S. Jung · S. Hoernes · K. Mezger

Synorogenic melting of mafic lower crust: constraints from geochronology, petrology and Sr, Nd, Pb and O isotope geochemistry of quartz diorites (Damara orogen, Namibia)

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Abstract Quartz diorites represent the earliest (ca. 540 Ma) and most primitive plutonic rocks in the Pan African Damara belt and they pre-date the main phase of high-T regional metamorphism. Two suites of synorogenic quartz diorites are unusual among Damaran intrusive rocks in their elemental and isotopic features. Comparison of the diorite compositions with melts from amphibolite-dehydration melting experiments points to a garnet-bearing meta-tholeiite, probably enriched in K_2O , as a likely source rock. Partial melting processes generated mafic (ca. 50 wt% SiO₂) quartz diorites in the deep crust at temperatures of between 1,000 and 1,100 °C, based on comparison with experimental results and similar temperature estimates based on P₂O₅ solubility in mafic rocks. Subsequently, the quartz diorites evolved by multistage, polybaric differentiation processes including fractional crystallization of mainly hornblende and plagioclase and assimilation of felsic basement gneisses. Although their chemical characteristics (high LILE, low HFSE) resemble those of other quartz diorites with calc-alkaline affinities, they differ in their enriched Sr (initial ⁸⁷Sr/⁸⁶Sr: 0.70943–0.71285), Nd (initial ϵ Nd: -9.1 to -15.2) and O (δ^{18} O: 6.8-8.1%) isotope compositions. Neodymium model ages (T_{DM})

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that range from 1.7 to 2.2 Ga and large variation in ²⁰⁷Pb/²⁰⁴Pb relative to ²⁰⁶Pb/²⁰⁴Pb indicates involvement of ancient crustal material. Lead (²⁰⁶Pb/²⁰⁴Pb: 17.08– 17.23, ²⁰⁷Pb/²⁰⁴Pb: 15.53–15.62, ²⁰⁸Pb/²⁰⁴Pb: 37.71– 38.16) isotope compositions are strongly retarded, indicating that the source underwent a pre-Pan-African U/Pb fractionation and U depletion. It is proposed that the quartz diorites originated by synorogenic high temperature melting of mafic lower crust. This contrasts with previous suggestions favouring an origin of these rocks by melting of an enriched mantle during Pan-African times with characteristics modified by subduction of oceanic crust and sedimentary rocks.

Introduction

Granite batholiths are an important and characteristic component of continental crust. They form from granitic melts that are often not primary melts, but are modified by differentiation processes prior to emplacement and crystallization. The relative importance of fractional crystallization versus partial melting and the role of open system processes (magma mixing, assimilation) in the genesis of these differentiated melts is still being debated. In areas where granite batholiths consist not only of granites sensu stricto, but also of more mafic rocks (gabbros, diorites), a fundamental role has been assigned to these mafic magmas. In general, they may be mantle-derived parental magmas (LeBel et al. 1985), end members in mixing and fractionation-assimilation processes (DePaolo 1981; Reid et al. 1983; Kistler et al. 1986; Eggins and Hensen 1987), material from lower crustal source regions (Gromet and Silver 1987; Tepper et al. 1993) and/or heat sources that drive melting of overlying continental crust (Pitcher 1987). Additionally, their petrological diversity can also be the result of source heterogeneity (Noves et al. 1983).

Granitoid intrusions are an important rock type throughout the Damara Belt (Namibia). Chemical and isotopic data accumulated in recent years indicate that these granitoids consist of several distinct suites. These suites include (1) synorogenic S-type granites that originated by partial melting of upper crustal metasedimentary rocks (Haack et al. 1982; McDermott et al. 1996; Jung et al. 2000a, 2001), (2) synorogenic granites that were derived by melting of undepleted metasedimentary and meta-igneous basement rocks (McDermott et al. 1996; Jung et al. 2002) and (3) late-orogenic A-type granites, some of them represent mixtures between moderately depleted meta-igneous source rocks and a component from the lithospheric mantle (Jung et al. 1998b). One objective of this study is to unravel the petrogenesis of synorogenic dioritic intrusions.

High-precision Sm-Nd and Pb-Pb (garnet, zircon) and U-Pb ages (monazite, titanite) are now available from a number of migmatite and granite occurrences throughout the high-grade central part of the Pan-African Damara orogen of Namibia (Briqueu et al. 1980; Kukla et al. 1991; Jung et al. 1998b, 2000a, 2000b, 2001; Jung and Mezger 2001, 2002). These ages provide a temporal framework for the structural and petrological evolution of the orogen. Thus, another objective of this study is to determine the temporal relationships between the time of intrusion of the quartz diorites as constrained by new high-precision U-Pb titanite ages relative to the time of high-grade metamorphism and melting. This comparison is particularly relevant for the quartz diorites because they may represent an important thermal component during orogeny.

The Damara Belt is interpreted to have formed mainly by continental collision including subduction, although the significance of the ocean floor involved in the orogeny is still a matter of debate (Kröner 1982; Miller 1983). In contrast to many typical collision belts (e.g. Caledonides, Alps), the Damara Belt is characterized by the emplacement of large volumes of granitoids similar to that observed in some active continental margins (e.g. Andes) or in broad ensialic orogens (e.g. Lachlan Fold belt). However, the general pattern of the Damaran plutonism contrasts with Cordilleran-type plutonism, both in its spatial distribution and bulk chemistry. In the Damara orogen, the proportions of gabbro:diorite-granodiorite:tonalite-granite:leucogranite contrast markedly with those found in continental margin environments. In the Andes, the proportions gabbro:diorite-granodiorite:tonalite-granite of are 16:58:26 (Pitcher 1978), whereas in the Damara Belt the proportions are 2:2:96 (Miller 1983), broadly similar to the proportions found in the Himalayas (0:0:100; Le Fort et al. 1987). Furthermore, the plutonic rocks in the Damara orogen are not arranged in linear belts, and leucocratic rocks are far more abundant than in subduction zone settings. Hawkesworth and Marlow (1983) have, based on trace element arguments, questioned the derivation of most Damaran granites from a subduction zone. Nonetheless, several workers (Kasch 1983; Miller 1983) have suggested that the Pan-African orogeny, and hence the associated plutonism of the Damara orogen of Namibia, is related to subduction zone processes. The

paucity of geological evidence for a fossil subduction zone, e.g. the lack of eclogites and blueschists indicating a high P–low T regime casts some doubts on this hypothesis and alternative models involving ensialic tectonism have been proposed (e.g. Kröner 1982). Because Kasch (1983) and Miller (1983) used the occurrence of quartz diorites and associated granodiorites and granites in the southern central Zone of the Damara Belt (Fig. 1) as an argument for a subduction zone setting, an evaluation of the isotope systematics of these rocks highlights the relative importance assigned to the mantle or to the crust for melt production during orogeny. Such an isotope study can provide valuable constraints on the relative role of crustal recycling and new mantle additions to the continental crust during orogenic processes.

Analytical techniques

Whole rock powders were prepared using a jaw crusher, a ball mill and an agate mortar. Major and some trace elements (except for REE) were determined on fused lithium-tetraborate glass beads using standard XRF techniques. Rare earth elements were analysed by inductively coupled plasma emission spectrometry following separation of the matrix elements by ion exchange (Heinrichs and Herrmann 1990). Loss on ignition (LOI) was determined gravimetrically after heating the samples at 1,050 °C for 1 h (Lechler and Desilets 1987). FeO was measured titrimetrically with standard techniques. Accuracy has been controlled by repeated measurements against several international and in-house standards.

For the Rb–Sr and Sm–Nd whole rock isotope analyses, the samples were spiked with a $^{149}\rm{Sm}/^{150}Nd$ and a $^{85}\rm{Rb}/^{84}Sr$ tracer and digested in concentrated HF-HNO₃ in 3-ml screw-top Teflon vials inside Krogh-style Teflon bombs at 200 °C for 3 days. After complete dissolution, the samples were dried down and redissolved in 2.5 N HCl. Rb, Sr and REE were separated by using standard cation exchange columns with a Dowex AG 50 W-X 12 resin using 2.5 N HCl for Rb and Sr and 6 N HCl for the REE. Nd and Sm were separated from the other REE by using HDEHP-coated Teflon columns and 0.12 N HCl for Nd and 0.3 N HCl for Sm. Isotope analyses were carried out at the Max-Planck-Institut für Chemie at Mainz using thermal ionization mass spectrometry with a Finnigan MAT 261 multicollector mass spectrometer operating in the static mode. Rb, Sm and Nd were run on Re double filaments and Sr was run on W single filaments. Nd isotopes were normalized to 146 Nd/ 144 Nd = 0.7219. The total procedural blank for Nd was < 40 pg and is considered to be negligible. Repeated measurements the La Jolla Nd standard gave 143 Nd/ 144 Nd = of 0.511854 ± 0.000031 (2σ ; n=24). The reproducibility of the Sr standard (NBS 987) is 87 Sr/ 86 Sr = 0.710218 ± 0.000029 (2 σ ; n = 24) and the fractionation was corrected to ${}^{86}Sr/{}^{88}Sr$: 0.1194. Uncertainties in the ${}^{87}Sr/{}^{86}Sr$ and ${}^{143}Nd/{}^{144}Nd$ are reported in the last two digits. Typical analytical errors in the 87 Rb/ 86 Sr and 147 Sm/ 144 Nd ratios are equal or better than 0.5 and 0.1%, respectively. Between 30 and 50 mg of high-purity K-feldspar separates were washed with a mixture of 3:1 HCl/HNO₃ to remove surface contamination and were subsequently rinsed three times with ultrapure water. After this treatment, the separates were leached three times in a mixture of concentrated HF/HNO₃, which resulted in a weight loss of \sim 70–80%. Subsequently, the feldspars were dissolved in concentrated HF and after evaporation redissolved in 2.5 N HCl and 0.6 N HBr and loaded on Teflon columns filled with Dowex AG 1 X 8 anion exchange resin (100-200 mesh) in chloride form (Mattinson 1986). The Pb was extracted using conventional HBr/HCl techniques and was loaded on Re single filaments following the H₃PO₄-silica gel method (Cameron et al. 1969). Pb analyses were corrected for mass fractionation by a factor of 0.11%

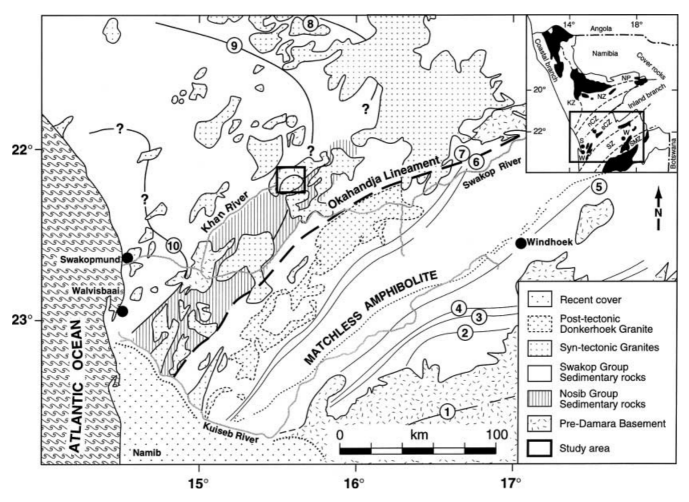


Fig. 1. Generalized geological map showing the study area within the Central Zone of the Damara orogen, Namibia. *Abbreviations* in *inset:* KZ Kaoko Zone; NP Northern Platform; NZ Northern Zone; nCZ northern Central Zone; sCZ southern Central Zone; SZ Southern Zone; SMZ Southern Margin Zone. Isograd map (Hartmann et al. 1983) gives the distribution of regional metamorphic isograds within the southern and central Damara orogen. Isograds: (1) biotite-in, (2) garnet-in, (3) staurolite-in, (4) kyanitein, (5) cordierite-in, (6) andalusite-sillimanite, (7) sillimanite-in according to staurolite-breakdown, (8) partial melting because of muscovite + plagioclase + quartz + H₂O \leftrightarrow melt + sillimanite, (9) K-feldspar + cordierite-in, (10) partial melting because of biotite + K-feldspar + plagioclase + quartz \leftrightarrow cordierite-melt + garnet. W Walvisbay; S Swakopmund; Wh Windhoek

per amu. The reproducibility of the standard NBS 982 was estimated to be 0.068, 0.064 and 0.071% for the ${}^{206}\text{Pb}/{}^{204}\text{Pb}$, ${}^{207}\text{Pb}/{}^{204}\text{Pb}$ and ${}^{208}\text{Pb}/{}^{204}\text{Pb}$ ratio, respectively. The total procedure blank was <100 pg Pb and, therefore, was, considered negligible.

After sieving the samples through a 40–80-mesh fraction, the mineral separates were purified by magnetic separation, separated by methylene iodide and handpicked, which produced high-purity separates. Only minerals free of inclusions were used. The titanites were washed in warm deionized water to remove surface contamination and were spiked with a ²⁰⁵Pb/²³⁵U spike before digesting in concentrated HF/HNO₃ in 3-ml screw-top Teflon vials inside Krogh-style Teflon bombs at 200 °C for several days. The titanites were boiled three times in 7 N HNO₃ and were subsequently dissolved in 3 N HCl for ion exchange chromatography. The Pb was separated using HCl–HBr chemistry and the U was separated using Eichrome resin and an improved 2 N HNO₃/0.02 N HNO₃ chemistry. Lead isotope analyses were carried out as described

above. Low U concentrations, combined with elevated common Pb concentrations, result in rather large uncertainties for the analyses. In order to correct for initial Pb composition, the Pb composition of leached K-feldspar from each sample was measured and was taken as the initial Pb composition during titanite growth.

Oxygen isotope analyses were performed at the University of Bonn on ~10-mg aliquots of powdered whole-rock samples, using purified fluorine for oxygen extraction, followed by conversion to CO₂ (Clayton and Mayeda 1963). ¹⁸O/¹⁶O measurements were made on a SIRA-9 triple-collector mass spectrometer by VG-Isogas. Analytical uncertainties are < 0.2%.

Geological setting and rock types

The Damara orogen exposes a deeply eroded section through a Pan African mobile belt that can be divided into a N–S-trending coastal branch, the Kaoko belt, and a NE–SW-trending intracontinental branch (see inset to Fig. 1). This mobile belt has been divided into several zones based mainly on stratigraphy, metamorphic grade, structure and geochronology (Miller 1983). The Central Zone is characterized by a large number of granitoids that record mineral and Rb/Sr whole rock ages mostly between 650 and 460 Ma and outcrop over an area of approximately 75.000 km² (Miller 1983; Fig. 1). Pre-Damara basement gneisses are overlain by Neoproterozoic to Palaeozoic metasedimentary sequences comprising quartzose sandstones, arkoses, mica schists, calc-silicate rocks, marble, metamorphosed glaciogenic diamictites, banded iron-stones, Al-rich metapelites, migmatites, metacarbonates and conglomerates (Miller 1983).

In the Central Zone (Fig. 1), previous estimates for the peak metamorphic temperatures are 560-650 °C at 3 ± 1 kbar for impure marbles (Puhan 1983). Metamorphic conditions based on oxygen isotope fractionation in a variety of meta-sedimentary and meta-igneous rocks yielded similar temperature estimates of 570-650 °C (Hoernes and Hoffer 1979). The metamorphic grade increases from east to west reaching high-grade conditions with local partial melting in the coastal area (Hartmann et al. 1983). However, more recent results indicated high-grade conditions that culminated in medium-pressure high-temperature granulite facies conditions with temperatures estimated to have been in excess of 700 °C at 4-6 kbar (Masberg et al. 1992; Jung et al. 1998c, 2000b; Jung and Mezger 2002). To the south-east, there is a gradation into the Okahandja Lineament Zone that separates the Central Zone from the Southern Zone (Fig. 1). In the Southern Zone, regional metamorphism is characterized by a Barrovian-type sequence with a general increase in the metamorphic grade from south to north. The metamorphic conditions range from low to medium pressures, and reached up to 8 kbar at maximum temperatures of 600 °C. Intrusions of large granitic bodies are absent even in the highest grade zone.

Important bodies of quartz diorites are exposed in the Goas (15.50°E/22.15°S) and Okongava (15.55°E/ 22.00°S) areas, which are located within the southern Central Zone (Fig. 1). Additional intrusions of quartz diorite include the nearby Palmental diorite (15.50°E/ 22.20°S) and the Mon Repos diorite (15.45°E722.05°S) both of which are not treated in this study. Plutons in the Goas and Okongava area are unzoned and range from quartz diorite through granite; leucogranites are less abundant. The plutonic rocks are generally medium grained, equigranular and hypidiomorphic with 50-70% plagioclase, 10-20% each hornblende and biotite, 5-15% guartz and 10–20% K-feldspar. Hornblende and plagioclase are the earliest-formed minerals and are virtually unzoned. Subhedral to anhedral hornblende can have rare inclusion of relict clinopyroxene and is usually associated with titanite and opaque minerals. Biotite may replace hornblende, quartz and K-feldspar occur mostly as interstitial phases. Accessory minerals include titanite, apatite, opaque minerals and zircon, in order of decreasing abundances. Alteration is weak or absent. Minor amounts of secondary minerals include mostly chlorite, muscovite and haematite. Enclaves of metasedimentary country rocks or other igneous rocks are absent.

Geochronology

In slowly cooled high-grade metamorphic terranes, U–Pb titanite ages are generally interpreted as dating the

time of cooling through the closure temperature for Pb volume diffusion (T_c). Estimates of T_c based on slowly cooled natural samples with a small grain size are ca. 600 °C (Tucker et al. 1986; Heaman and Parrish 1991) or 660-700 °C (Mezger et al. 1993; Scott and St-Onge 1995). Experimentally determined Pb diffusion parameters (Cherniak 1993) indicate a closure temperature for a grain size of ca. 200 µm of ca. 630 °C for a cooling rate of 10 °C/Ma and ca. 600 °C for a cooling rate of 2 °C/ Ma. Recent studies provide evidence that an inherited Pb component can be preserved in titanite incorporated into syenite melts (Pidgeon et al. 1996; Zhang and Schärer 1996). This indicates T_c in excess of 710 °C for titanite, but only for very short-lived magmatic events. In slowly cooled high-grade metamorphic terranes titanite ages probably date cooling during the waning stages of metamorphism (Mezger et al. 1991), whereas in shortlived igneous bodies U-Pb titanite ages can provide prograde ages that constrain the time of intrusion (Verts et al. 1996).

Titanite was recovered from both quartz diorites at Goas and Okongava. Dark brown titanite grains were generally sub-spherical, anhedral to euhedral and 100 to 500 µm in diameter. All selected fractions were free of visible inclusions. Most U-Pb analyses of titanite samples from Goas are concordant with ²⁰⁷Pb/²⁰⁶Pb ages of between 533 ± 4 and 538 ± 2 Ma (Tables 1 and 2, Fig. 2). One sample is normally discordant and yielded a similar 207 Pb/ 206 Pb age of 533 ± 3 Ma. One sample yielded a younger ${}^{207}\text{Pb}/{}^{206}\text{Pb}$ age of 503 ± 3 Ma. Most titanite fractions extracted from quartz diorites from Okongava are concordant, with 207Pb/206Pb ages of between 524 ± 2 and 557 ± 1 Ma. Normal discordant titanites yielded 207 Pb/ 206 Pb ages between 538 ± 2 and 551 ± 1 Ma. One fraction gave an apparently older 207 Pb/ 206 Pb age of 579 ± 4 Ma, whereas another fraction yielded a younger 207 Pb/ 206 Pb age of 520 ± 2 Ma. A regression line through all samples in 206 Pb/ 204 Pb versus 207 Pb/ 204 Pb space yielded a 207 Pb/ 206 Pb age of 551.9 ± 0.7 Ma.

Geochemistry

Major and trace elements

The quartz diorites have SiO₂ concentrations between 56.3 and 61.8 wt% (Goas) and between 51.6 and 59.5 wt% (Okongava) (Tables 3 and 4). All samples are metaluminous with alumina saturation index [ASI; = molar A/CNK: $(Al_2O_3/CaO + Na_2O + K_2O)$] values ranging from 0.88 to 1.01 at Goas and from 0.79 to 0.92 at Okongava. Mafic samples have $K_2O < Na_2O$, whereas more evolved samples have $K_2O > Na_2O$. There is a large variation in MgO, TiO₂, FeO_{total}, CaO, Al₂O₃ and K₂O among both suites of quartz diorites in which MgO, TiO₂, FeO_{total}, CaO, Al₂O₃ and K₂O increases with increasing SiO₂ (Fig. 3). The decrease in Al₂O₃ is more pronounced in samples from Okongava than in samples from Goas. Similarly,

Table 1. U–Pb tita	nite data from qu	Table 1. U-Pb titanite data from quartz diorites from the J	the Damara oroge	en (Namibia). Foi	Damara orogen (Namibia). For analytical details see Analytical techniques	see Analytical tec	hniques		
Sample name	Tnt M20	Tnt M21	Tnt M22	Tnt M24	Tnt M27	Tnt M29	Tnt M31	Tnt M33.1	Tnt M33.2
U (ppm)	143.0	138.1	134.0	297.4	312.6	227.7	142.2	411.6	340.9
Pb (ppm)	21.9	21.7	20.1	43.2	44.1	29.6	18.2	41.2	81.4
$^{206}\mathrm{Pb}/^{204}\mathrm{Pb}$	169.6	176.1	193.4	93.2	105.8	185.0	169.9	474.3	69.3
$^{208}{ m Pb}/^{206}{ m Pb}$	0.48113	0.53946	0.49757	0.14724	0.13825	0.22397	0.14658	0.61807	0.59990
207 Pb $/^{206}$ Pb	0.057661	0.058170	0.058138	0.058081	0.057303	0.058228	0.057845	0.058348	0.059319
Error $(\%)$	0.7759	0.2624	0.2451	0.5267	0.4564	0.2790	0.3367	0.7023	0.7162
$^{207}\text{Pb}/^{235}$ U	0.70361	0.71440	0.71056	0.62901	0.64623	0.70831	0.70993	0.50981	0.74548
Error (%)	1.1173	1.7544	1.7488	1.8593	1.8216	1.7671	1.7631	2.1307	1.9333
$^{206}\text{Pb}/^{238}$ U	0.08850	0.08907	0.08864	0.07855	0.08179	0.08822	0.08901	0.06337	0.09115
Error (%)	0.7538	1.7330	1.7300	1.7756	1.7579	1.7427	1.7272	1.9950	1.7818
²⁰⁶ Pb/ ²³⁸ U age	547	550	548	487	507	545	550	396	562
(million years)									
²⁰⁷ Pb/ ²³⁵ U age	541	547	545	495	506	544	545	418	566
(million years) 207pb/ ²⁰⁶ 11 aue	517	536	535	533	503	538	574	543	570
		0	0			0	110		

trace element abundances of V, Cr, Ni, Zr, Y and Sr decrease with increasing SiO₂ (Fig. 4). Again, for samples from Okongava, the decrease in Sr is more pronounced than in samples from Goas. Rubidium and Th increase with increasing SiO₂. Barium shows a different behaviour. In samples from Okongava, Ba first increases between ca. 52 and 57 wt% SiO₂ and then decreases from 57 to 60 wt% SiO₂ whereas in samples from Goas, Ba decreases from 56 to 62 wt% SiO₂. For both suites of quartz diorites, rare earth elements (REE) are enriched with chondrite-normalized La abundances of between 129 and 184 (Goas) and between 95 and 222 (Okongava). Chrondrite-normalized Lu abundances are similar in both suites and range from 7.6 to 12.4 (Fig. 5). Both suites of quartz diorites show no Eu anomaly in the mafic rock types. With increasing SiO₂, negative Eu anomalies develop with maximum Eu/Eu* values of 0.67 for Goas and 0.60 for Okongava. Chondrite-normalized La_n/Yb_n ratios and Rb, Zr and Nb concentrations are similar to most crust-derived granites from the Damara orogen (McDermott et al. 1996; Jung et al. 1998b), but are higher than in quartz diorites derived from an upper mantle modified by a subduction zone component (e.g. Shaw et al. 1993).

Strontium, Nd, Pb and O isotopes

The results of the Sr, Nd, Pb and O isotope analyses are reported in Table 5. The quartz diorites at Goas have an unradiogenic Nd isotopic composition with initial ϵ_{Nd} values ranging from -12.8 to -14.6 and initial ${}^{87}Sr/{}^{86}Sr$ ratios ranging from 0.71081 to 0.71285. Quartz diorites from Okongava have a larger spread in Nd isotopic composition, with initial ϵ_{Nd} values ranging from -9.1 to -15.2. Initial ${}^{87}Sr/{}^{86}Sr$ ratios have values between 0.70943 and 0.71251 (Fig. 6). For both suites, initial ϵ_{Nd} values become less radiogenic and Sr isotope compositions become more radiogenic with decreasing MgO and increasing SiO₂.

The leached feldspars from Goas have ²⁰⁶Pb/²⁰⁴Pb ratios that range from 17.16 to 17.23 and ²⁰⁷Pb/²⁰⁴Pb ratios ranging from 15.53 to 15.56 (Fig. 7; Table 5). At Okongava, ²⁰⁶Pb/²⁰⁴Pb ratios are less radiogenic ranging from 17.08 to 17.17, but ²⁰⁷Pb/²⁰⁴Pb ratios are more radiogenic with values between 15.56 and 15.63. The ²⁰⁸Pb/²⁰⁴Pb ratios show considerable overlap with ratios from 37.71–37.90 at Goas and from 37.78–38.16 at Okongava. In ²⁰⁷Pb/²⁰⁴Pb vs. ²⁰⁶Pb/²⁰⁴Pb space both suites plot above the Pb evolution curve of Stacey and Kramers (1975; Fig. 7a) indicating strongly retarded Pb isotope compositions. In the ²⁰⁸Pb/²⁰⁴Pb vs ²⁰⁶Pb/²⁰⁴Pb vs ²⁰⁶Pb/²⁰⁴Pb diagram the samples plot above the Th/U evolution curve (Fig. 7b). Like the Nd and Sr isotope composition, ²⁰⁷Pb/²⁰⁴Pb ratios show some variation with major element composition in which the Pb isotope ratios decrease with decreasing MgO for both suites. Similarly, for both suites, ²⁰⁷Pb/²⁰⁴Pb ratios decrease

Sample name	Tnt M34.1	Tnt M34.2	Tnt M35.1	Tnt M35.2	Tnt M38.1	Tnt M38.2	Tnt M39.1	Tnt M39.2
U (ppm)	339.9	274.4	322.5	429.3	265.3	265.5	225.1	185.5
Pb (ppm)	37.7	50.3	43.0	75.4	39.2	55.2	40.9	58.5
²⁰⁶ Pb/ ²⁰⁴ Pb	438.5	140.4	565.6	139.5	414.9	146.8	243.6	70.3
²⁰⁸ Pb/ ²⁰⁶ Pb	0.80147	0.78730	0.62360	0.63276	1.02569	1.01996	1.60322	1.69092
²⁰⁷ Pb/ ²⁰⁶ Pb	0.058564	0.057746	0.058738	0.058279	0.0582095	0.0585063	0.0584058	0.0585126
Error (%)	0.3469	0.3533	0.2182	0.3548	0.2555	0.3223	0.4101	0.8404
207 Pb/ 235 U	0.51087	0.69716	0.69109	0.71892	0.60723	0.73827	0.56814	0.71849
Error (%)	1.8870	1.8867	1.9305	1.8482	1.9539	1.8526	1.9034	1.9906
206 Pb/ 238 U	0.06327	0.08756	0.08533	0.08947	0.07566	0.09152	0.07055	0.08906
Error (%)	1.8521	1.8500	1.9167	1.8104	1.9351	1.8216	1.8531	1.7832
²⁰⁶ Pb/ ²³⁸ U age	395	541	528	552	470	565	439	550
(million years)								
²⁰⁷ Pb/ ²³⁵ U age	419	537	533	550	482	561	457	550
(million years)								
²⁰⁷ Pb/ ²⁰⁶ U age	551	520	557	540	538	549	545	549
(million years)								

Table 2. U-Pb titanite data from quartz diorites from the Damara orogen (Namibia). For analytical details see Analytical techniques

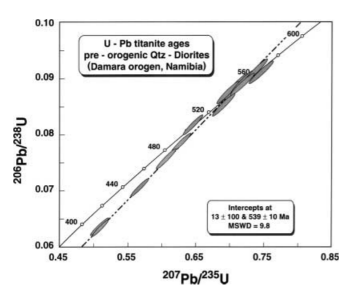


Fig. 2. U/Pb concordia diagram showing titanite analyses from quartz diorites

with decreasing ϵ_{Nd} values and increasing initial ${}^{87}Sr/{}^{86}Sr$ values.

Generally, all quartz diorite samples show rather low δ^{18} O values (Table 5). The variation of δ^{18} O for quartz diorites from Goas is between 7.5 and 8.1% and between 6.8 and 7.8% for samples from Okongava. Most of these values are lower than those reported for other Damaran quartz diorites (7.8–9.9%; Haack et al. 1982) and other granitoids from the Damara belt (Haack et al. 1982; Jung et al. 1998b, 1999, 2001). Generally, they are more typical of mafic crustal rocks than of mafic rocks that contain a mantle component.

Discussion

The quartz diorites from Goas and Okongava are among the least evolved granitoids of the Damara orogen with low SiO₂, relatively high MgO and low Sr

isotope compositions. Geochemical features diagnostic for primary mafic magmas (e.g. Ni > 300 ppm, Cr: > 500 ppm) are not applicable for the quartz diorites because the variation in Nd, Pb and O isotope composition precludes a derivation of the quartz diorites from common upper mantle sources. In principle there are three main possibilities for the derivation of the primitive quartz diorite magmas: (1) melting of upper mantle previously enriched by subduction, (2) melting of less enriched or even depleted mantle followed by crustal contamination and (3) melting of mafic lower crust. However, Sr, Nd, Pb and O isotope compositions are heterogeneous, thus precluding a simple closed-system partial melting/fractionation model for the origin of the quartz diorite suites. Therefore, in order to constrain the partial melting process and possible sources of the quartz diorites, it is necessary to evaluate simple fractionation processes with and without assimilation of crustal material.

Fractional crystallization processes

Both suites of quartz diorites are distinct, but a general fractional crystallization trend is indicated by decreasing TiO₂, MgO, FeO, CaO, Al₂O₃, P₂O₅, V, Cr, Ni, Y, Zr, Sr, Ba concentrations and U/Pb ratios and increasing K_2O , Rb and Th concentrations, and Th/U ratios with increasing SiO_2 (Figs. 3 and 4). These trends can be interpreted to indicate fractionation of amphibole, Fe-Ti-oxides, plagioclase and minor amounts of apatite. Figure 8 is an evaluation of the most likely fractionating mineral assemblage in terms of Rb and Sr concentrations. The smooth negative correlation between Rb and Sr can be explained by fractionation of 30% amphibole and 70% plagioclase. In the quartz diorites, the negative Eu anomaly (expressed as Eu/Eu*) increases with decreasing Sr (Fig. 9) and Ga concentrations, indicating that plagioclase plays a major role during fractional crystallization. There is also a positive correlation of Ga

Table 3. Major (in wt%) and trace element (in ppm) composition of quartz diorites from the Damara orogen (Namibia). *LOI* Loss on ignition; FeO^* all Fe as Fe²⁺; *ASI* alumina saturation index; *X Mg* MgO/(MgO + FeO*); Eu/Eu^* europium anomaly; *n.d.* not determined

Location of	Goas								
sample	M20	M21	M22	M24	M25	M26	M27	M29	M30
SiO ₂	58.56	57.96	56.32	61.17	58.76	61.75	58.65	57.93	56.93
TiO ₂	0.92	0.93	1.18	0.73	0.92	0.77	0.92	0.92	1.08
Al_2O_3	17.34	17.60	17.64	16.86	17.48	16.98	17.50	17.42	16.99
Fe ₂ O ₃ FeO	1.89	2.44	2.76	1.38	2.28	2.08	2.52	2.29	2.34
FeO	5.08	4.64	5.50	4.44	4.73	3.60	4.29	4.78	5.78
MnO	0.14	0.14	0.20	0.13	0.14	0.13	0.16	0.16	0.19
MgO	2.74	2.83	3.37	2.36	2.88	2.33	2.68	3.01	3.60
CaO	6.15	6.24	6.75	5.31	5.99	5.44	5.64	6.49	6.72
Na ₂ O	2.68	2.78	3.06	2.64	2.66	2.79	2.85	2.62	2.39
K ₂ O	2.13 0.23	2.25 0.24	2.42 0.32	2.47 0.20	2.52 0.24	2.69 0.20	2.93 0.27	2.36 0.22	2.19 0.28
P ₂ O ₅ LOI	0.23	0.24	0.32	0.20	0.24 0.83	0.20	0.27 0.77	0.22	0.28 0.80
Total	99.11	99.13	100.35	98.46	99.43	99.48	99.18	99.42	99.29
Sc	13	19	20	14	14	13	16	17	20
V	113	126	136	95	116	89	115	128	163
Ċr	n.d.	n.d.	36	n.d.	44	32	39	30	43
Co	4	31	18	25	8	11	32	20	9
Ni	n.d.	n.d.	22	n.d.	18	18	21	20	27
Zn	83	83	114	71	90	76	94	86	97
Ga	20	20	22	14	19	22	24	23	19
Rb	86	94	101	99	99	100	112	92	88
Sr	517	531	506	472	503	488	487	554	511
Y	21	23	32	16	23	18	24	24	25
Zr	228	223	271	190	214	180	253	186	242
Nb	12	15	19	14	11	19	20	10	16
Ba	811	848	889	733	886	791	939	784	906
Pb	24	25	29	26	22	25	27	25	27
Th	9 3	11	8 3	12 2		15 2	13	11 4	9 2
$U Fe^{2+}/Fe^{3+}$	3 0.37	3		2 0 21		0.58	3 0.59		
Fe /Fe FeO*	0.37 6.78	0.53 6.83	0.50 7.98	0.31 5.68	0.48 6.78	0.58 5.47	0.39 6.56	$\begin{array}{c} 0.48\\ 6.84\end{array}$	$0.40 \\ 7.88$
ASI	0.78	0.85	0.88	1.01	0.78	0.98	0.30	0.84	0.92
Na_2O/K_2O	1.26	1.24	1.26	1.07	1.06	1.04	0.97	1.11	1.09
XMg	0.85	0.82	0.83	0.87	0.83	0.82	0.81	0.84	0.86
Rb/Sr	0.17	0.18	0.20	0.21	0.20	0.20	0.23	0.17	0.17
Rb/Ba	0.11	0.11	0.11	0.14	0.11	0.13	0.12	0.12	0.10
Sr/Ba	0.64	0.63	0.57	0.64	0.57	0.62	0.52	0.71	0.56
La	50.9	46.2	40.0	43.9	45.7	50.2	55.6	39.9	57.1
Ce	97.0	92.6	82.9	93.1	91.5	99.6	111.1	85.1	119.0
Nd	34.6	36.6	36.5	31.2	34.8	36.7	39.0	33.8	50.5
Sm	6.51	7.48	7.43	4.86	5.54	7.07	6.78	6.20	8.19
Eu	1.74	2.34	2.27	1.04	1.30	1.58	1.67	1.76	2.68
Gd	4.92	7.74	7.20	4.19	4.83	5.22	4.17	4.92	8.34
Dy	4.73	6.00	7.21	3.58	3.91	4.05	3.52	4.63	6.45
Er	2.80	3.00	3.48	2.18	2.20	2.20	2.00	2.76	3.19
Yb	2.20	2.40	2.78	2.03	2.10	2.00	1.78	2.30	2.61
Lu	0.31	0.34	0.37	0.30	0.30	0.30	0.24	0.35	0.40
Eu/Eu*	0.96	0.88	0.91	0.67	0.74	0.82	1.03	0.97	0.94

concentrations with the size of the negative Eu anomaly; samples with low Ga concentrations display the strongest Eu anomaly. Fractionation between LREE and HREE can also monitor fractional crystallization processes. The Ce/Yb_N ratio increases and the Yb_N concentration decreases with decreasing TiO₂ (Fig. 10), suggesting modification of the Ce/Yb_N ratio and the Yb concentration during fractional crystallization. Enrichment of LREE, together with depletion of HREE during fractional crystallization, are compatible with fractionation of amphibole, which tends to concentrate the HREE.

Contamination

From Fig. 11, it is evident that the Sr, Nd, Pb and O isotope composition show a systematic behaviour with respect to major element chemistry. Initial ⁸⁷Sr/⁸⁶Sr and δ^{18} O values increase whereas initial ϵ_{Nd} , ²⁰⁷Pb/²⁰⁴Pb and ²⁰⁸Pb/²⁰⁴Pb decrease with decreasing MgO concentration (Fig. 11). These features indicate that during crystal fractionation processes (i.e. decreasing MgO), assimilation of country rock material substantially modified the isotopic composition. There is also a positive correlation between δ^{18} O and initial ⁸⁷Sr/⁸⁶Sr and between ϵ_{Nd} and

Location of	Okongaw	va								
sample	M31	M33	M34	M35	M37	M38	M39	M40	M41	M42
SiO ₂	55.59	57.58	59.47	58.94	57.58	57.76	55.81	55.66	57.58	51.57
TiO ₂	1.17	1.08	0.87	1.04	0.97	0.99	1.07	1.08	1.12	1.29
Al_2O_3	17.77	15.05	15.78	15.02	16.62	15.84	16.52	17.22	17.04	17.53
Fe ₂ O ₃	2.32	2.58	1.91	2.41	2.41	2.37	2.13	2.72	2.97	3.72
FeO	5.71	5.13	4.42	5.27	4.61	4.84	5.69	5.32	4.51	5.78
MnO	0.17	0.18	0.13	0.17	0.14	0.14	0.15	0.14	0.14	0.18
MgO	3.13	4.04	2.84	3.52	3.37	3.69	3.78	2.98	2.95	4.66
CaO	6.40	6.64	5.42	5.80	6.14	6.19	6.88	6.47	6.15	8.35
Na ₂ O	2.93	2.39	2.89	2.38	2.73	2.71	2.70	2.91	2.91	2.78
K ₂ O	2.67	2.86	3.12	2.95	3.17	2.55	2.42	2.26	2.69	1.70
P_2O_5	0.34	0.23	0.20	0.26	0.24	0.25	0.25	0.27	0.26	0.30
LOI	0.95	0.89	2.25	1.07	0.91	1.25	1.06	1.22	0.47	0.75
Total	99.15	98.65	99.30	98.83	98.89	98.58	98.46	98.25	98.79	98.61
Sc	11	28	18	25	26	18	23	20	19	23
V	119	176	137	164	150	161	180	171	155	227
Cr	65	50	n.d.	52	62	n.d.	n.d.	n.d.	41	76
Co	4	46	35	20	22	32	43	41	16	40
Ni	35	33	n.d.	32	26	n.d.	n.d.	n.d.	21	38
Zn	116	91	71	97	86	83	90	89	87	105
Ga	23	16	18	19	22	21	17	18	21	22
Rb	100	97	118	130	117	102	91	78	94	60
Sr	489	437	447	372	494	534	485	496	480	578
Y	22	29	21	27	26	25	23	22	20	33
Zr	245	195	189	221	182	208	222	235	241	280
Nb	16	18	15	18	14	14	15	14	14	14
Ва	850	856	789	789	930	711	731	764	916	708
Pb	29	28	30	31	31	31	23	28	34	23
Th	10	16	21	17	12	16	13	10	14	10
U	3	2	2	3	2	2	3	2	3	2
Fe^{2+}/Fe^{3+}	0.41	0.50	0.43	0.46	0.52	0.49	0.37	0.51	0.66	0.64
FeO*	7.80	7.45	6.14	7.44	6.78	6.97	7.60	7.77	7.18	9.12
ASI	0.92	0.79	0.88	0.85	0.87	0.86	0.84	0.91	0.90	0.81
Na ₂ O/K ₂ O	1.10	0.84	0.93	0.81	0.86	1.06	1.12	1.29	1.08	1.64
XMg	0.84	0.86	0.85	0.85	0.85	0.86	0.88	0.81	0.80	0.83
Rb/Šr	0.20	0.22	0.26	0.35	0.24	0.19	0.19	0.16	0.20	0.10
Rb/Ba	0.12	0.11	0.15	0.16	0.13	0.14	0.12	0.10	0.10	0.08
Sr/Ba	0.58	0.51	0.57	0.47	0.53	0.75	0.66	0.65	0.52	0.82
La	68.8	42.6	43.7	50.5	29.5	34.6	35.3	40.7	48.1	35.6
Ce	119.5	89.2	84.3	102.3	69.8	70.9	73.7	84.9	97.3	73.2
Nd	41.8	39.5	33.3	45.4	30.8	32.4	33.2	33.7	36.6	33.6
Sm	8.09	9.32	6.05	8.23	6.66	6.47	6.74	6.40	7.78	6.70
Eu	1.99	1.88	1.43	1.36	1.13	1.43	1.38	1.49	1.91	1.86
Gd	5.56	7.56	5.52	6.27	4.92	5.63	5.15	5.43	6.41	6.07
Dy	4.55	5.86	3.54	4.48	4.24	4.31	4.78	4.31	4.84	5.04
Er	2.38	2.93	1.80	2.10	2.04	2.20	2.50	2.10	2.50	2.50
Yb	2.50	2.44	1.50	2.00	1.68	1.60	2.00	1.80	2.00	2.40
Lu	0.40	0.37	0.25	0.36	0.24	0.25	0.26	0.25	0.31	0.35
Eu/Eu*	0.97	0.71	0.72	0.60	0.62	0.71	0.73	0.76	0.84	0.86
	0.27	5.71	5.72	5.00	5.02	5.71	5.75	5.70	5.61	0.00

Table 4. Major (in wt%) and trace element (in ppm) composition of quartz diorites from the Damara orogen (Namibia). *LOI* Loss on ignition; *FeO** all Fe as Fe^{2+} ; *ASI* alumina saturation index; *XMg* MgO/(MgO + FeO*); *Eu/Eu** europium anomaly; *n.d.* not determined

 207 Pb/ 204 Pb, and a negative correlation between δ^{18} O and 207 Pb/ 204 Pb and ϵ_{Nd} and also between 87 Sr/ 86 Sr and ϵ_{Nd} , which further indicate contamination processes.

The composition of the quartz diorites and two different Damara country rocks (Jung et al. unpublished) are shown in a diagram of ⁸⁷Sr/⁸⁶Sr vs Sr concentration (Fig. 12). The country rocks are a typical Damara metapelite (⁸⁷Sr/⁸⁶Sr: 0.720, Sr: 200 ppm) and an evolved basement gneiss of inferred Proterozoic to Archaean age (⁸⁷Sr/⁸⁶Sr: 0.740, Sr: 500 ppm) because it can be expected that such lithologies are the most fusible parts of a crustal segment and, thus, are most likely involved in any assimilation scenario. The most mafic quartz diorite with ca. 50 wt% SiO₂ has ca. 600 ppm Sr, a primitive Sr isotopic composition and only a small negative Euanomaly; therefore, it is considered to represent one of the least contaminated end members. The most likely AFC curves to fit the data are those of the metapelitic end member with 200 ppm Sr and ⁸⁷Sr/⁸⁶Sr: 0.7200 (case A) and the pre-Damara orthogneiss (case B). Decreasing Al₂O₃, Sr and Ga concentrations and increasing Eu/Eu* shows that plagioclase strongly controls the fractionating assemblage and, therefore, the bulk K_d for Sr has to be >1. The value of r (r= ratio of assimilation to

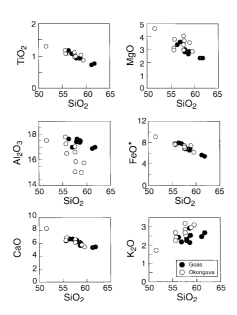


Fig. 3. Major element plots for both types of quartz diorite

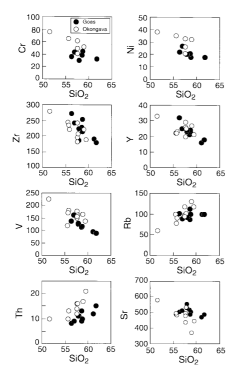


Fig. 4. Selected trace element plots for both types of quartz diorite

fractionation) was set at 0.8 and the amount of assimilation is estimated to be <10%, suggesting that only a small fraction of crustal material was assimilated. In contrast to previous estimates for the ratio of mass assimilated to mass fractionated (r: <<1; DePaolo 1981) higher values of r (r: 1-2; Reiners et al. 1995) may be possible. High values of r further suggest that assimilation took place in the deep crust at high temperatures.

Processes of crustal assimilation can also be illustrated by using different isotope systems. Figure 13

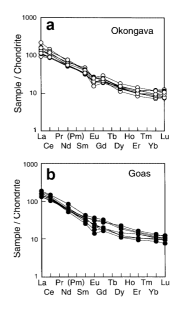


Fig. 5. Chondrite-normalized rare earth element plots for both types of quartz diorite. Normalization factors according to Boynton (1984)

shows that the negative correlation between Sr and Nd and the positive correlation between Sr and O isotope composition is best monitored by using an orthogneiss from the pre-Pan African basement as the contaminant. Lead isotope compositions are not available from basement gneisses; however, the Pb isotope composition is particularly susceptible to extreme changes during AFC as a result of the high concentration of Pb in felsic crustal rocks relative to mafic rocks. Therefore, it is very likely that compositional differences in the contaminant are responsible for changes in Pb isotope composition. Lead isotope composition correlates with Nd and Sr isotope compositions and increasing contamination is accompanied by decreasing $\epsilon_{\rm Nd}$, increasing ${}^{87}{\rm Sr}/{}^{86}{\rm Sr}$, but decreasing ${}^{207}{\rm Pb}/{}^{204}{\rm Pb}$ and ${}^{208}{\rm Pb}/{}^{204}{\rm Pb}$ ratios. This relationship indicates that the source of the contaminant has a lower μ and κ value than the quartz diorites.

Partial melting processes and magma sources

Sill-like intrusions are thought to be a major component of mid- to lower crustal regions (Bohlen 1987) although, in the Damara orogen, geophysical evidence (Green 1983) argue against the existence of large volumes of mafic rocks in the lower crust. However, recent work by Petford and Gallagher (2001) indicates that these sills need not be very thick in order to melt mafic lower crust. Experimental evidence (Helz 1976; Beard and Lofgren 1991; Rapp et al. 1991; Rushmer 1991; Wolf and Wyllie 1994; Rapp and Watson 1995) and theoretical considerations (Roberts and Clemens 1993) suggests that dehydrating basaltic material within the lower crust can produce significant volumes of mafic partial melts, particularly in regions with high heat flow. Depending on

e ⁸⁷ Sr/ ⁸⁶ Sr values of values of	²⁰⁸ Pb/ ²⁰⁴ Db/
nties for th resent-day esent-day	$^{207}_{204} Pb/$
Uncertain JR with p air and pr	$^{206}_{204}Pb/$
echniques /e to CHU tle reservo	δ^{18} O
uted relativ ted relativ eted mant	Sr (mmm)
mibia. <i>n.d.</i> Not determined. See <i>A</i> ion (Tables 3 and 4) are calcula lel ages calculated with a depl	87 Rb/ ⁸⁶ Sr 87 Sr/ ⁸⁶ Sr Rb
Table 5. Rb–Sr, Sm–Nd, Pb and O isotope data from quartz diorites from the Damara orogen Namibia. <i>n.d.</i> Not determined. See Analytical techniques. Uncertainties for the ⁸⁷ Sr/ ⁸⁶ Sr and ¹⁴³ Nd/ ¹⁴⁴ Nd are 2σ (mean) errors in the last two digits. ϵ Nd values at the time of intrusion (Tables 3 and 4) are calculated relative to CHUR with present-day values of ¹⁴³ Nd/ ¹⁴⁴ Nd = 0.512638 and ¹⁴⁷ Sm/ ¹⁴⁴ Nd = 0.1966 (Jacobson and Wasserburg 1980). Nd model ages calculated with a depleted mantle reservoir and present-day values of ¹⁴³ Nd/ ¹⁴⁴ Nd = 0.5121648 and ¹⁴⁷ Sm/ ¹⁴⁴ Nd = 0.512164 and ¹⁴⁷ Sm/ ¹⁴⁴ Nd = 0.222 (Michard et al. 1985). <i>m</i> Measured; <i>i</i> initial	ϵ Nd T _{DM} ⁸⁷ Sr/ ⁸⁶ Sr (i) (Ga) (m)
from quartz d ast two digits. 966 (Jacobson 2 (Michard et	$^{143}_{144}{ m Nd}/$
otope data ors in the 1 $^{44}Nd = 0.1$ $^{4}Nd = 0.22$	(maa)
b and O is nean) erro d ¹⁴⁷ Sm/ ¹ d ¹⁴⁷ Sm/ ¹⁴	Sm (mmm)
Sm–Nd, P 1 are 2σ (r 512638 an 513144 an	$^{143}_{144}Nd/$
Table S. Rb–Sr, Sm–Nd, Pb and O isotope data from quartz di and ¹⁴⁵ Nd/ ¹⁴⁴ Nd are 2σ (mean) errors in the last two digits. ¹⁴³ Nd/ ¹⁴⁴ Nd = 0.512638 and ¹⁴⁷ Sm/ ¹⁴⁴ Nd = 0.1966 (Jacobson ¹⁴³ Nd/ ¹⁴⁴ Nd = 0.513144 and ¹⁴⁷ Sm/ ¹⁴⁴ Nd = 0.222 (Michard et at a second	Sample ${}^{147}_{144NIA}Sm/ {}^{143}Nd/ {}^{144}_{NIA}$

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$^{204}\overline{Pb}$	37.804	37.901	37.903	37.707	n.d.	37.726	37.774	37.853	37.799	37.797	38.156	38.053	37.994	38.107	38.075	38.051	38.051	37.857	37.957	
$^{204}\tilde{Pb}$	15.556	15.553	15.563	15.535	n.d.	15.541	15.556	15.546	15.545	15.576	15.611	15.586	15.572	15.593	15.591	15.625	15.594	15.558	15.615	
$^{204}\tilde{\mathbf{P}}\mathbf{\dot{b}}$	17.223	17.210	17.230	17.180	n.d.	17.185	17.172	17.170	17.157	17.119	17.121	17.126	17.109	17.139	17.161	17.170	17.110	17.083	17.160	
(‰)	7.78	7.65	7.50	7.90	8.00	8.02	7.75	8.14	7.90	7.50	7.73	7.80	7.50	7.20	7.20	6.80	7.45	7.43	6.90	
(mqq)	532.1	545.9	504.6	487.2	503.5	497.2	494.0	549.1	524.0	480.2	439.9	431.4	391.6	516.1	540.8	485.1	491.6	482.6	609.5	
(mdd)	87.20	89.72	98.91	100.9	96.85	99.84	108.2	92.22	87.54	100.0	102.0	120.3	126.8	118.7	96.29	88.53	72.89	91.17	61.13	
(i)	0.712412	0.712449	0.712254	0.711189	0.712848	0.711233	0.711340	0.711211	0.710807	0.712509	0.710776	0.711057	0.710839	0.711123	0.709429	0.710069	0.711867	0.712185	0.710319	
	0.4745	0.4759	0.5675	0.5993	0.5013	0.5814	0.6340	0.4862	0.4836	0.6029	0.6700	0.8072	0.9375	0.6657	0.5154	0.5283	0.4292	0.5470	0.2903	
(m)	0.716132 (16)	\sim	\sim	0.715888 (14)	\sim															
(Ga)	1.9	2.0	2.2	1.9	2.0	2.0	2.0	2.1	2.0	2.1	1.9	1.8	1.7	2.0	1.7	1.8	2.0	2.0	1.8	
(i)	-13.0	-13.7	-13.2	-14.6	-14.6	-14.1	-14.6	-13.7	-12.8	-15.2	-10.5	-10.5	-10.5	-10.8	-9.1	-9.2	-13.8	-13.9	-9.7	
(i)	0.511264 (12)	\sim	\sim	0.511183(13)	\sim															
(mqq)	39.56	39.19	36.01	30.74	37.83	41.40	40.30	32.74	49.24	45.22	42.86	34.29	53.84	31.39	34.18	35.22	36.70	39.11	34.38	
(mqq)	6.724	6.981	7.349	4.926	6.563	7.303	6.801	6.214	8.362	8.041	8.499	6.137	9.199	6.375	6.281	6.753	6.657	7.147	6.520	
(m)	0.51163	0.51162	0.51170	0.51153	0.51156	0.51159	0.51155	0.51164	0.51167	0.51154	0.51825	0.51178	0.51176	0.51182	0.51186	0.51188	0.51162	0.51162	0.51185	
¹⁴⁴ Nd	0.1027	0.1077	0.1234	0.0968	0.1049	0.1066	0.1020	0.1147	0.1093	0.1075	0.1199	0.1082	0.1033	0.1227	0.1111	0.1159	0.1069	0.1105	0.1146	
	M 20	M 21	M 22	M 24	M 25	M 26	M 27	M 29	M 30	M 31	M 33	M 34	M 35	M 37	M 38	M 39	M 40	M 41	M 42	

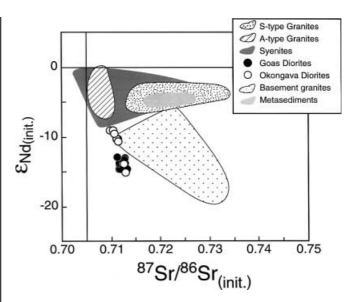


Fig. 6. Initial $\epsilon_{\rm Nd}$ versus initial $^{87}{\rm Sr}/^{86}{\rm Sr}$ diagram for quartz diorites

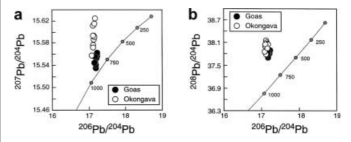


Fig. 7. Plot of **a** ${}^{207}\text{Pb}/{}^{204}\text{Pb}$ and **b** ${}^{208}\text{Pb}/{}^{204}\text{Pb}$ versus ${}^{206}\text{Pb}/{}^{204}\text{Pb}$ isotope ratios of leached K-feldspar from quartz diorites. The curve represent the average Pb growth curve according to Stacey and Kramers (1975). *Tick marks* represent 250-Ma intervals

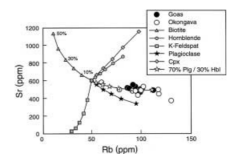


Fig. 8. Variation in Rb and Sr concentration within the quartz diorites. *Symbols* as in Fig. 3. Mineral vectors calculated according to partition coefficients compiled in Rollinson (1993). Compositional trend of the quartz diorites indicate fractionation of ca. 30 wt% of plagioclase and ca. 70 wt% of hornblende

bulk composition, dehydration melting of amphibolite yields 10-60% quartz diorite to tonalite melt at temperatures of 900–1,000 °C, whereas H₂O-saturated melting yields similar amounts of granite melt at lower temperatures of between 850 and 900 °C.

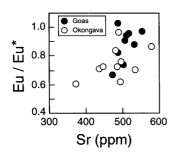


Fig. 9. Plot of Eu/Eu* as a measure of the negative Eu anomaly vs. Sr concentrations for both suites of quartz diorites. For discussion see text

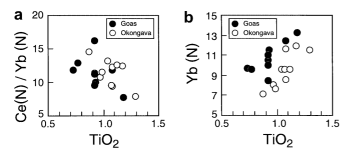


Fig. 10. Plot of a Ce/Yb_N and b Yb_N vs TiO₂. Correlation between the parameters indicate strong influence of crystal fractionation processes upon REE systematics

Some of the quartz diorites have chemical features that are characteristic of partial melts derived from metabasaltic rocks. They have low SiO₂, high Al₂O₃, moderately high Na₂O/K₂O, high LREE concentrations, moderately low HREE concentrations and the most primitive rocks do not display a negative Eu anomaly. Based on previous work (Rapp et al. 1991; Wolf and Wyllie 1993; Rapp and Watson 1995), the highest pressure partial melts should have the lowest Y and HREE concentrations, but the highest Al₂O₃ and Sr

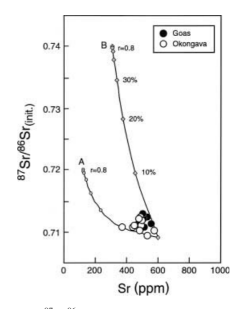
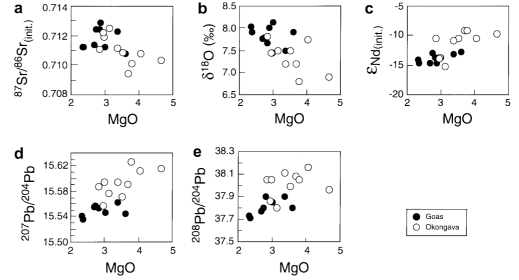


Fig. 12. Plot of ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ vs. Sr concentration for quartz diorites and calculated AFC curves using two hypothetical country rocks (*A* pelite with 200 ppm Sr and ${}^{87}\text{Sr}/{}^{86}\text{Sr}$: 0.7250, and *B* metaigneous basement gneiss with 500 ppm Sr and ${}^{87}\text{Sr}/{}^{86}\text{Sr}$: 0.7400). Bulk K_D (Sr) was 1.5. The hypothetical source rock contains 600 ppm Sr and ${}^{87}\text{Sr}/{}^{86}\text{Sr}$: 0.7090 similar to the most primitive quartz diorite sample M42

concentrations and the highest Sr/Y ratios. For the quartz diorites from the Damara orogen, those with the lowest Y concentration and highest Sr/Y have the highest Al_2O_3 and least negative Eu anomaly. All these characteristics would be expected for partial melts from a basaltic protolith, most probably a garnet-bearing amphibolite at high pressures. An amphibole-bearing residue for the quartz diorites of the Damara orogen is consistent with the low TiO₂ concentrations and low Rb/ Sr and K/Rb ratios, despite their high absolute K and Rb concentrations (Petford and Atherton 1996).

Fig. 11. Variation of 87 Sr/ 86 Sr, δ^{18} O, ϵ Nd, 207 Pb/ 204 Pb, 208 Pb/ 204 Pb vs. MgO. The trends are best explained by assimilation of evolved country rocks during fractional crystallization. Note that the contaminant must be more evolved in Sr, Nd and O isotope composition, but must have lower μ and κ values



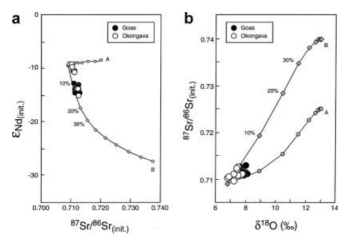


Fig. 13. Plot of **a** ⁸⁷Sr/⁸⁶Sr vs. $\epsilon_{\rm Nd}$ and **b** ⁸⁷Sr/⁸⁶Sr vs. δ^{18} O for quartz diorites and calculated AFC curves using the same two hypothetical country rocks as in Fig. 12. *A* pelite with 200 ppm Sr ⁸⁷Sr/⁸⁶Sr: 0.7250, $\epsilon_{\rm Nd}$: -8.4, δ^{18} O: 13% and *B* meta-igneous basement gneiss with 500 ppm Sr, ⁸⁷Sr/⁸⁶Sr: 0.7400, $\epsilon_{\rm Nd}$: -29, δ^{18} O: 13% Bulk K_D (Sr) was 1.5 and bulk K_D (Nd) was 0.8. A *r*-value (ratio of assimilation to fractionation) of 0.8 was used. The hypothetical source rock has 600 ppm Sr, ⁸⁷Sr/⁸⁶Sr: 0.7090, $\epsilon_{\rm Nd}$: -9.4, δ^{18} O: 6.8% similar to quartz diorite sample M42

The quartz diorites from the Damara orogen have $K_2O > 2.5$ wt%, MgO > 3 wt%, CaO > 6 wt%, FeO_{total} > 7 wt% and TiO₂ > 1.0 wt%. Such chemical characteristics are observed only in experiments performed between 1,000–1,100 °C at high pressures (Fig. 14). Abundances of MgO are similar in the quartz diorites and in the experimental melts derived by melting of low

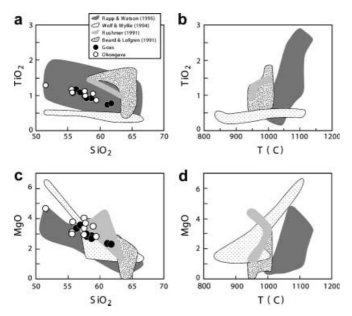


Fig. 14. Plot of **a** TiO_2 and **b** MgO vs. SiO_2 , **c** TiO_2 and **d** MgO vs. temperature for quartz diorites and experimental investigations. Note that the composition of the quartz diorites is likely reproduced by high-temperature melting of a source similar to the low-K (medium-K?) ol-tholeiite of Rapp and Watson (1995). For further discussion see text

K-basaltic rocks (Wolf and Wyllie 1994; Rapp and Watson 1995), but substantially higher than in melts generated by melting of alkali basaltic and high-Al basaltic sources. Similarly, the high CaO concentrations in the quartz diorites precludes alkali basaltic sources and the low TiO_2 abundances suggests that high-Al basaltic sources are also unlikely sources. Overall, the source rock of the quartz tholeiites is most likely a high-grade metamorphic, medium-K basaltic rock that melted at high temperatures and high pressures in the lower crust.

At 10-15 kbar, calc-alkaline liquids coexist with garnet under both vapour-saturated and undersaturated conditions (Huang and Wyllie 1986; Rutter and Wyllie 1988; Carrol and Wyllie 1990). None of the most primitive quartz diorites show the significant HREE depletion (i.e. Yb_N < 10; Wareham et al. 1997) predicted for melts that equilibrated with residual garnet, suggesting that they formed alternatively under lower pressures, with less garnet in the residuum or that garnet was melted to a large extent during formation of the melts. However, the high temperatures required for amphibolite melting restrict this process to the deep crust. Temperatures in the deep crust during generation of the quartz diorites cannot have exceeded 1,070 °C at 10-15 kbar, which is the thermal stability limit of amphibole in basalt (Allen and Boettcher 1978). Comparison with experimental results (Fig. 14) support the idea of high-temperature generation of the quartz diorites because the major element concentrations are only reproduced in high-temperature runs between 1,000-1,100 °C. The temperature at which a mafic melt separated from its source may be estimated from its P₂O₅ concentration, using the apatite solubility expression of Harrison and Watson (1984). This approach assumes that the melt formed in equilibrium with residual apatite and has not undergone subsequent modification by processes related to fractional crystallization or assimilation. The first requirement may be satisfied during melting, especially at low degrees of melting, but the second requirement is more difficult to evaluate. However, the most mafic quartz diorites have the most primitive isotopic composition indicating less modification by AFC processes. For these samples P_2O_5 concentrations are between 0.32-0.34 wt%, indicating temperatures of between 1,060 and 1,070 °C, similar to those given by the experimental investigations.

Summary and discussion of model

Miller (1983) and Kasch (1983) used the occurrence of quartz diorites as an argument for an island arc setting of the southern part of the Central Zone of the Damara orogen. Despite the striking absence of geological evidence for a subduction zone setting, it is pertinent to compare the chemistry of the quartz diorites with that of magmatic rocks from arc environments. The low concentrations of Ni and Cr argue against a pure mantle derivation of the quartz diorites; however, similar low abundances have been reported in the least evolved members of some island arc tholeiite and calc-alkaline series (e.g. Perfit et al. 1980). A number of trace element characteristics are typical for arc magmatism, i.e. enrichment of LILE (Ba, Rb, K), relative depletion of HFSE (e.g. Nb) and, consequently, high LILE/HFSE ratios. Enrichment of LILE might be caused by transfer of volatiles from dehydrating subducted oceanic crust to the mantle wedge (e.g. Hawkesworth et al. 1979a, 1979b; Thirlwall and Graham 1984, among many others). In the absence of a subduction zone setting, enrichment of LILE may be caused by preferential tapping of enriched mantle sources (e.g. Saunders et al. 1980) or may be caused by AFC (assimilation-fractional crystallization) or MASH (melting-assimilation-storage-homogenization) processes (e.g. Hildreth and Moorbath 1988). Despite the striking evidence for AFC processes, the least evolved quartz diorite samples have rather low SiO₂ contents, but unradiogenic Nd and radiogenic Sr isotope composition. While large-scale crustal contamination of magma derived from upper mantle sources could explain some of the isotope variation, such contamination would likely be accompanied by strong fractional crystallization, which is not observed. Furthermore, the strongly retarded Pb isotope compositions precludes their derivation from a common or even moderately enriched upper mantle source at 550 Ma. The Sr-Nd isotope composition of the quartz diorites is unlike the Sr-Nd isotope composition of other Damara granites for which a upper crustal derivation (McDermott et al. 1996; Jung et al. 2001), a lower crustal derivation (Jung et al. 2002) or even a contribution from the lithospheric mantle (Jung et al. 1998b) has been inferred. It is also unlike the Nd and Sr isotope composition of other quartz diorites from the Damara orogen. Although similar in age (ca. 550 Ma), these rocks are situated farther north and have different isotope compositions $({}^{87}\text{Sr}/{}^{86}\text{Sr}: 0.7045 - 0.7053; \epsilon_{Nd}: -0.1; \overline{\delta}{}^{18}\text{O}: 7.8 - 9.1\%$ Hawkesworth et al. 1981; Haack et al. 1982). Therefore, it is very likely that the isotope composition of the quartz diorites studied here matches the composition of the mafic lower crust beneath the Damara orogen.

Previous studies have shown that Pb isotopes from leached feldspars provide useful information for the understanding of crustal evolution and often record evidence for an ancient crustal memory. Magmatic differentiation and metamorphic events can produce significant variations in μ and κ values, which lead to highly variable Pb isotope ratios. Lead isotope ratios from leached feldspars are useful, because feldspars have low μ and κ values and their Pb isotope compositions can be used to estimate the initial Pb isotope ratios at the time of the last equilibration event. The quartz diorites have the most retarded Pb isotope compositions so far recorded in Damaran plutonic rocks. The samples plot well above the Stacey and Kramers (1975) Pb evolution curve at a Pb model age of ca. 1 Ga. Although strongly retarded, these isotopic features suggest that the diorites cannot come from a MORB-type source. The large

variation in ²⁰⁷Pb/²⁰⁴Pb relative to ²⁰⁶Pb/²⁰⁴Pb indicate the involvement of old reworked crust and is probably also a result of the influence of an ancient sedimentary component. High-grade metamorphism during ancient events in deep crustal sources can lead to strongly retarded Pb isotope compositions because of fractionation of U from Pb and Th from U. Although both elements, U and Th, are highly incompatible, U is more soluble than Th and is preferentially incorporated into a fluid phase whereas Th is practically insoluble in a fluid phase, but strongly partitioned into a melt (e.g. Taylor and McLennan 1985). Therefore, it is very likely that the observed Pb isotope composition is the result of both high-grade metamorphism and melting. Considering the retarded composition these depletion events must be significantly older than 550 Ma.

The most plausible model for the generation of the quartz diorites that accounts for the observed geochemical features involves partial melting of pre-Pan African mafic lower crust probably because of underplating of mantle-derived magmas. Melting of this lower crust would be most efficient if the partial melting event immediately followed the underplating event and probably close to the onset of high-grade metamorphism because high temperatures are required for such melting events (Petford and Gallagher 2001). It is important to note that the oldest Sm-Nd whole rock garnet ages and U-Pb monazite ages are ca. 540 Ma for the central part of the Damara orogen (Jung and Mezger 2000; Jung et al. 2000b; Jung and Mezger 2002). Therefore, it is reasonable to assume that the crust was already hot at 540 Ma, but it is also very likely that similar quartz diorites are a precursor to high-grade metamorphism, melting and granite formation. Intrusion of large volumes of basaltic rocks are not reported from the Damara orogen, but close to the occurrence of the quartz diorites and a number of independent syenites, some of them ne-normative, intruded the area close to a major shear zone. This shear zone, or parts of it, might have promoted the ascent of mantle-derived syenites, but also lower crustal melts at a time when most of the orogen underwent compression because of thickening and thrusting. New high precision U-Pb titanite ages seem to indicate that the syenites intruded between 590 and 490 Ma; however, the majority of the samples record ages between 520 and 540 Ma (Jung et al. 1998a: unpublished data). This observation outlines a thermal role for mantle-derived magmas in the production of lower crustal melts, but also implies that thick and hot crust is a requirement for the generation of quartz diorites from lower crustal sources. In this model, the crust acts as a barrier to magmas ascending from the mantle, causing them to pond and crystallize, and leading to the development of a zone of lower crustal melting. The overlying crust must have been thick, probably because of thickening during the early stages of orogeny because the quartz diorites evolved by AFC rather than pure FC processes. During intrusion, the quartz diorites assimilated crustal material, which was more radiogenic in terms of the Sr isotope composition, but less radiogenic in terms of Nd and Pb isotope composition (Fig. 6). These Sr and Nd isotope features are so far only monitored by pre-Pan African basement gneisses as inferred from the isotope composition of basement-derived granites (Jung et al. 2002). The correlation between Sr, Nd, Pb and O isotopes and MgO concentration suggests that during evolution of the quartz diorites more and more evolved (older?) rocks contributed to the isotopic signature of the melts. This behaviour is the opposite of what is expected if an igneous body moves from the site of melting through a crust with the oldest, most evolved rocks underlying the site of melt generation. Therefore, it is reasonable to assume that the area consists of an inverted stratigraphy at depth, in which crustal slices of pre-Pan African basement are intercalated with Pan African rocks. A similar scenario has been suggested for other basement-derived granites elsewhere in the Damara orogen (Jung et al. 2002).

Concluding remarks

Quartz diorites from Okongava and Goas (Damara orogen, Namibia) are the most mafic, but SiO₂-saturated rocks so far discovered in the orogen. Based on temperature estimates based on P₂O₅ solubility in mafic rocks and comparison with results from amphibolitedehydration melting experiments, melting of a medium-K garnet-bearing metatholeiite at temperatures of between 1,000 and 1,100 °C is likely. The major- and trace element variation is best explained by crystal fractionation processes involving mainly plagioclase (ca. 70%) and hornblende (ca. 30%). Isotope variation suggests that the quartz diorites were modified by AFC processes. Increasing initial Sr isotope ratios, but decreasing Nd isotope ratios, indicate that the contaminant is probably a felsic meta-igneous basement gneiss. The model calculations of AFC require moderately high K_d Sr (>1), but low K_d Nd (<1), which is supported by the inferred fractionated mineral assemblage of plagioclase and hornblende. However, during AFC processes $^{207}Pb/^{204}Pb$ and $^{208}Pb/^{204}Pb$ ratios decrease with increasing degree of assimilation, suggesting that the source of the contaminant has lower μ and κ values. A high ratio of mass assimilated to mass fractionated further supports lower crustal AFC processes at high temperatures. Although the lower Paleozoic was a time during which a large amount of igneous material was introduced into the upper crust of the Damara orogen it was not a major crust-forming period because most granites studied so far, and also the most mafic rocks such as the quartz diorites studied here, are dominated by recycled crustal material.

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