

# THE KAMAFUGITE-CARBONATITE ASSOCIATION IN THE ALTO PARANAÍBA IGNEOUS PROVINCE (APIP) SOUTHEASTERN BRAZIL

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**ABSTRACT** The Late-Cretaceous Alto Paranaíba Igneous Province (APIP) comprises a variety of ultrapotassic rock-types, including kimberlites, lamproites and large volumes of kamafugites, in addition to a number of carbonatite-bearing plutonic alkaline complexes. Phlogopite-picrites are ultramafic fine-grained rocks typically composed of olivine phenocrysts set in a groundmass of phlogopite, carbonate, perovskite, apatite and chromite. They occur as dyke swarms in the carbonatite complexes, but are also scattered throughout the Province. The phlogopite-picrites represent the peralkaline, ultrapotassic, carbonate-rich, silicate magmas parental to the carbonatite-bearing complexes, and have strong chemical affinity with kamafugites. Together with petrographic similarities observed between silicate rocks from the carbonatite complexes and xenoliths occurring in APIP kamafugites, this provides a strong link between kamafugitic and carbonatitic magmatism in the Province.

**Keywords:** kamafugite, carbonatite, phlogopite-picrite, ultrapotassic, APIP

## INTRODUCTION AND GEOLOGICAL SETTING

Voluminous magmatism occurred in central and southern Brazil, from Early Cretaceous until Eocene times. This includes the extensive Early-Cretaceous Paraná continental flood basalt (CFB) province and a number of Early-Cretaceous to Eocene Alkaline Igneous Provinces that surround the Paraná Basin. All of this magmatism has been associated with the thermal and/or chemical influence of mantle-plumes (Tristan and Trindade) impacting on the base of the continental lithosphere (e.g. Herz 1977, Toyoda *et al.* 1994, Gibson *et al.* 1995a,b, 1997, VanDecar *et al.* 1995; Thompson *et al.* 1998).

The products of the intense Late-Cretaceous alkaline magmatism at the northeastern border of the Paraná Basin (Fig.1), in south-east Brazil, are collectively known as the Alto Paranaíba Igneous Province (APIP, Gibson *et al.* 1995b). The magma types include kimberlites, madupitic olivine lamproites and kamafugites, in addition to large intrusive carbonatite-bearing alkaline complexes. Rocks of kamafugitic affiliation are by far the most abundant in the APIP. Madupitic olivine lamproites are restricted to the NW portion of the province and diamondiferous (Gonzaga and Tompkins 1991) kimberlites, containing xenoliths of garnet lherzolite (Leonardos *et al.* 1993, Carvalho and Leonardos 1995) occur at Três Ranchos (Goiás State), in the north of APIP.

## ALTO PARANAÍBA IGNEOUS PROVINCE CARBONATITE COMPLEXES

A number of relatively large (up to 65 km<sup>2</sup>), intrusive, carbonatite-bearing ultramafic complexes are found in the province. These occurrences comprise Catalão I and Catalão II in southern Goiás State, and Serra Negra, Salitre I, Salitre II, Araxá and Tapira in western Minas Gerais State (Fig. 1). The complexes are multi-stage intrusions emplaced into Late-Proterozoic metamorphic rocks of the Brasília mobile belt, which are typically deformed into dome structures.

The silicate-rock association in the carbonatite complexes includes dunites, wehrlites, clinopyroxenites, bebedourites, phoscorites and syenites. Rare melilitolite (uncompahgrite) has been described from the Tapira Complex (Guimarães *et al.* 1980, Brod 1999). A striking feature of the APIP carbonatite complexes is the absence of nepheline-rich rocks, such as those of the ijolite series, which are commonly associated with many carbonatite complexes worldwide. Instead, a major primary silicate rock-type in the APIP carbonatite complexes is bebedourite, that varies from fine- to coarse-grained (locally pegmatitic) and is sometimes strongly foliated. It consists mainly of diopsidic pyroxene with variable amounts of phlogopite, perovskite, apatite, magnetite, melanite and sphene. Modal variations in the essential mineral phases can produce facies rich in olivine (dunites, wehrlites), perovskite, magnetite or apatite. The Serra do Salitre (Salitre Mountains), within the APIP, is the type area for this rock, as defined by Tröger (1928).

The petrographic and chemical characteristics of syenitic rocks associated with the APIP carbonatite complexes indicate an ultrapotassic affinity. In thin section, altered patches composed of a

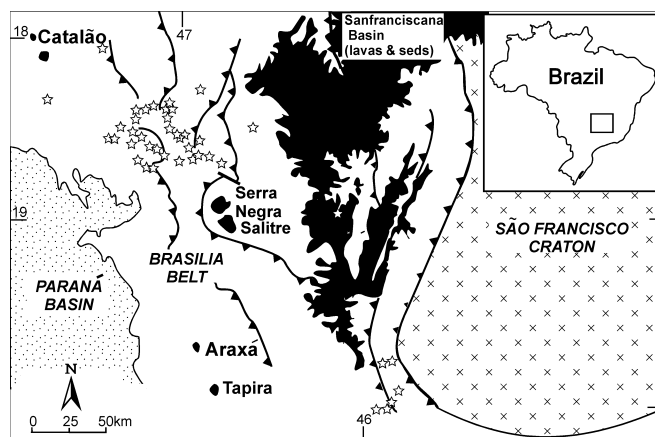


Figure 1-Location and geological setting of the Alto Paranaíba Igneous Province. The carbonatite-bearing complexes of Catalão, Serra Negra, Salitre, Araxá and Tapira are indicated individually. Smaller occurrences of APIP ultrapotassic rocks are shown by stars. The Sanfranciscana Basin includes both the Mata da Corda volcanics (not individualized) and the Areado sediments. Adapted from Gibson *et al.* (1995b).

mixture of carbonate, sericite and clay minerals often resemble the outlines of leucite crystals, whereas others resemble nepheline or kalsilite. An XRD reconnaissance of Tapira syenites yielded a set of peaks consistent with the presence of leucite or analcime. In samples from Salitre the XRD study indicated the presence of kalsilite (Brod 1999). These rocks are currently undergoing microprobe investigation.

Sovite and beforosite are dominant among the carbonatites, with subordinate silico-carbonatite (e.g. Catalão II, Machado Jr. 1992).

Ultramafic dykes are intimately associated with all APIP carbonatite complexes. The dykes are usually a few centimetres or tens of centimetres thick, rarely reaching more than one metre. These have been traditionally described as lamprophyres, on the basis of the abundant olivine and subordinate phlogopite phenocrysts. On other occasions they have been mistaken for metasomatic glimmerites, because of the high modal percentages of phlogopite and carbonate. Gibson *et al.* (1995b) and Brod (1999) demonstrated the magmatic character of these rocks. Flow differentiation, flow structures, porphyritic textures with euhedral phenocrysts, discordant tabular (dyke) shape of the intrusions, and carbonate ocelli are common primary magmatic features. Chilled margins are rare, but have been observed in dykes emplaced away from the carbonatite complexes.

Gibson *et al.* (1995b) stressed that such rocks do not fit the established nomenclature, and proposed the name phlogopite-picrite. No suitable alternative name appeared in the IUGS guidelines issued shortly after by Woolley *et al.* (1996) for the classification of alkaline potassic rocks. Therefore, the nomenclature proposed by Gibson *et al.*

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(1995b) is adopted here; the name phlogopite-picrite is used for dyke-rocks essentially composed of phenocrysts of olivine and subordinate phlogopite, set in a groundmass of phlogopite, chromite, magnetite, perovskite, apatite and carbonate. The MgO contents of many samples certainly fit the latest IUGS definition of picrite (Le Bas, 2000). Although phlogopite-picrite dykes are typically associated with the carbonatite complexes, they are also found scattered throughout the province, with no obvious spatial relationship to carbonatites. Besides the obvious potassic character of their mineralogical assemblage, phlogopite-picrites also have textural similarities to other types of ultrapotassic rocks. Olivine phenocrysts may be coated by aggregates of equant small crystals of perovskite and oxides (chromite). This feature is similar to that described from kimberlites (e.g. Reid *et al.* 1975, Dawson and Hawthorne 1973), or by subhedral laths of phlogopite, similarly to olivine from lamproites (Mitchell and Bergman 1991). Mantling of olivine by phlogopite laths has also been reported by Gibson *et al.* (1994) from the Serra do Bueno kamafugitic diatreme, in the APIP. Phlogopite phenocrysts may be zoned towards tetra-ferriphlogopite, in samples where carbonate is modally abundant in the groundmass, a feature also observed in orangeites (Mitchell 1995). Carbonate-rich zones in the groundmass of phlogopite-picrite dykes may locally evolve to carbonate "pockets" of irregular or globular shape, suggesting that immiscibility of carbonate-rich liquid from the phlogopite-picrite magma may have occurred as early as during the formation of olivine. Carbonate immiscibility from unevolved silicate liquids has been documented from kimberlites and ultramafic lamprophyres (e.g. Dawson and Hawthorne, 1973; Pearce and Leng, 1996).

**LINKS WITH ALTO PARANAÍBA IGNEOUS PROVINCE KAMAFUGITES** The chemical and mineralogical affiliation of the Mata da Corda volcanic and pyroclastic rocks with the kamafugite series are now firmly established (Seer and Moraes 1988, Seer *et al.* 1989, Sgarbi and Valença 1993, 1994, Sgarbi and Gaspar 1995, Gibson *et al.* 1995a). In addition, Gibson *et al.* (1995b) demonstrated the kamafugitic affinity of most of the alkaline silicate-rock intrusions in the remainder of the APIP.

Kamafugite lavas and tuff horizons of the Mata da Corda often contain xenoliths of plutonic rocks such as dunite, pyroxenite, melilitite and syenite. This rock association is also common in the carbonatite complexes of APIP. Seer and Moraes (1988) suggested that large amounts of these xenoliths in the volcanic breccias may indicate the presence of intrusive carbonatite-bearing complexes at depth. This is consistent with geophysical evidence (Bosum 1973), where some of the dipole anomalies observed in aeromagnetometry maps coincide with the known intrusive alkaline bodies, whilst numerous others may represent unexposed intrusions.

The composition of clinopyroxenite xenoliths in the Mata da Corda kamafugites (calcic augite, phlogopite, perovskite, apatite, melanite, opaque minerals and occasionally Ba-rich feldspar) closely resembles that of the bebedourites. In addition, nearly all of the textural features observed in bebedourites (Brod 1999) are also present in the Mata da Corda xenoliths (Seer and Moraes 1988, Lloyd and Bailey 1991, Brod 1999). It would therefore seem entirely appropriate to classify these rocks as bebedourite. In this context, another crucial piece of evidence comes from the presence of kalsilite in some of the Mata da Corda xenoliths, recently reported by Sgarbi *et al.* (1998). So far, kalsilite has not been positively identified in connection with the carbonatite complexes, but XRD evidence (Brod 1999) suggests the presence of kalsilite, in addition to nepheline, in syenitic dykes from Salitre, and the recognition of this mineral in bebedourites within the carbonatite complexes may be just a matter of time. Finally, phlogopite-picrites dykes, which are abundant in APIP carbonatite complexes, are also found cutting the Mata da Corda volcanic rocks (Gibson *et al.* 1995b).

The combined field, petrographic and geophysical evidence presented above, together with the obvious spatial and temporal association, strongly suggest that the plutonic counterparts of the Mata da Corda kamafugites are bebedourites. These are analogous to those occurring in the carbonatite complexes and thus provide an important link between the carbonatites and the kamafugitic magmatism in APIP.

**GEOCHEMISTRY** The Late Cretaceous magmatism in the Alto Paranaíba Igneous Province (APIP) has been the subject of numerous recent chemical and isotopic studies, which have attempted to identify the geochemical characteristics of the underlying mantle regions (e.g.

Toyoda *et al.* 1994, Bizzi *et al.* 1995, Gibson *et al.* 1995b, McDonald *et al.* 1995, Carlson *et al.* 1996). With the exception of Toyoda *et al.* (1994), a general consensus seems to emerge from these studies that the vast majority of APIP parental melts originated from metasomatised sub-continental lithospheric mantle (SCLM). The predominance of rocks with mineralogical and/or geochemical kamafugitic affiliation in both plutonic and volcanic settings of APIP, is also now widely recognised (e.g. Sgarbi and Valença 1993, Sgarbi and Gaspar 1995, Sgarbi *et al.* 1998, Gibson *et al.* 1994, 1995b, Tompkins *et al.* 1998, Araújo *et al.* 1998).

Brod (1999) demonstrated that the phlogopite-picrite dykes are primitive magmas and that all of the silicate and carbonate primary (magmatic) rocks in the Tapira complex may be derived from them, through a complex interplay of liquid immiscibility and fractional crystallisation processes. This is likely to be the case in the other APIP carbonatite complexes as well, given their strong petrographic similarity and the intimate association of phlogopite-picrite dykes with all of them. Phlogopite-picrites thus provide an excellent opportunity for investigation of the geochemical affiliation of the parental magmas of APIP carbonatite complexes.

**Major-element oxides** Major-element analysis of the phlogopite-picrites (and also of some more-evolved ultramafic, syenitic and carbonatitic dykes) must take into account the abundance of some elements that are usually regarded as traces in other igneous rocks. For instance, in some samples, Ba and Sr can reach percentage levels and must be included in the major-element analyses, in order to achieve acceptable analytical totals. A second problem posed by these rocks is that Loss on Ignition (LOI) values are mostly related to the presence of groundmass carbonate and/or carbonate globules (ocelli). Since in most cases this is a magmatic feature of phlogopite-picrites (Brod 1999), the usual technique of recalculating the analyses on a "dry" basis would induce unrealistic interpretation of the data. Therefore, data for major-element oxides are reported here on a "wet" basis.

Phlogopite-picrites display progressive variation in many of the major-element oxides, and are probably best considered as representing several stages of a differentiating series. The following general features can be regarded as typical of the phlogopite-picrite series. All samples are extremely silica-undersaturated ( $\text{SiO}_2 = 24.30 - 33.40$  wt. %), carbonate-rich ( $\text{CO}_2 = 3.29 - 21.9$  wt. %) and strongly potassic ( $\text{K}_2\text{O} = 2.68 - 5.87$  wt. %). They typically have high CaO ( $9.31 - 19.41$  wt. %) and  $\text{TiO}_2$  ( $3.24 - 5.37$  wt. %) contents and are low in  $\text{Al}_2\text{O}_3$  ( $1.63 - 4.69$  wt. %). The  $\text{K}_2\text{O}/\text{Na}_2\text{O}$  ratio varies from 2.14 to 14.86, on a wt. % basis. The high MgO ( $8.73 - 21.34$  wt. %) testifies to the primitive character of these rocks. Their mg-number varies from 42 to 62, and correlates positively with strongly compatible trace elements, such as Cr and Ni.

**Trace elements** The high Cr (134 - 1012 ppm) and Ni (82-932 ppm) contents are consistent with the ultramafic character of these rocks and suggest that the least evolved members of the phlogopite-picrite series can be considered as primitive magmas. Scandium is moderately high, varying from 22 to 37 ppm. Phlogopite-picrites are strongly enriched in incompatible elements. In particular, Ba, Sr and the sum of the rare-earth elements (REE) often reach concentrations of a few tenths of a percent. Concentrations of the high-field-strength elements (HFSE) are relatively high: Nb (160 - 360 ppm); Ta (10 - 22 ppm); Zr (470 - 1000 ppm) and Hf (10 - 23 ppm). The trace-element distribution is illustrated in the chondrite-normalised spider diagram of Fig. 2.

It should be noted that all samples behave coherently in Fig. 2. Each pattern peaks at La, with the most enriched samples showing La concentration over 1600 times the chondritic abundance. Negative anomalies at Rb, K and Sr, together with a less pronounced negative Ti anomaly, are also present. Few samples show an incipient phosphorus negative anomaly. The chondrite normalised ratios of the "paired" elements, such as Zr/Hf, Y/Ho and Nb/Ta show little variation, in the ranges 1.16-1.52, 8.6-9.9 and 0.58-1.34, respectively.

The REE patterns of the phlogopite-picrites are shown in Fig. 3. The patterns are smooth and steep, showing strong enrichment of the light rare-earth elements (LREE) relative to the heavy rare-earth elements (HREE). Chondrite-normalised La/Lu ratios range from 69 to 210, but most are concentrated between 90 and 160. La/Sm(N) ratios vary from 16.7 to 19.3.

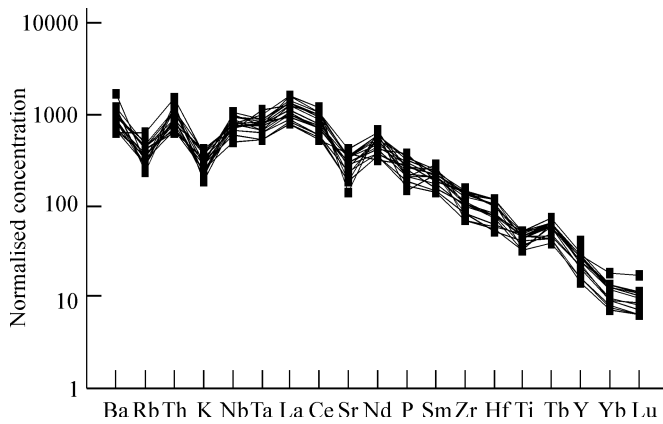


Figure 2-Trace-element diagram showing the composition of phlogopite-picrites, normalised to chondrite (except Rb, K and P) according to Thompson *et al.* (1984).

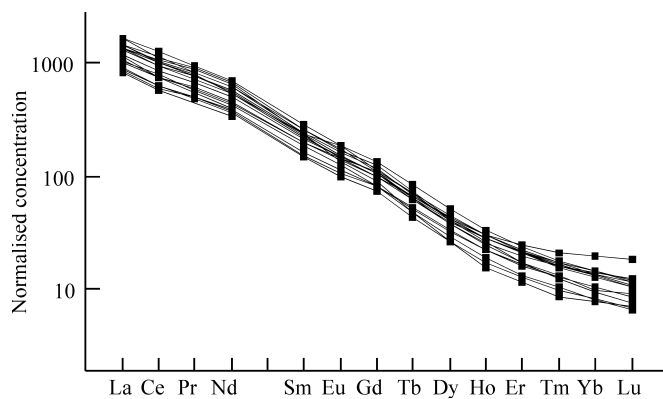


Figure 3-Chondrite-normalised REE patterns of phlogopite-picrites. Normalisation factors are from Boynton (1984).

**Chemical classification of the phlogopite-picrites** Any attempt at a chemical classification of the phlogopite-picrites has to account for the peralkaline (mol. ratio of alkalis to aluminium from 1.04 to 3.02) and ultrapotassic ( $K_2O/Na_2O$  from 2 to 15) character of these rocks, as well as their extreme silica-undersaturation. Classification diagrams involving the relative abundance of silica and alkalis, such as those of Cox *et al.* (1979) and Le Maitre *et al.* (1989) are difficult to apply, because of the excessively low silica content of the phlogopite-picrites. Mitchell (1996) emphasised that the total alkalis vs. silica (TAS) diagram is "totally inadequate" for the classification of several types of alkaline rocks, such as kimberlites, lamproites, kamafugites, melilitites and nephelinites.

A chemical scheme for the classification of ultrapotassic rocks was devised by Foley *et al.* (1987). Before the system is applied, the whole-rock analyses must pass a chemical screen ( $MgO > 3$  wt. %,  $K_2O > 3$  wt. %, and  $K_2O/Na_2O > 2$ ), in order to confirm the ultrapotassic character of the rock and to concentrate the classification on mafic varieties only. All the studied samples of phlogopite-picrite have  $MgO$  higher than 8 wt. %, and only one fails to satisfy the minimum  $K_2O$  limit, despite its relatively high  $K_2O/Na_2O$  ratio. This sample shows textural evidence of alteration, which could be responsible for changes in the original  $K_2O$  content.

Fig. 4 shows plots of phlogopite-picrite analyses from the Tapira complex in a number of the classification diagrams proposed by Foley *et al.* (1987). The phlogopite-picrites invariably plot within or near the field of Group II ultrapotassic rocks (kamafugites). Although the mineralogy of these rocks does not warrant their classification as kamafugites in a petrographic sense, their geochemical affinity with the kamafugitic series is obvious, at least in terms of major elements. Data from Gibson *et al.* (1995b) for kamafugitic rocks of the APIP and for phlogopite-picrites associated with other carbonatite complexes in the province are also plotted in Fig. 4, for comparison. Phlogopite-picrites from Tapira have slightly lower  $SiO_2$  and  $Al_2O_3$  than the APIP kamafugites and higher  $CaO$  than phlogopite-picrites from other APIP carbonatite complexes. The  $K_2O/Al_2O_3$  ratio is higher in Tapira phlogopite-picrites than in both groups of APIP rocks.

The geochemical affinity of phlogopite-picrites with kamafugitic rocks is also observed in trace-element ratio diagrams, such as  $Th/Yb$  vs.  $Ta/Yb$  and  $Th/Zr$  vs.  $Nb/Zr$  (Fig. 5). Both diagrams show that phlogopite-picrites and APIP kamafugites plot at the higher end of the array of rocks derived from non-subduction-related mantle sources.

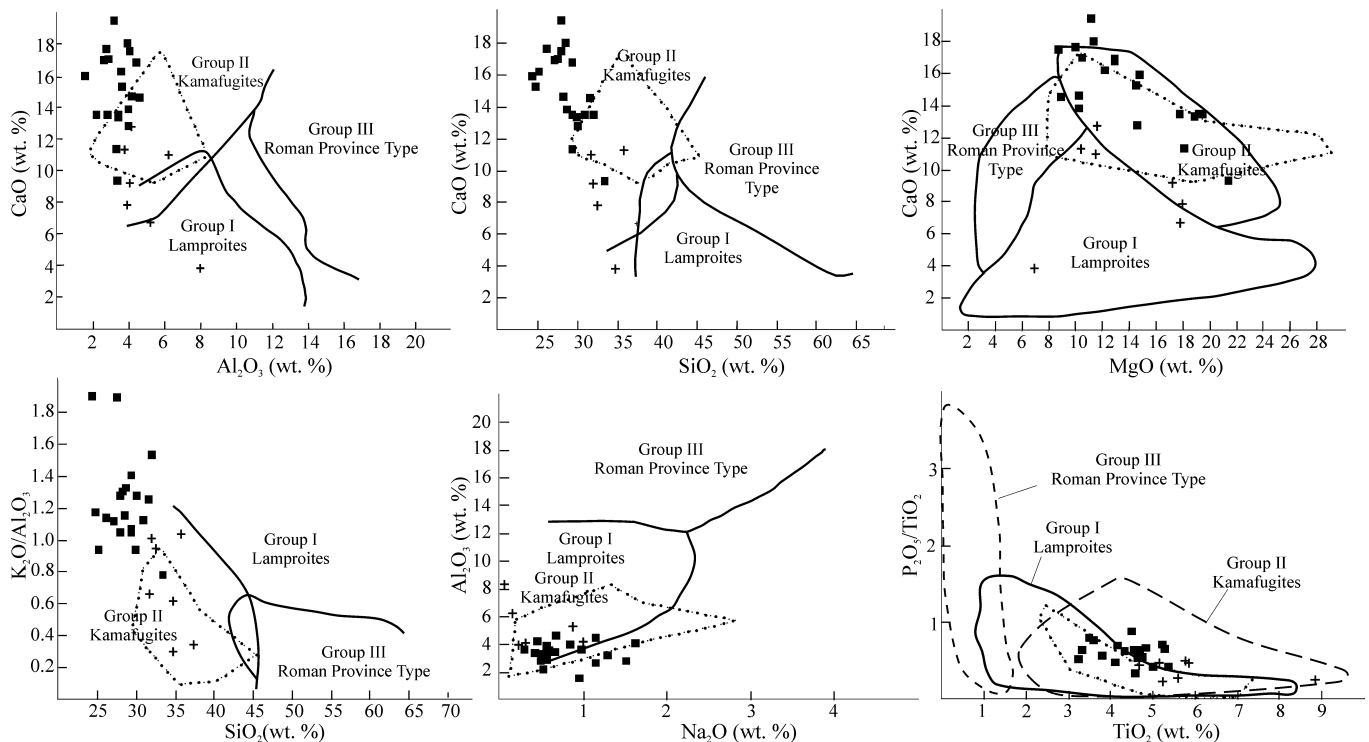


Figure 4-Major-element variation in the classification diagrams of Foley *et al.* (1987), showing the composition of Tapira phlogopite-picrites (squares). Data for other APIP phlogopite-picrites (crosses) and APIP kamafugites (fields outlined by dotted lines) are from Gibson *et al.* (1995b) and Brod (1999).

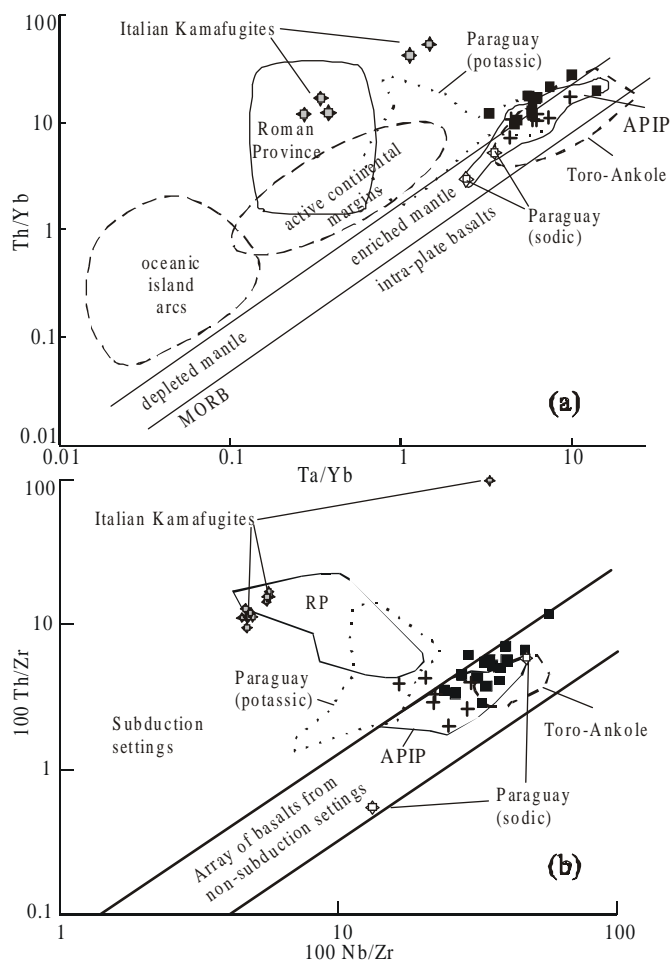


Figure 5—A) Th/Yb vs. Ta/Yb diagram showing the composition of Tapira phlogopite-picrites (solid squares), compared with other rock types. Data for potassic and sodic rocks from Paraguay are from Comin-Chiaromonti *et al.* (1997), Italian kamafugites from Conticelli and Peccerillo (1992) and from Peccerillo *et al.* (1988), APIP rocks from Gibson *et al.* (1995b). Other fields are from Wilson (1989), and references therein. B) Th/Zr vs. Nb/Zr diagram, showing the composition of phlogopite-picrites from Tapira (solid squares) and other APIP phlogopite-picrites (crosses), compared with potassic rocks from other provinces. Data for APIP rocks are from Gibson *et al.* (1995b), Italian kamafugites are from Conticelli and Peccerillo (1992) and Peccerillo *et al.* (1988). Other fields are after Comin-Chiaromonti *et al.* (1997).

The chemical similarity of APIP rocks to the Toro-Ankole kamafugites is also evident. The APIP rocks are distinct from other potassic rocks, such as those from the Roman Province-type, Italian kamafugites and Eastern Paraguay, which have lower relative concentrations of Nb and Ta. Sodic alkaline igneous rocks from Paraguay, however, have compositions analogous to APIP, a feature also noted by Comin-Chiaromonti *et al.* (1997), who assumed that their mantle source was similar to that postulated by Gibson *et al.* (1995b) for the Alto Paranaíba ultrapotassic magmas.

Fig. 6 compares the chondrite-normalised ratios of trace elements in the Tapira phlogopite-picrites with those of other examples of alkaline rocks. The pattern for Tapira samples agrees well with those of the APIP kamafugites and shows a nearly perfect coincidence with phlogopite-picrites from other carbonatite complexes in the province. Both the Ugandan and Italian examples of kamafugites are less enriched in most elements than the APIP rocks. Additionally, the Italian kamafugites exhibit characteristic negative anomalies at Nb and Ta, which are lacking in the Ugandan and APIP rocks. The negative anomalies of K and Rb are more pronounced in the sample of APIP kimberlite than in the phlogopite-picrites. APIP lamproitic rocks are distinguished from the phlogopite-picrites by their lower contents of LREE, and by the absence of negative Rb and K anomalies.

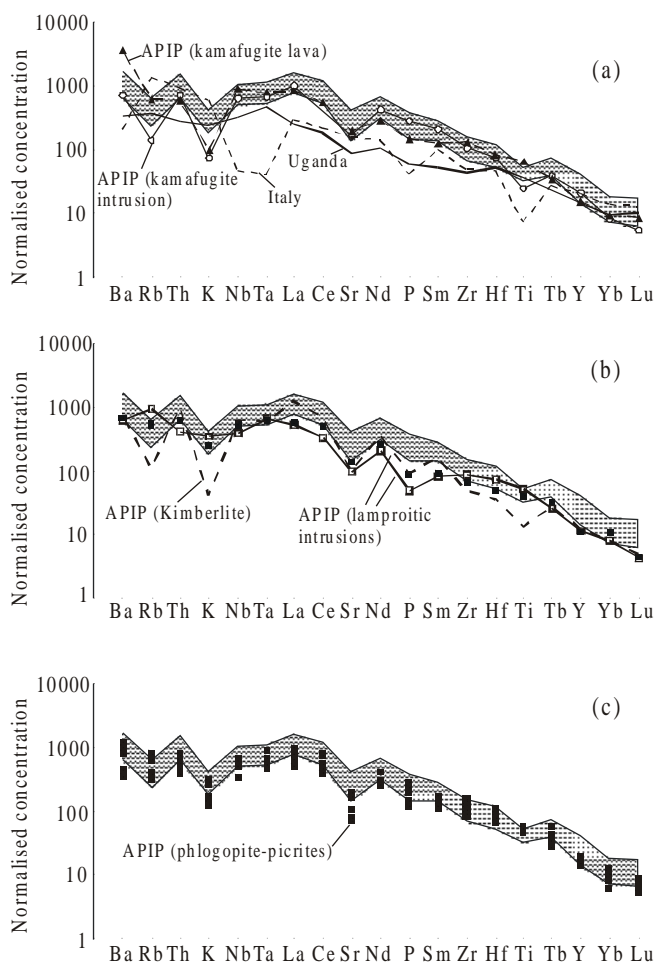


Figure 6—Spider diagrams comparing the composition of Tapira phlogopite-picrites (shaded field) with: (a) kamafugites from APIP (Gibson *et al.* 1995b), Italy (Peccerillo *et al.* 1988) and Uganda (Thompson *et al.* 1984); (b) kimberlite and lamproitic rocks from APIP (Gibson *et al.* 1995b); (c) phlogopite-picrites associated with other APIP complexes (Gibson *et al.* 1995b). Concentrations normalised to chondrite (except Rb, K and P) according to Thompson *et al.* (1984).

Finally, initial  $^{87}\text{Sr}/^{86}\text{Sr}$  and  $^{143}\text{Nd}/^{144}\text{Nd}$  ratios obtained by Brod *et al.* (1999) for carbonatites and silicate rocks of the Tapira complex are within the overall range of the APIP rocks (Gibson *et al.* 1995b) suggesting a broadly similar mantle source region for the carbonatite and ultrapotassic magmatism.

**DISCUSSION AND CONCLUSIONS** Detailed variations in the chemical properties of kamafugites, lamproites and kimberlites from APIP led Gibson *et al.* (1995b) to suggest heterogeneities in the melt source regions of these rocks. Thus, they interpreted the high CaO and Sc of the kamafugites as indicative of a clinopyroxene-rich mantle source. The favoured mechanism for the production of the postulated source mineralogy is the conversion of mantle harzburgite or lherzolite to wehrlite by reaction with dolomitic carbonatite melt (e.g. Dalton and Wood 1993, Moore and Wood 1998), which is believed to take place at relatively low pressures (< 25 kbars). On the other hand, high-pressure assemblages in xenoliths (Carvalho and Leonards 1995) from the Três Ranchos kimberlite suggest a much deeper (> 150 km), clinopyroxene-depleted source for this kimberlite. The mantle source of APIP lamproites is interpreted as somewhat depleted in clinopyroxene with respect to the kamafugites (Gibson *et al.* 1995b).

The evidence presented in this paper strongly suggests that the origin of the APIP carbonatites is inextricably linked with the ultrapotassic magmatism. The bulk-rock chemistry in the phlogopite-picrites from Tapira and other carbonatite complexes most closely resemble the chemical features of the APIP kamafugites (e.g. Mata da Corda volcanics). In particular, similarities in trace-element and REE

ratios suggest that the phlogopite-picrites and the kamafugites share a common source, and underwent comparable post-melting processes.

The link between the kamafugitic rocks and the carbonatite complexes is further emphasised by the common presence of xenoliths of pyroxenite (bebedourite) and ultrapotassic syenites within the Mata da Corda pyroclastics, indicating the presence of magma chambers similar to the carbonatite complexes at depth. This is consistent with the suggestion by Lloyd and Bailey (1991), that kamafugites may be the parental magmas for APIP bebedourites.

The association of carbonatites with potassic igneous rocks of kamafugitic affinity is well known from the Late Pleistocene Umbria-Latium ultra-alkaline district, Central Italy (Stoppa and Cundari 1995). However, it should be noted that the origin and tectonic environment of these rocks is distinct from those of the APIP complexes. For

instance, Stoppa and Lavecchia (1992) and Lavecchia and Stoppa (1996) suggested that the primary melts of the Italian kamafugites and carbonatites were derived from a source region in the asthenosphere. Also, some of the key features in mineral chemistry and whole-rock geochemistry of the Italian kamafugites and carbonatites are distinct from those of the APIP (Brod 1999). In this respect, the former more closely resemble the Early-Cretaceous mafic-potassic/carbonatite associations of Eastern Paraguay.

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