

GEOOTHERMOMETRY OF THE POÇOS DE CALDAS NEPHELINE SYENITES, SOUTHERN BRAZIL*

MABEL NORMA COSTAS ULBRICH, presented by A. C. ROCHA-CAMPOS — Instituto de Geociências, Universidade de São Paulo, São Paulo, SP — Nepheline syenites (NeS), the only phaneritic rocks outcropping in the Poços de Caldas alkaline massif, southern Brazil, are leucocratic to hololeucocratic varieties. The main felsic minerals are K-feldspar (KF, 40-60%) and nepheline (Ne, 20-36%); the former is K-rich (>80% Or), mostly structurally inhomogeneous (highly ordered microcline and orthoclase sometimes coexisting in the same grain, or in different grains of the same sample). Ne is mediopotassic. Sodic pyroxene (soda augite, aegirine-augite or aegirine) is usually subordinate, joined sometimes by minor biotite or Mg-arfvedsonite (this last one in agpaitic rocks); biotite is the only mafic mineral in some facies. Agpaitic rocks may show up to 10-15% rare-metal silicates (mainly eudialite); miaskitic or intermediate facies (which are predominant) are characterized by sphene, fluorite and magnetite. NeS, although monotonous, appear as several different facies, identified by variations in texture and accessory mineralogy; they crop out mainly in the northern half of the district, as small to medium-sized magmatic bodies (which rarely cover more than 5 or 6 km²). Geothermometry of NeS was based on Ne and Ne-KF diagrams of Hamilton (1959, *J. Geol.*, **69**, 321-329) and Powell & Powell (1977, *Contr. Min. Petrol.*, **62**, 193-204) using electron microprobe data (ARL 3-channel manual equipment).

The coarser-grained NeS (both agpaitic and miaskitic) show Ne with slight excess SiO₂ (<5%), with data clustering around the Morozewicz-Buerger convergence field (limited by the Buerger, Ne₇₃Ks₂₇, and Morozewicz, Ne_{75.1}Ks_{20.9}Qz_{4.0}, compositions; cf. Tilley, 1954, *Am. J. Sci.*, **252**, 65-75). KF (usually, Or 80%) is unzoned or weakly zoned, with Na-richer centers. Three cases are identified in these rocks: 1) agpaitic facies, with coexisting orthoclase and microcline, with geothermometric T of 500-540°C, thus defining probably true magmatic crystallization T; 2) facies with highly ordered microcline as the predominant KF, with T less than 500°C, recording re-equilibration during cooling, with Na-K magmatic exchange reactions; 3) some facies show Ne with near-Buerger compositions accompanied by maximum microcline, attesting to submagmatic re-equilibration.

Finer-grained NeS are generally more rapidly quenched border facies. Ne are typically Qz-rich (5% or over), and KF is mainly orthoclase. In miaskitic varieties, Ne is compositionally homogeneous; KF is zoned (e.g. Or₇₆Ab₂₄ at the center, and Or₉₅Ab₅ at the border). Textures are usually poikilitic, with larger KF plates enclosing small Ne euhedra. Ne composition, plotted on a Hamilton diagram, shows crystallization T of 730-800°C for miaskitic types. In similar finer-grained agpaitic facies, KF is weakly zoned, and textures indicate KF-Ne joint crystallization

along cotectic lines; compositions of the two phases suggest crystallization T of 580-600°C. — (13 de novembro de 1984).

THE ULTRAMAFIC-ALKALINE AND GABBROIC ALKALINE LINEAGES: DERIVATION BY FRACTIONATION OF ULTRABASIC ALKALINE MAGMAS?*

HORSTPETER H. G. J. ULBRICH, presented by A. C. ROCHA-CAMPOS — Instituto de Geociências, Universidade de São Paulo, Cidade Universitária, São Paulo, SP — Most nepheline syenites (NeS) are associated with ultramafic rocks (peridotites, pyroxenites, jacupirangites, rocks of the ijolite series), both with and without accompanying alkaline gabbroic rocks. If the main difference between these rock associations is given by the presence, or absence, of calcic plagioclase, then two distinct lineages can be set up: the gabbroic-alkaline and the ultramafic-alkaline, as already pointed out in the literature (Sheynmann *et al.*, 1963, *Internat. Geol. Rev.*, **5**, 451-457). The anorthite component of plagioclase is unstable at higher pressures and crystallizes only from magmas which are relatively Ca- and Al-rich. It is thus tempting to propose a common origin for the two lineages, whose main difference would be given by the pressure level at which Ca is fractionated out of the parental magmas.

There is little doubt that most NeS are rocks derived from parental basic or ultrabasic magmas. Possible compositional ranges of these parents, although varied, can be determined both from actual examples (e.g., possible primary magmas) and on theoretical grounds (e.g., what melts can be derived from a peridotitic mantle?): these parental magmas have to be relatively SiO₂-poor, rich in MgO and CaO, with low Al₂O₃ and probably also very poor in alkalis.

Chemically, conversion of these magmas into NeS magmas means enrichment in SiO₂, Al₂O₃ and alkalis, and very strong depletion both in MgO and CaO. These chemical trends can be duplicated if wehrlitic fractions can be separated from the parental magmas (i. e., separation of Mg-olivine and Ca-pyroxene). Wehrlitic fractionation opens up two possibilities: a) efficient early separation of a wehrlitic fraction, depleting the parental magma strongly in Ca and inhibiting a lower-pressure crystallization of Ca-plagioclase, thus giving rise to the ultramafic-alkaline lineage; b) enhanced early separation of Mg-olivine, over that of Ca-pyroxene, may drive the compositions of derived magmas into the plagioclase phase volume, allowing this mineral to crystallize at lower pressures, thus reproducing a gabbroic-alkaline trend.

A general problem remains, mainly related to the very low alkali content of most ultrabasic parental magmas (e.g., komatiites, picrites). The problem may be solved if it is assumed that the primary ultrabasic magmas of alkaline trends are already alkali-enriched (e.g., by melting of alkali-rich mantle metasomatites). Rocks which represent such hypothetical magmas are unknown. The

* Financed by FINEP, Proc. n.º 3.3-82.0381.00; FAPESP, Proc. 79/206-5; CNPq, Proc. 40.1412/81.

* Financed by FAPESP and CNPq.