocks are exposed only in the southern part of the Vardar Zone, in the Kozuf-Voras Massif. They are mildly alkaline to transitional (Fig. 3), with potassic affinity and a K_2O/Na_2O ratio generally less than 1.8 (Fig. 4, Table 1). These rocks are considered to form a shoshonitic association; they are silica-oversaturated, except samples KZ7 and KZ14, which are olivine- and hypersthene-

Table 2 Summary of petrological and chemical characteristics of the studied rocks

Locality	Rock type	Group	Texture	Phenocrysts/ microphenocryst	Groundmass
Cer	Shoshonite	HMg-K	Porphyritic	Ol, Cpx, Phl	Na-San, Pl, Ti-Mg, Ilm, Ap
	$K_2O/Na_2O=1.8$		Gm-poikilitic		
Slavujevci	shoshonite	HMg-K	Porphyritic	Ol, Cpx, Phl, Pl	Pl, Mg, Ap, analcime
	$K_2O/Na_2O=1.1$		Gm-poikilitic		
	Ne (norm) 8.3				
M. Nagorichane	phonotephrite	UK	Porphyritic	Ol, Cpx, Lc	Na-San, Ba-Ti Phl, Cpx, Ti-Mg, Ap, Lc altered into zeolite
	K ₂ O/Na ₂ O=2.8-4.5		Gm-poikilitic		
	Ne (norm) 8-11+ Lc				
	(norm) 1.0				
Gradishte	Latite to UK-latite	UK	Porphyritic	Ol, Cpx, Phl	Cpx, Na-San, Phl, Ti-Mg, Ap
	$K_2O/Na_2O=2.0-2.1;$		Gm-trachytic		
	Ne (norm) 0.6-1.7		-		
Djurishte	Latite to shoshonite	HMg-K	Porphyritic	Ol, Cpx, Phl	Cpx, Phl, Na-San, Ti-Mg, Ilm, Ap
	K ₂ O/Na ₂ O=1.4-1.5		Gm-poikilitic		· · · · · · ·
	Ne (norm) 0.3–1.8 or hy 0.8		-		
Jezevo Brdo	Phonotephrite	UK	Microporphyritic	Ol, Cpx, Ti-Phl	Anorth, Pl, Phl, Na-San, Mg- bearing carbonate, Cpx, Lc (with Cpx inclusion)
	$K_2O/Na_2O = 1.9 - 2.9$		Gm-poikilitic		· ·
	Lc (norm) 0.9–6.6		Ĩ		
Kishino	phonotephrite	UK	Porphyritic	Ol, Analcime	Cpx, Ol, Na-San, Anorth, Ti-Phl, Ti-Mg, Ap, Carbonate, zeolite
	$K_2O/Na_2O=2.0$		Gm-microlitic		
	Lc (norm) 5.7				
K. Krasta	Phonotephrite to UK-shoshonite	UK	Porphyritic	Ol, Cpx, Phl	Na-San, Phl, Cpx, Ol, Lc (altered into zeolites+clays), Ti-Mg, Ap
	$K_2O/Na_2O=3.5-5.5$		Gm-poikilitic		
	Ne (norm) 1.5-5.5				
Kozuf	Shoshonites to rhyolites	LMg-K	Porphyritic	Pl, San, Amph, Bt	Pl, San, Bt, Mg
	K ₂ O/Na ₂ O=0.8-1.6		Gm–holohyaline to holocrystalline ipidiomorphic		
	Ne (norm) 0–3; Q 0–21.7				

Ol olivine, *Cpx* clinopyroxene, *Phl* phlogopite, *Amph* amphibole, *San* sanidine, *Anorth* anorthoclase, *Gm* groundmass, *Lc* leucite, *Ne* nepheline, *Mg* magnetite, *Ilm* ilmenite, *Ap* apatite, *Pl* plagioclase, *Bt* biotite

normative, and samples KZ21 and KZ22, which are nepheline-normative. All rocks are porphyritic, with variable amounts of glass in the groundmass; in some cases a porphyritic holohyaline texture is observed. The groundmass is about 50–70 vol.% of the total rock volume and generally contains the same minerals present as phenocrysts. The phenocryst assemblage consists of zoned plagioclase (from labradorite to oligoclase), hornblende and biotite; biotite becomes more abundant in the more evolved rocks. Sanidine occurs sporadically in latites and becomes significantly more abundant in trachytes, where it can locally reach centimeter size. Scattered clinopyroxene is mainly observed in shoshonites and latites. Opaque minerals are represented by Ti-magnetite and rare ilmenite.

HMg-K Group This group shows a shoshonitic affinity, like the Kozuf samples, but with higher alkalinity (Na₂O+K₂O> 7 wt%), a K₂O content generally >4 wt%, and a K₂O/Na₂O ratio of 1.0 to 1.8 (Figs. 3 and 4). The rocks are generally saturated or slightly undersaturated in SiO₂ (highest normative Ne content=8.3%, Slavujevci). In addition, they exhibit relatively high MgO contents with Mg# [100 Mg/(Mg+ Fe²⁺)] >70, in contrast to Mg# <61 of the Kozuf rocks.

These rocks show a porphyritic texture; in shoshonites and latites, the phenocrysts are represented by the associ-



Fig. 6 Al_2O_3 vs TiO₂ (wt%) diagram for phogopites from HMg-K and UK rocks from southern Serbia and Macedonia. Group III (Roman Province-type) and Group I (Lamproites) fields redrawn after Prelević et al. (2005)

 Table 3
 T and P estimates

Locality	Rock type	<i>T</i> (°C)	P (GPa)
M. Nagorichane	Phonotephrite	1,298 (1)	
		1,312 (2)	
Kishino	Phonotephrite	1,259 (1)	
		1,284 (2)	
Jezevo Brdo	Phonotephrite	1,287 (1)	
		1,262 (2)	
		1,144 (3)	0.7
Gradishte	UK-Latite	1,259 (1)	
		1,267 (2)	
		1,298 (3)	1.7
Djurishte	K-Latite	1,189 (1)	
		1,210 (2)	
		1,198 (3)	0.8
K.Krasta	UK-Sho	1,310 (1)	
		1,290 (2)	
		1,278 (3)	1.2

(1) Putirka et al. (2007); (2) Beattie (1993); (3) Putirka et al. (2003)

ation of Mg-rich olivine (Fo₈₀₋₈₈), phlogopite, clinopyroxene and rare andesine plagioclase. The groundmass includes small amounts of recrystallized glass, sanidine, pyroxene, phlogopite, Ti-magnetite and apatite; Djurishte lavas also contain ilmenite. In the literature these rocks have been classified as UK rocks with lamproitic affinity (e.g. Prelević et al. 2005; 2007; Cvetković et al. 2004; Altherr et al. 2004).

UK Group These rocks are highly primitive, with MgO generally >7 wt% and Mg# ranging between 71 and 79. Although the rocks of this group have been considered lamproites, mainly on a mineralogical basis, i.e. the absence of plagioclase (Altherr et al. 2004), or on a chemical basis, i.e. CaO vs MgO diagram (Prelević et al. 2007), we consider them to have a Roman Type (Group III of Foley et al. 1987)-affinity because of their CaO/SiO2 and CaO/ Al_2O_3 ratios (Fig. 5; see following section) and the occurrence of groundmass feldspar in some lavas (i.e. Jezevo Brdo). The majority of studied samples show a porphyritic texture; phenocrysts are characterized by the presence of olivine, clinopyroxene (except Kishino lavas) and phlogopite. Leucite has been found in UK phonotephrites from Mlado Nagorichane. In some samples phenocrysts form glomeroporphyritic aggregates. Olivine, up to 1-2 mm in size, is generally slightly rounded and zoned (e.g. in Kureshnichka Krasta: cores Fo₈₁₋₉₃, rims Fo₇₄₋₈₃). Olivine rims and cracks are locally altered to serpentine minerals. Clinopyroxene has an average grain size of 1 mm and is slightly zoned (e.g. in Gradishte and Kureshnichka Krasta); it has a diopside-augite composition, with Al contents ranging from 0.43 to 4.95 wt% (Jezevo Brdo and Djurishte). Phlogopite, zoned and with Ba-rich cores, is

Table 4 K-Ar data

Locality	Material	K (wt%)	⁴⁰ Ar _{rad} (ccSTP/g)	$^{40}\mathrm{Ar_{rad}}$ %	K–Ar age (Ma) $\pm 1\sigma$
M. Nagorichane	WR	5.46	3.856×10^{-7}	40.4	1.81 ± 0.07
Kishino	WR	3.87	2.221×10^{-7}	24.3	$1.47 {\pm} 0.09$
Jezevo Brdo	WR	5.05	6.369×10^{-7}	56.2	3.24±0.11
Gradishte	WR	4.78	3.161×10^{-7}	32.8	$1.70 {\pm} 0.08$
Djurishte	WR	4.24	5.259×10^{-7}	42.8	3.19±0.12
K. Krasta	WR	4.85	3.848×10^{-7}	27.0	2.04 ± 0.10

WR whole rock

often extensively replaced by a magnetite and clinopyroxene assemblage. It has an Mg# of 67 (rim) to 92 (core); the Al₂O₃ content is relatively high (12.6–15.5 wt%). The latite and shoshonite from Gradishte and Kishino, contain phlogopite crystals with low Ba and Ti (\approx 0.3 wt% and \approx 1.7 wt%, respectively) and relatively high Cr (up to 1 wt% for Gradishte rocks) contents; in contrast, the TiO₂ content of phlogopite from the phonotephrites is high (\approx 10 wt%). Leucite occurs as microphenocrysts, except in the Gradishte UK latites. This phase is sometimes partially or totally replaced by analcime, zeolites and clay minerals.

The groundmass consists of the same minerals occurring as phenocrysts with the addition of feldspars, represented by Na-sanidine and oligoclase. Sanidine locally displays a poikilitic texture and has BaO contents of up to ≈ 5 wt% (Mlado Nagorichane). In Jezevo Brdo, Mlado Nagorichane and Kishino, there are patches of calcite with MgCO₃ contents of around 5–7 mole %.

Table 2 summarizes the main petrographical and chemical features of the studied rocks.

Evidence for the Roman-type affinity of the UK group

The petrogenetic affinity of UK rocks from Macedonia is poorly defined on the basis of chemical parameters alone.

Taking into account the distribution of major elements and, in particular, the CaO/MgO ratio, these rocks are intermediate between those of the Group III (Roman Province type) and Group I (lamproites) of Foley et al. (1987; see also Prelević et al. 2005; 2007). However, based on the relatively high Al₂O₃ content, the molar K₂O/Al₂O₃<0.8 and La content <130 ppm (see below and Table 1), these rocks are collectively considered to have a Roman-type affinity. Their mineralogical composition further supports this petrogenetic attribution. In particular, the conspicuous presence of sodic plagioclase in some lavas (i.e. Jezevo Brdo), Fe-poor sanidine (Fe₂O₃<0.8 wt%) and leucite (Fe₂O₃<1 wt%) together with relatively Al-rich clinopyroxene (typically Al₂O₃>1 wt%) is noteworthy. Lastly, the high Al₂O₃ content (>12 wt%) of phlogopite, a typical primary phase of UK Macedonian rocks, distinguishes these rocks from the lamproite phlogopites (Fig. 6).

Pressure and temperature estimate

Crystallization temperatures were assessed on the basis of the partitioning of Fe and Mg between olivine and the whole rock using the geothermometer of Beattie(1993) and Putirka et al. (2007). Crystallization pressures were determined using the thermobarometer of Putirka et al. (2003), which yields T and P estimates for clinopyroxene in equilibrium with coexisting





liquid. Results are reported in Table 3. The crystallization temperature of mafic phenocrysts in the UK group ranges between 1,300 and 1,250°C. The lowest T was calculated for the latite of Djurishte (about 1,200°C). The clinopyroxene segregation pressure of 1.7 to 0.7 GPa corresponds to depths well within the lithospheric mantle.

K-Ar ages

Table 4 reports six new K–Ar age determinations for UK and HMg-K rocks from Macedonia; they range between 3.24 Ma (Jezevo Brdo) and 1.47 Ma (Kishino; Fig. 2). Our data enlarge the existing data set for the younger magmatic



Fig. 8 Primordial mantlenormalized incompatible element and CI chondritenormalized REE patterns (McDonough and Sun 1995). a and b LMg-K Group; c and d, HMg-K Group; e and f, UK Group



Fig. 9 ¹⁴³Nd/¹⁴⁴Nd vs ⁸⁷Sr/⁸⁶Sr plot for the studied samples. The data for the Roman Magmatic Province (*RMP*; Conticelli et al. 2002) and for the Tuscan lamproites (Conticelli et al. 2002; Peccerillo and Martinotti 2006) are also plotted for comparison. Symbols as in Fig. 3

products occurring in the Vardar Zone, south of the Scutari-Peć fault zone. Moreover, the new determinations modify the K–Ar ages reported by Terzić and Svešnikova (1991) of 9.5 Ma for the lava of Kureshnichka Krasta and of 5.5 Ma for that of Jezevo Brdo, which instead yield Late Pliocene isotopic ages of 2.04 ± 0.10 and 3.24 ± 0.11 Ma, respectively. The Djurishte latite yields an age of 3.19 ± 0.12 Ma, whereas the UK rocks have K–Ar ages of 1.81 ± 0.07 Ma at Mlado Nagorichane, 1.70 ± 0.08 Ma at Gradishte and 1.47 ± 0.09 Ma at Kishino, respectively. The UK group is thus representative of the youngest magmatic activity in the Macedonia volcanic belt. Older ages were determined for the volcanics from Slavujevci (6.57 Ma) and Cer, in southernmost Serbia (3.86 ± 0.17 Ma, Cvetković et al.





2004), as well as in the Kozuf Massif (\approx 6.5 Ma, Kolios et al. 1980; Boev and Lepitkova 1991).

Geochemistry

Major and trace element data are listed in Table 1. The Kozuf samples (LMg-K group) show relatively evolved compositions, as indicated by their low Mg# and low MgO, Ni and Cr concentrations (Fig. 7). They show good negative correlations between SiO₂ and TiO₂, Fe₂O₃, MgO, CaO and P₂O₅, whereas Al₂O₃, Na₂O and K₂O show scattered values. Overall, major element variations are consistent with a fractional crystallization process mainly involving the mafic phases. However, the spread of data, mostly alkali and alumina contents, implies the occurrence of other concomitant processes.

The HMg-K group is characterized by relatively high MgO (5.3–7.6 wt%), Ni and Cr (194–131 and 314–180 ppm, respectively); however, these elements are not correlated with MgO (Fig. 7). SiO₂ varies from 51.6 to 55.8 wt% and shows no significant correlation with other major or trace elements.

Rocks belonging to the UK group have primitive compositions with high MgO (6.9 to 9.9 wt%) and Cr and Ni contents (243–100 and 420–170 ppm, respectively); the decrease in MgO is not coupled with a decrease of compatible elements (Fig. 7). Overall, the UK rocks show high dispersion in the Harker diagrams. SiO₂ contents range from 46 to 54 wt% and are significantly lower than those measured in other UK Mediterranean rocks (see Altherr et al. 2004). Note that the TiO₂ contents of UK rocks range from 1.5 to 1.1 wt%, whereas those in Jezevo Brdo are significantly higher, ranging from 2.3 to 2.5 wt%.

Figure 8 reports the primordial mantle-normalized incompatible element and chondrite-normalized rare earth element (REE) patterns for selected LMg-K, HMg-K and UK rocks. All the investigated samples show arc-type



Fig. 11 Rb/Nb vs K/P plot for studied samples. The diagram also shows the field of evolved feldspathic rocks and the average value of the upper continental crust (Taylor and McLennan 1995). K/P=8 is the average value of HMg-K and UK rocks (excluding LMg-K samples). Symbols as in Fig. 3

incompatible element distributions; they are generally enriched in LILE, with high LILE/HFSE ratios and typical negative Ta, Nb and Ti anomalies, along with a positive Pb spike which decreases from the LMg-K samples to the UK rocks. Moreover, the LMg-K samples are characterized by positive Th and U spikes and negative Hf and Zr anomalies. These anomalies are reduced in the HMg-K rocks and almost absent in the UK rocks. In contrast to other UK Mediterranean rocks, no negative Ba anomaly was observed (Altherr et al. 2004).

The REE patterns show a pronounced and variable enrichment of LREE over HREE, with La_N/Yb_N ranging from 13 (UK rocks from Mlado Nagorichane) to 42 (LMg-K group). The LREE display different degrees of fractionation; the highest degree of fractionation is observed in the LMg-K rocks (La_N/Sm_N from 2.6 to 7.7) and the lowest in the UK group (La_N/Sm_N from 2.7 to 4.6). Note that the UK rocks show convex-upward LREE patterns, which become more pronounced in the most La-enriched samples (Fig. 8). Very similar REE patterns are found among the Circum-Mediterranean ultrapotassic rocks, such as in southeastern Spain (Venturelli et al. 1984a), Tuscany (Peccerillo 2005), northwestern Alps (Venturelli et al. 1984b) and Serbia (Prelević et al. 2005). Convex-upward LREE patterns may be obtained through the metasomatic enrichment of a strongly LREE depleted mantle source.

Heavy REE show the same degree of fractionation in LMg-K rocks (Tb_N/Yb_N from 1.7 to 1.9), whereas they show variable and greater degrees of fractionation in HMg-

K and UK lavas $(Tb_N/Yb_N \text{ from 1.3 and 2.5})$. Overall, the slight U-shape of HREE patterns becomes more pronounced from LMg-K to UK rocks. The negative Eu anomaly is ubiquitous, even in the UK rocks showing no evidence for early plagioclase crystallization.

The investigated rocks are characterized by relatively high ⁸⁷Sr/⁸⁶Sr and low ¹⁴³Nd/¹⁴⁴Nd ratios (Table 1) and fall in the enriched quadrant of the Sr-Nd isotopic diagram (Fig. 9). Sr and Nd isotope variability is quite large and comparable to that observed by Altherr et al. (2004). With respect to K and UK rocks from other Mediterranean occurrences (e.g. SW Spain, Central Italy, Sisco), these rocks are characterized by lower 87Sr/86Sr and higher 143 Nd/ 144 Nd ratios. The HMg-K rocks (87 Sr/ 86 Sr= 0.70768 - 0.70818 and 143 Nd/ 144 Nd=0.51243 - 0.51238) show the highest Nd and lowest Sr isotope ratios. The LMg-K group varies in a narrow range (⁸⁷Sr/⁸⁶Sr=0.7087-0.7093 and 143 Nd/ 144 Nd=0.51233-0.51229), whereas the UK rocks display large isotope variations (87Sr/86Sr= 0.70869-0.7104 and ¹⁴³Nd/¹⁴⁴Nd=0.51237-0.51215), with a negative correlation between Sr and Nd.

Discussion

Role of low pressure processes

The formation of the Tertiary and Quaternary Balkan UK and K province has been ascribed to partial melting of a mantle metasomatized by subducted components during the Mesozoic (Altherr et al. 2004; Boev and Yanev 2001; Yanev et al. 2003 for Macedonian lavas; Prelević et al. 2001; 2005; 2007; Cvetković et al. 2004 for Serbian lavas). Low pressure processes such as fractional crystallization and/or interaction with the continental crust have also been proposed to explain several geochemical features and, in particular, the large range of isotopic variations. The Sr and Nd isotope ratios of the studied rocks with respect to MgO contents are reported in Fig. 10. The most primitive samples, represented by UK group (MgO>8 wt%), almost span the whole range of ¹⁴³Nd/¹⁴⁴Nd and ⁸⁷Sr/⁸⁶Sr values and show no significant correlation with MgO contents. This suggests that fractional crystallization accompanied by crustal assimilation cannot explain the observed variations in UK rocks. In contrast, the LMg-K volcanics show relatively small variations in both Sr and Nd isotope ratios with a significant decrease in MgO contents. This is consistent with the occurrence of fractional crystallization processes, probably combined with low degrees of crustal assimilation. The HMg-K group displays relatively small isotope variability; however, the data points are well correlated with MgO and define a trend line which extends that defined by the LMg-K samples. To better highlight the

interaction of magmas with crustal materials, we plotted data in the K/P vs Rb/Nb diagram (Fig. 11), being K/P and Rb/Nb elemental ratios very sensitive to the upper crustal contamination. The LMg-K samples clearly point towards evolved feldspathic crustal materials, whereas the HMg-K and UK rocks show a strong increase in the Rb/Nb ratio at an almost constant K/P ratio (about 8, i.e. very close to the average value of mantle-derived basalts). Geochemical signatures thus suggest that these rocks were affected by a fractional crystallization process associated with crustal contamination. The most primitive LMg-K lavas display geochemical characteristics very similar to those of HMg-K group cropping out in the investigated area. The relatively small Sr and Nd isotope variations are explained by the buffering effect of the high Sr and Nd contents of the analyzed samples (1,100–1,300 ppm Sr and 35–60 ppm Nd, considerably higher than the average upper continental crust).

In conclusion, the collected geochemical data indicate that rocks from the LMg-K group were significantly affected by crustal contamination, whereas the HMg-K

Fig. 12 ¹⁴³Nd/¹⁴⁴Nd vs Zr/Nb (a), Th/Ta (b), Pb/Ce (c), and Pb/Th (d). MORB values from Niu and O'Hara (2003). Symbols as in Fig. 3

and UK rocks were not; we therefore believe that geochemical features of the latter rocks reflect mantle heterogeneity, as also suggested by Prelević et al. (2005) and Prelević and Foley (2007).

Geochemical evolution of the source region

The UK rocks have a large isotope variability which is not correlated with the emplacement age, the geographic position or, as mentioned earlier, the MgO contents (Fig. 10). The low Al_2O_3 contents, the relatively low CaO/Al_2O_3 ratios (average of 0.57), and the refractory nature of some minerals such as olivine and spinel (Prelević et al. 2005) are consistent with a residual mantle source (Cvetković et al. 2007). P and T estimates (Table 3) are also consistent with a lithospheric mantle source. However, the enrichment in LILE, the radiogenic Sr and non-radiogenic Nd of this source suggest that it has been modified by a metasomatic event (Yanev et al. 2003; Altherr et al. 2004; Prelević et al. 2005; 2007). In the Zr/Nb vs ¹⁴³Nd/¹⁴⁴Nd



diagram (Fig. 12a), the UK rocks show a negative correlation: samples with the highest Zr/Nb ratios have the lowest Nd ratios, whereas both UK and HMg-K group fall in a field characterized by high Nd isotope ratios and small variations in the Zr/Nb ratio. An analogous trend is observed in the ¹⁴³Nd/¹⁴⁴Nd versus Th/Ta diagram (Fig. 12b). The similar behavior of the Zr/Nb and Th/Ta ratios suggests that the enrichment of Zr and Th, coupled with the low ¹⁴³Nd/¹⁴⁴Nd and high ⁸⁷Sr/⁸⁶Sr ratios, is linked to metasomatism of a mantle domain by an agent having a crustal affinity. The nature of this agent can be investigated considering the relative enrichment of fluidmobile elements (Cs and Pb) with respect to relatively fluid-immobile elements (Zr and Th). In the ¹⁴³Nd/¹⁴⁴Nd versus Pb/Ce and Pb/Th diagram (Fig. 12c and d) all the studied samples are relatively depleted in Pb (and Cs, not shown) with respect to Ce and Th (and Zr not shown); the UK latites and shoshonites of the Kureshnichka Krasta and Gradishte are strongly depleted in Pb (and Cs), with Pb/Th ratios lower that those of average MORB and Pb/Ce ratios close to MORB values (Niu and O'Hara 2003). These features suggest that the metasomatized component was essentially a melt rather than a fluid. Note that the UK rocks from Mlado Nagorichane and Jezevo Brdo (phonotephrites, Table 1 and Fig. 12), along with the HMg-K group, define a subset with a relatively homogeneous geochemical imprint. We tentatively interpret these lavas as derived from a mantle domain in which the metasomatized component played a minor role with respect to the source region that fed the volcanism of Kureshnichka Krasta and Gradishte.

Geodynamic implications

The geochemical features of the investigated HMg-K to UK rocks suggest that they originated from a refractory lithospheric mantle (see also Cvetković et al. 2007) metasomatized by an agent essentially consisting of silicate melts derived from a subducted component already depleted in fluids. This implies that the metasomatic episode occurred during a waning phase of the subduction process. The activation of a relatively shallow lithospheric source is linked to the extensional process affecting the region at least since the Oligocene.

Volcanic activity (Fig. 13) was probably the result of three concomitant processes: (1) Dinaric subduction and extension, active since at least the Paleogene (Dumurdzanov et al. 2005); (2) the northwestern extension of the Aegean Sea rift; and (3) the southern termination of Pannonian Basin rifting in the hanging wall of the eastward retreating Carpathian subduction zone. The rifting episodes led to the collapse and stretching of the thickened lithosphere, allowing the uplift of the previously metasomatized underlying mantle. This opposite movement created favorable conditions for the



Fig. 13 Sketch map showing the geodynamic setting of the Aegean-Balkanic region. The study area (*box*) is located in the hangingwall of the Mesozoic to present-day Dinaric subduction (*small black arrows*), which underwent widespread late and syn-subduction Cenozoic extension associated with the Aegean rift and, to the north, with the Pannonian rift. The origin of the HMg-K und UK volcanic rocks of Macedonia is consistent with a subduction setting and coeval extensional tectonics

generation of ultrapotassic to potassic magmas, firstly (Oligocene–Early Miocene) in central Serbia and lately (Late Miocene to Pleistocene) in southernmost Serbia and Macedonia. We believe that this subduction is the Late Cretaceous northeastward subduction of the Western Vardar Ocean, with the Apulian plate under the edge of the Eurasian one (Carminati et al. 2004). The remnants of the subducted Apulian plate are visible on tomographic profiles (Piromallo and Morelli 2003; Bennet et al. 2008). Magmatism possibly ceased due to a decrease in the subduction rate caused by the docking of the thicker Adriatic-Apulian lithosphere at the Dinaric subduction trench.

Summary and conclusions

In Macedonia, Late Miocene to Pleistocene potassic and ultrapotassic volcanic rocks are exposed in the Vardar zone, from south of Scutari-Peć fault zone to the Greek border. The erupted products show a subduction-related signature (enrichment in LILE, high LILE/HFSE ratios and typical negative Ta, Nb and Ti anomalies) but variable potassiumenrichment and petrogenetic affinity. Three main rock groups have been identified: the first two groups display shoshonitic features and are represented by: (1) LMg-K rocks forming the Kozuf Massif which extends also in Greece and (2) HMg-K scattered small volcanic centers extending from the Serbia-Macedonia border to central Macedonia, which are silica saturated or slightly undersaturated lavas with more alkaline and K-rich character. The last group is formed by ultrapotassic, primitive, frequently leucite-bearing lavas, which represent the youngest volcanism of the area and range in age from 1.8 to 1.5 Ma. Based on chemical and mineralogical composition these rocks are inferred to have a petrogenetic affinity akin to the Group III (Roman Province type) ultrapotassic rocks of Foley et al. (1987). All the studied rocks are characterized by relatively high ⁸⁷Sr/⁸⁶Sr and low ¹⁴³Nd/¹⁴⁴Nd ratios and fall in the enriched quadrant of the Sr-Nd isotopic diagram. Isotopic and geochemical data reveal that LMg-K lavas were affected by upper crustal contamination processes, whereas HMg-K and UK groups display variations mainly reflecting geochemical heterogeneity of the mantle source; these rocks are interpreted as the result of partial melting of a refractory source whose geochemical composition was modified by the addition of a metasomatic agent essentially constituted by melts as stressed by the relative depletion of fluid mobile with respect to fluid immobile elements (i.e. Cs and Pb versus Zr and Th). The heterogeneous nature of the refractory mantle source as well as geothermometric calculations support the hypothesis that this source was localized in the lithospheric mantle.

The eruption of the HMg-K and UK rocks is related to rifting episodes that occurred in the region since at least the Oligocene. The overall arc-type geochemical signature of the Macedonia Mio-Pleistocene lavas confirms their relation with Dinaric subduction, active since the Late Eocene– Oligocene. The later stretching of this area is considered the result of two concurrent and coeval processes, the dominant southwestern extension of the Aegean Sea rift and, to the north, the Pannonian rifting due to the retreat of the Carpathian subduction zone.

Acknowledgments This research was financially supported by PRIN (Cofin-MIUR) and by CNR grants. We thank *Dejan Prelević* and *Hilary Downes* for their constructive reviews; the editorial handling of *Johann Raith* was greatly appreciated.

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