

Whole-rock geochemistry of the Hili Manu peridotite, East Timor: implications for the origin of Timor ophiolites*

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The Hili Manu peridotite occupies a key position at the outer limit of continental crust on the north coast of East Timor. Most models for the tectonic evolution of the Outer Banda Arc interpret peridotite bodies on Timor, such as Hili Manu, as fragments of young oceanic lithosphere from the Banda Arc (upper plate). However, recent workers have used major-element geochemistry to argue that the peridotite bodies on Timor were derived from the Australian subcontinental lithosphere. Our major, trace and isotopic geochemical study of the Hili Manu peridotite body supports a supra-subduction origin from either a forearc or backarc position for the Hili Manu peridotite. In particular, the wide range in Nd and Sr isotopic compositions, overlapping that of arc volcanics from the Sunda–Banda Island arc, and highly fractionated Nb/Ta values indicate a supra-subduction setting. As there is no evidence for subduction beneath the rifted Australian continental margin, it is unlikely that the Hili Manu peridotite is Australian subcontinental lithosphere. This result, along with the clear supra-subduction setting of the Ocuzy peridotite and associated volcanics in West Timor, gives support to the interpretation that the Miocene collision between the Banda Arc and the Australian continental margin has produced widespread 'Cordilleran'-style ophiolites on Timor.

KEY WORDS: Australia, geochemistry, Hili Manu, ophiolite, peridotite, Timor.

INTRODUCTION

Fault-bounded peridotite bodies are common in suture zones and along major transcurrent faults. However, the tectonic significance of these bodies is debatable (Coleman 1971; Hamilton 1979; Dilek *et al.* 2000; Beccaluva *et al.* 2004). In East Timor, a peridotite body outcrops as a fault block on the north coast, on the margin of continental crust in a modern island arc–continent collision zone. This body is of particular interest, because later events have not obscured its position in the orogenic belt, and also because it has retained its high-temperature mineralogy. The peridotite (Hili Manu peridotite) is remarkably fresh and has a complex history of high-temperature processes recorded in its disequilibrium textures and mineral assemblage.

The most common rock type in this body is a clinopyroxene-poor lherzolite, but there are smaller proportions of clinopyroxene-rich lherzolite and harzburgite. The dominant mineral assemblage is olivine, orthopyroxene, clinopyroxene, spinel and calcic amphibole. Low-temperature hydrous minerals are restricted in distribution. The chemical composition of the peridotite is very similar to mantle-derived spinel lherzolite

nodules and some alpine peridotites (Berry 1981). The internal variation of the peridotite suggests variable depletion by some combination of partial melting and liquid contamination of the residua.

Berry (1981) recognised three solid-state events from the mineral chemistry and texture. The earliest event is recorded by coarse exsolution lamellae of orthopyroxene in clinopyroxene porphyroclasts. These grains formed at 1250°C. A later granuloblastic texture equilibrated at 1100°C, and finally the rocks were mylonitised at 800–1000°C and 800–2000 MPa.

The Hili Manu peridotite body could be part of either the Banda upper plate (the Banda terrane) or the lower Australian Plate. If the Hili Manu peridotite is derived from the Banda Plate, we should expect to see a supra-subduction zone geochemical signature, as the Banda terrane is made up of volcanic and sedimentary sequences associated with the Sunda and Banda volcanic arcs that overlie a polyphase metamorphic complex (Audley-Charles & Harris 1990). In contrast to the Banda terrane, the lower Australian Plate is made up of passive-margin rift-related sequences associated with the breakup of Gondwanaland beginning in the Permian (O'Brien *et al.* 1993; Harris *et al.* 1998). As there is no

Appendix 1 [indicated by an asterisk () in the text and listed at the end of the paper] is a Supplementary Paper; copies may be obtained from the Geological Society of Australia's website (<www.gsa.org.au>) or from the National Library of Australia's Pandora archive (<<http://nla.gov.au/nla.arc-25194>>).

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evidence for any active subduction beneath this margin, we would not expect to find any subduction zone geochemical features in peridotite bodies derived from the lower Australian Plate, as a result of Miocene collision. Harris and Long (2000) argued on the basis of major elements only that the Hili Manu peridotites were relatively fertile compared with refractory forearc peridotites (Parkinson & Pearce 1998) and therefore most likely to be derived from the lower Australian Plate. In contrast, Berry (1981) argued the Hili Manu peridotite is a sample of the oceanic mantle trapped between the Java Trench and the Inner Banda Arc and hence is part of the upper Banda Plate. The aim of this report is to use the detailed trace-element and isotopic composition of the Hili Manu peridotite to resolve these conflicting interpretations.

REGIONAL SETTING

The island of Timor was formed by the Miocene collision between the northwest margin of the Australian continent and an island arc–subduction complex (Berry & McDougall 1986). Timor is separated from the North West Shelf of Australia by a 3-km-deep trough that is underlain by crustal rocks characteristic of continental areas (Curry *et al.* 1977). Gravity modelling confirms that continental crust continues to the northern edge of Timor (Chamalaun *et al.* 1976). About 100 km north of Timor is a line of volcanic islands known as the inner Banda Arc. This arc is continuous with the volcanic chain through Java and Sumatra

(Sunda Arc) and has active andesitic volcanoes along most of its length, except north of Timor.

Timor is presently rising rapidly (Chappell & Veeh 1978). The geological structure is dominated by strong faulting, and a large range of rock types occurs on the island, commonly as small, fault-bounded blocks. Many are similar to rocks of the same age on the continental shelf to the south, whereas others have been correlated with lithological associations found in Sundaland (Carter *et al.* 1976; Milsom 2000) or have been identified as abyssal sediments.

The steep gradient in the gravity field across Timor is due to the rapid transition from continental to oceanic crust (Chamalaun *et al.* 1976). In addition to this north–south gradient, a local gravity high occurs 30 km east of Dili (Milsom & Richardson 1976) centred on the Hili Manu peridotite. The extent of this anomaly suggests a substantial extension of ultramafic rocks at depth.

GEOLOGY

The Hili Manu peridotite body abuts a large block of amphibolite-facies schists, the Aileu Formation, along a near-vertical fault on its southern margin (Figure 1). To the east, there is also a small block of amphibolite of ambiguous association (Berry 1979; Harris & Long 2000), but the nature of this eastern contact is unknown. Berry and McDougall (1986) suggested that this massive amphibolite could be correlated with the Aileu Formation, but recently Charleton (2002) interpreted the amphibolite as allochthonous.

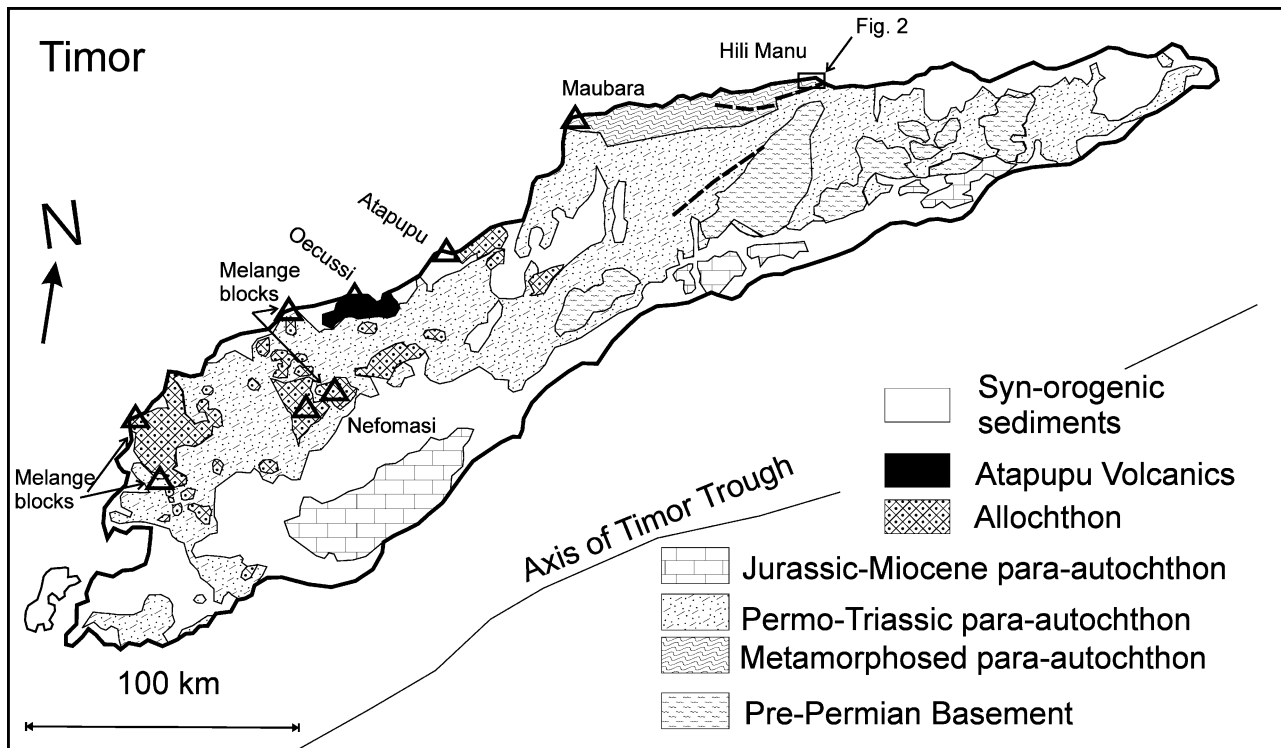


Figure 1 Location and field relationships of the Hili Manu peridotite, East Timor (geology slightly modified from Charleton 2002).

The most common rock type of the Hili Manu peridotite body is a very fine-grained, brown-weathering lherzolite, in which orthopyroxene occurs as large, tabular porphyroclasts standing out on the weathered surface. A compositional layering, commonly 2–10 cm thick, is defined by concentrations of orthopyroxene. The layering dips at a moderate angle to the south. There is also a strong foliation subparallel to the layering, which is defined by the dimensional preferred orientation of the tabular orthopyroxene grains. In a few areas, the peridotite is altered to massive, pale-green or brown serpentinite.

For this project, we have reanalysed the original samples reported by Berry (1981). These samples have been moved to the University of Tasmania (UTAS) and are reported here with the new UTAS catalogue numbers.

SAMPLE SELECTION AND ANALYTICAL METHODS

The samples are the same as those of Berry (1981) and were collected from the peridotite where it crops out along the road and along the foreshore (Figure 2). Samples were ground in an agate mill to avoid any possible trace-element contamination. Some of the samples collected were from boulders along the foreshore. Major- and some trace-element analyses (Table 1) were performed at the School of Earth Sciences (UTAS) using X-ray fluorescence spectrometry (XRF) and the methods of Robinson (2003). Additional trace-element geochemistry (Table 2) was performed at the School of Earth Sciences and Central Science Laboratories (UTAS) using inductively coupled plasma–mass spectrometry (ICP-MS) and the methods of Robinson *et al.* (1999) and Yu *et al.* (2000). ICP-MS analyses were performed on duplicate high-pressure HF–HClO₄ digestions. Sub-boiling double-distilled

acids and ultra-pure water were used, as were clean sampler and skimmer cones, ICP torch, spray chamber, nebuliser and sample-introduction tubes (including auto-sampler tubing). Prior to sample analysis, the instrument was purged for at least 24 h with 5% v/v HNO₃ and 0.05% v/v HF rinse solution. The international peridotite standard PCC-1 was also analysed along with the Hili Manu samples, and our results for this standard are compared with literature values in Appendix 1*. Isotope analyses (Table 3) were performed at the Max Planck-Institut für Chemie in Mainz by thermal ionisation mass spectrometry (TIMS) following methods outlined by Rampone *et al.* (1995). Both leached and unleached aliquots of whole-rock powders were analysed. Whole-rock powders were leached in hot 6 N HCl, and the residue was thoroughly washed in H₂O. During the period of analyses (1995), the La Jolla Nd standard and SRM-NBS 987 Sr standards yielded $^{143}\text{Nd}/^{144}\text{Nd} = 0.511859 \pm 11$ (five analyses) and $^{87}\text{Sr}/^{86}\text{Sr} = 0.710266 \pm 14$ (10 analyses). Rb, Sr, Sm and Nd concentrations were determined by isotope dilution (ID).

RESULTS

Major- and trace-element geochemistry

The major- and trace-element geochemistry of the Hili Manu peridotite is presented in Tables 1 and 2. Compared with peridotite bodies from alpine, ophiolitic and abyssal environments, the Hili Manu peridotite is exceptional in being relatively free from alteration. Our analysed Hili Manu peridotite samples have an average LOI of $0.04 \pm 0.29\%$. In comparison, the abyssal peridotites reported by Niu (2004) are >60 vol% serpentinised with LOI >10%. The geochemical variations observed in the Hili Manu peridotite can confidently be attributed to primary mantle processes of partial melting, and melt infiltration and impregnation. There is no evidence of mineralogical or chemical modification after the high-temperature mylonitic fabric was produced. Anhydrous MgO contents of the Hili Manu peridotite vary from 37 to 45 wt%, and many major and trace elements display a strong correlation with MgO (Figure 3). Geochemical modelling and experimental studies on a model Hili Manu peridotite composition demonstrate that these correlations are consistent with partial melting processes in the mantle (T. J. Falloon & R. F. Berry unpubl. data).

Many elements do not display good correlations with MgO, especially the low rare-earth elements (LREE) and high field strength elements (HFSE) such as Nb and Ta. Chondrite normalised REE patterns for the Hili Manu samples are presented in Figure 4. Hili Manu lherzolite samples show LREE-depleted patterns ($\text{La}_N/\text{Sm}_N < 1$), whereas the harzburgites show LREE enrichment patterns ($\text{La}_N/\text{Sm}_N > 1$). The REE contents of the Hili Manu peridotite, in common with other peridotite suites (Takazawa *et al.* 2000, 2003; Niu 2004), are inconsistent with a simple partial melting process and indicate that peridotite samples from the Hili Manu peridotite are not simple residues.

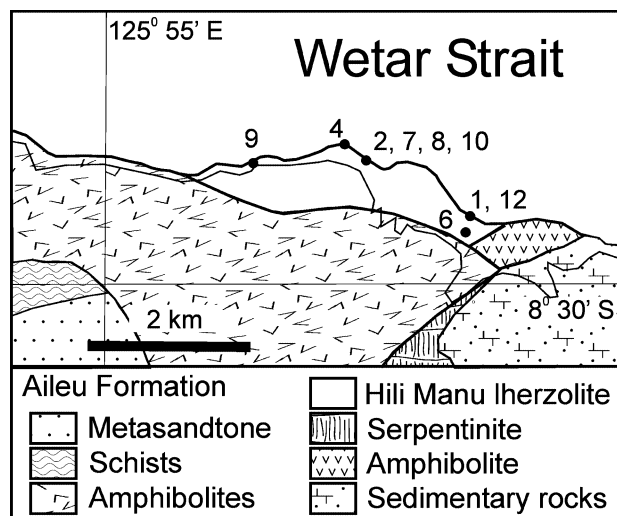


Figure 2 Sample locations of studied peridotite samples from the Hili Manu peridotite. Location of map shown on Figure 1.

Table 1 Whole-rock major- and trace-element geochemistry (XRF) and modal mineral abundances of the Hili Manu peridotite samples investigated in this study.

UTAS sample no.	155964	155968	155956	155976	155960	155971	155957	155970	155961
Sample no. (Berry 1981)	1	2	4	6	7	8	9	10	12
Latitude (°S)	8.492833	8.486	8.484167	8.493833	8.486	8.486	8.486167	8.486	8.492833
Longitude (°E)	125.9573	125.9455	125.943	125.9567	125.9455	125.9455	125.9318	125.9455	125.9573
Lithology	Harzburgite					Lherzolite			
Modal abundances (wt%)									
Olivine	73.6	66.8	64.4	68.6	60.4	60	61.2	59.3	48.3
Orthopyroxene	24.1	29.7	25.1	20.4	27.7	27.3	23.4	26.5	31.1
Clinopyroxene	1.8	2.4	8.8	9.4	10.1	11	13.5	12.3	17.8
Spinel	0.6	1.1	1.7	1.6	1.8	1.7	1.8	1.9	2.8
Major elements (wt%)									
SiO ₂	44.69	45.17	43.79	44.22	45.00	44.91	44.22	45.09	45.48
TiO ₂	0.02	0.02	0.05	0.08	0.11	0.10	0.09	0.10	0.14
Al ₂ O ₃	0.78	0.88	2.14	2.46	3.27	3.17	3.19	3.08	4.27
Cr ₂ O ₃	0.4	0.41	0.42	0.37	0.37	0.37	0.37	0.34	0.38
Fe ₂ O ₃	8.1	8.86	9.02	8.99	8.95	8.94	8.91	9.42	8.61
MgO	44.66	43.15	40.88	41.61	38.65	39.19	38.41	39.13	36.83
NiO	0.32	0.30	0.28	0.29	0.26	0.27	0.26	0.24	0.25
MnO	0.11	0.13	0.13	0.13	0.13	0.13	0.13	0.14	0.14
CaO	0.52	0.74	2.24	2.15	3.08	3.03	3.11	2.76	4.10
Na ₂ O	0.04	0.03	0.11	0.18	0.26	0.22	0.19	0.22	0.33
K ₂ O	0.01	0.00	0.01	0.01	0.01	0.00	0.00	0.01	0.01
P ₂ O ₅	0.01	0.00	0.01	0.01	0.01	0.01	0.01	0.01	0.02
S	0.01	0.00	0.03	0.01	0.02	<i>nd</i>	0.02	0.02	0.01
LOI	-0.26	0.07	0.69	-0.44	-0.06	-0.24	1.18	-0.21	-0.37
Trace elements (ppm)									
Cr	2880	3054	3200	2925	2775	2953	2924	2597	3074
Ni	2478	2387	2173	2214	1960	2105	2093	2120	1916
Sc	6	9	14	13	14	15	16	1	19
V	28	35	73	64	71	77	80	10	101

In Figure 5, we plot, in a similar manner to Niu (2004), the correlation coefficients (R^2) of the REE with MgO and compare these with R^2 values from abyssal peridotite (Niu 2004) and the Horoman peridotite body (Takazawa *et al.* 2000). As can be seen from Figure 3, the R^2 values for the MREE to the HREE are all above 0.9 for the Hili Manu peridotite, and together these REE (Lu through to Sm) show a distinct 'plateau'. This result indicates that the variations in HREE and MREE are consistent with partial melting (T. J. Falloon & R. F. Berry unpubl. data), whereas the LREE (Nd through to La) are not. It can be seen from Figure 5 that the R^2 values for the LREE drop dramatically in a systematic manner for Hili Manu peridotite, indicating that another process or processes is required to explain their variation. A similar result is also evident for the Horoman peridotite (Figure 5). The REE patterns for harzburgites from Hili Manu are similar to refractory peridotites from other mantle suites which show U-shaped normalised REE patterns (Prinzhofer & Allègre 1985; Jahn 1986; McDonough & Frey 1989; Sharma & Wasserburg 1996; Van der Wal & Bodinier 1996; Gruau *et al.* 1998; Parkinson & Pearce 1998; Godard *et al.* 2000; Takazawa *et al.* 2000, 2003; Girardeau *et al.* 2002; Niu 2004). The presence of U-shaped REE patterns in depleted residues suggests that they have been affected by an additional mantle process such as metasomatism or secondary fertilisation during the

melt/rock reaction (Navon & Stolper 1987; Bodinier *et al.* 1990; Kelemen *et al.* 1992; Takazawa *et al.* 1992; Godard *et al.* 2000; Niu 2004).

Neodymium and strontium isotopic data

The Nd and Sr isotopic results for the Hili Manu peridotite are presented in Table 3 and Figure 6. The samples have a large range in $^{143}\text{Nd}/^{144}\text{Nd}$ (0.51243–0.51326) and $^{87}\text{Sr}/^{86}\text{Sr}$ (0.70312–0.70942) values. The lherzolite samples (except sample 155956) all have $^{143}\text{Nd}/^{144}\text{Nd}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ values consistent with a depleted mantle source and fall within the range defined by natural glasses from Indian Ocean spreading centres. However, the two studied harzburgite samples have enriched $^{143}\text{Nd}/^{144}\text{Nd}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ values, and fall within the range of volcanics from the Sunda–Banda Island Arc. The Nd and Sr isotopic values, in general, are poorly correlated with other major and trace elements and ratios. In particular, a poor correlation of $^{143}\text{Nd}/^{144}\text{Nd}$ with Sm/Nd makes any calculation of model ages for the Hili Manu peridotite problematic (Rampone & Piccardo 2000). The range in Nd and Sr isotopic values displayed by the Hili Manu peridotite emphasises again that they are not simple residues. The range in Nd and Sr values of the Hili Manu peridotite is most likely explained by the infiltration of migrating melts of distinctively different

Table 2 Whole-rock trace-element geochemistry (ICPMS) of the Hili Manu peridotite samples investigated in this study.

UTAS sample no.	155964	155968	155956	155976	155960	155971	155957	155970	155961
Trace elements (ppm)									
Sc	4.6	9.5	13.4	13.6	16.7	16.6	16.4	16.2	19.6
Rb	0.077	0.074	0.040	0.037	0.184	0.086	0.045	0.073	0.028
Sr	0.309	1.549	0.614	2.123	2.622	1.751	1.195	2.262	2.324
Sn	0.01	0.0304	0.0336	0.0662	0.0483	0.0237	0.0180	0.0394	0.3375
Y	0.3489	0.2822	1.2616	1.7891	2.8025	2.7427	2.5308	2.4408	4.1902
Zr	0.1251	0.2253	0.2547	2.0016	2.0804	2.1254	0.8272	2.2070	2.8760
Nb	0.0300	0.0754	0.0190	0.0508	0.0824	0.0643	0.0246	0.0448	0.0191
Mo	0.0396	0.1460	0.2618	0.1743	0.1295	0.1700	0.1600	0.2034	0.0758
Ba	0.9408	0.5500	0.3000	0.2236	4.4978	0.5345	0.2979	0.4482	0.8003
Sb	0.0914	0.0077	0.0797	0.0049	0.0170	0.0081	0.0144	0.0092	0.0072
La	0.0342	0.0170	0.0088	0.0154	0.0611	0.0444	0.0072	0.0264	0.0195
Ce	0.0535	0.0285	0.0168	0.1029	0.1701	0.1467	0.0296	0.1237	0.1315
Pr	0.0051	0.0037	0.0056	0.0293	0.0370	0.0363	0.0154	0.0332	0.0461
Nd	0.0190	0.0192	0.0512	0.2511	0.3104	0.3339	0.1758	0.2975	0.4521
Sm	0.0050	0.0071	0.0473	0.1192	0.1666	0.1828	0.1338	0.1630	0.2512
Eu	0.0017	0.0027	0.0225	0.0542	0.0718	0.0788	0.0591	0.0714	0.1147
Gd	0.0098	0.0154	0.1108	0.1933	0.2996	0.3046	0.2549	0.2687	0.4467
Tb	0.0030	0.0039	0.0258	0.0411	0.0613	0.0643	0.0561	0.0556	0.0935
Dy	0.0315	0.0367	0.1976	0.2980	0.4579	0.4688	0.4101	0.4123	0.6849
Ho	0.0086	0.0103	0.0461	0.0689	0.1033	0.1083	0.0926	0.0933	0.1525
Er	0.0375	0.0383	0.1469	0.2192	0.3110	0.3332	0.2855	0.2924	0.4647
Tm	0.0066	0.0069	0.0226	0.0331	0.0456	0.0503	0.0426	0.0441	0.0680
Yb	0.0540	0.0589	0.1619	0.2334	0.3191	0.3494	0.3001	0.3123	0.4632
Lu	0.0079	0.0108	0.0261	0.0373	0.0495	0.0554	0.0466	0.0495	0.0717
Hf	0.0038	0.0082	0.0246	0.0967	0.1072	0.1333	0.0693	0.1161	0.1613
Ta	0.0003	0.0022	0.0004	0.0008	0.0018	0.0025	0.0004	0.0009	0.0008
Bi	0.0002	0.0004	0.0103	0.0018	0.0001	0.0004	0.0008	0.0014	0.0004
Th	0.0200	0.0044	0.0016	0.0046	0.0116	0.1040	0.0015	0.0046	0.0010
U	0.0482	0.0033	0.0007	0.0050	0.0028	0.0067	0.0008	0.0016	0.0005

Table 3 Whole-rock isotope geochemistry of the Hili Manu peridotite samples investigated in this study.

UTAS sample no.	Sr ⁸⁷ /Sr ^{86a}	Sr ⁸⁷ /Sr ⁸⁶	Nd ¹⁴³ /Nd ¹⁴⁴	Rb	Sr	Nd	Sm
155976	0.703841	0.70312	0.513258	0.04	2.17	0.244	0.115
155960	0.704252	0.703488	0.513187	0.645	2.37	0.32	0.133
155961	0.703595	0.703507	0.512975	0.106	2.28	0.322	0.215
155970	0.705928	0.704806	0.512902	–	3.58	0.341	0.142
155971	0.706369	0.70515	0.512957	0.077	1.11	0.328	0.16
155957	0.707736	0.705431	0.513119	0.126	1.19	0.164	0.067
155956	0.708469	0.708348	0.512943	0.053	0.71	0.057	0.046
155968	0.708868	0.708766	0.51247	–	2.54	0.018	0.008
155964	–	0.709421	0.512434	0.103	1.6	0.018	0.006

Rb, Sr, Nd and Sm contents are in ppm determined by ID.

^aUnleached powders.

isotopic composition which have reacted with the depleted peridotite. Many studies of peridotite bodies have demonstrated that the isotopic enrichments observed in some peridotites are directly related to proximity of sampling to melt channels. The former existence of melt channels is defined by either: (i) dunite bodies, with transitions to harzburgite and eventually lherzolite at greater distances (Quick 1981; Kelemen *et al.* 1992); or (ii) pyroxenite layers (Becker 1996). In the case of the Hili Manu peridotite, the harzburgites (which are the result of modally enriched bands of orthopyroxene) probably formed by melt

channelling and reaction by a subduction-related siliceous melt composition (T. J. Falloon & R. F. Berry unpubl. data).

DISCUSSION

Comparison with other peridotite suites

Harris and Long (2000), based on major elements, compared the geochemistry of Timor peridotites (Figure 1) with other peridotite suites including alpine peridotites,

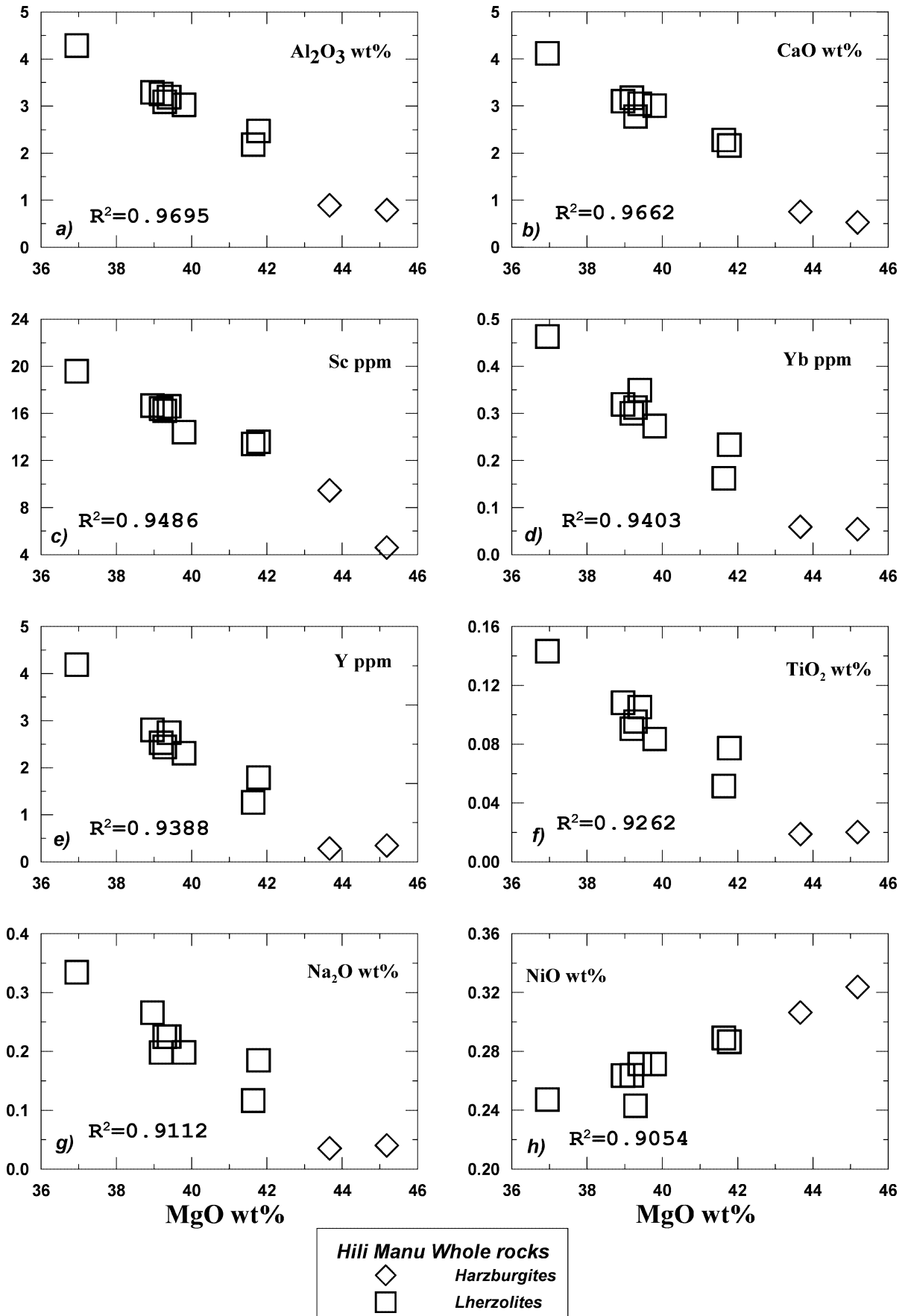


Figure 3 Selected major and trace elements plotted against MgO wt% for samples from the Hili Manu peridotite. R^2 values were determined by least-squares linear regression.

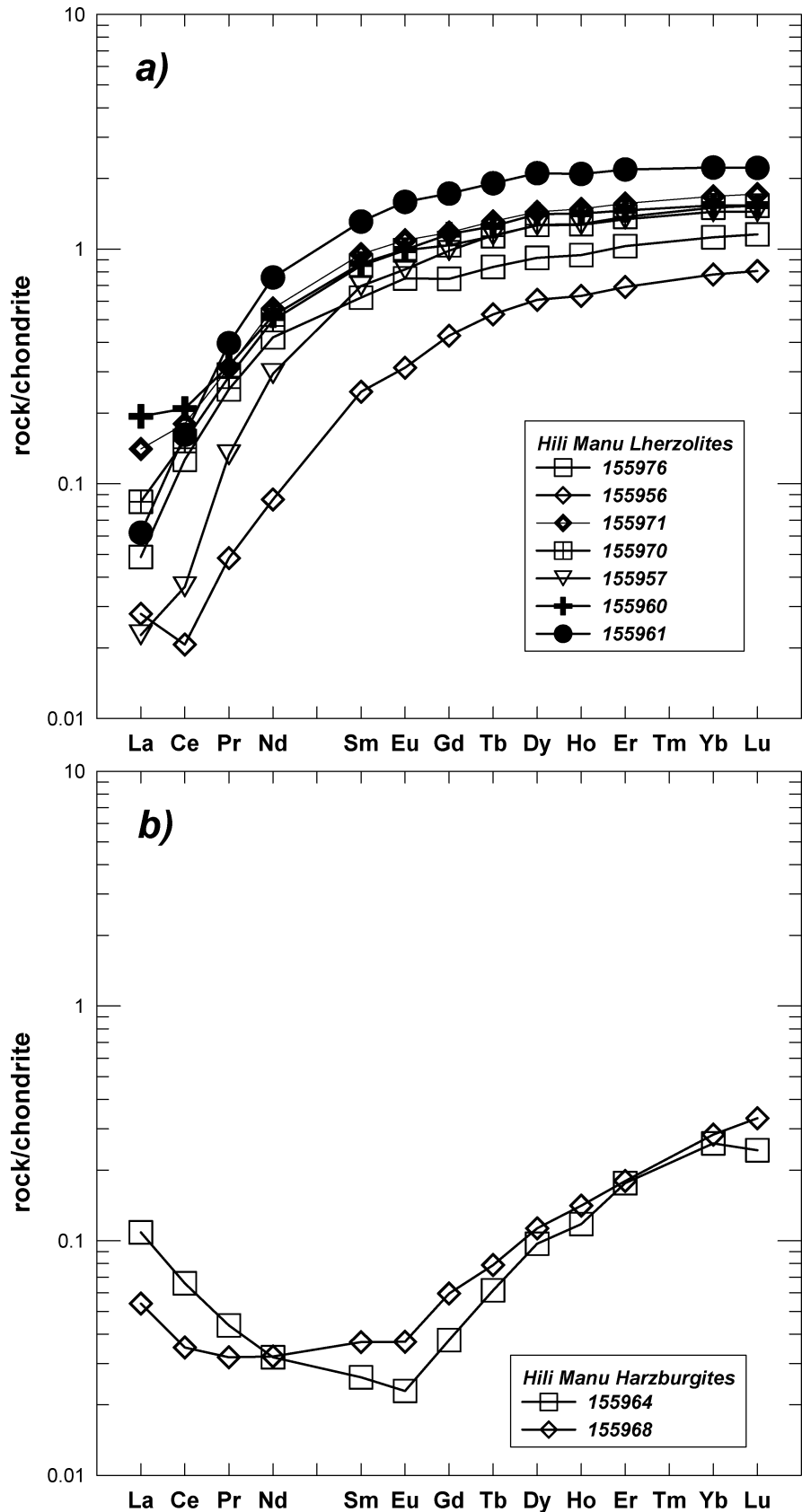


Figure 4 Chondrite-normalised REE patterns for (a) lherzolites and (b) harzburgites for the Hili Manu peridotite.

abyssal peridotites and supra-subduction-related peridotites from modern forearcs and trenches. On the basis of this comparison, they noted that the Timor peridotites including Hili Manu have compositions consistent

with undepleted to slightly depleted continental lithosphere similar to many alpine peridotite bodies (e.g. Ronda, Horomann). On the basis of this and other arguments, Harris and Long (2000) concluded that the

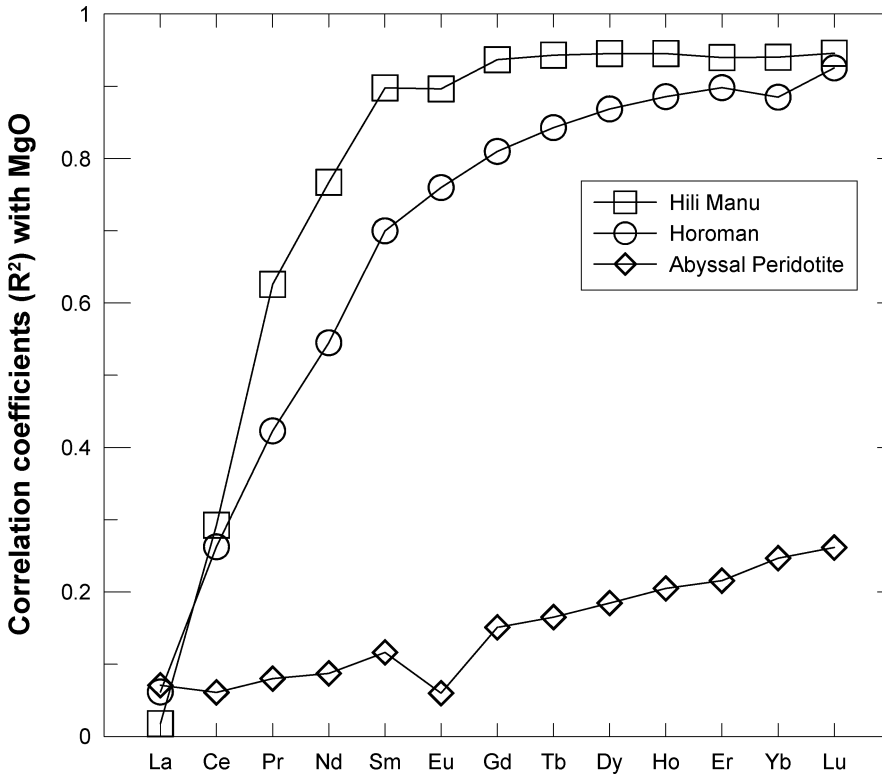


Figure 5 Correlation of REE elements with MgO wt% for the Hili Manu peridotite and a comparison with the same correlations obtained for the Horoman peridotite body (Takazawa *et al.* 2000) and abyssal peridotites (Niu 2004).

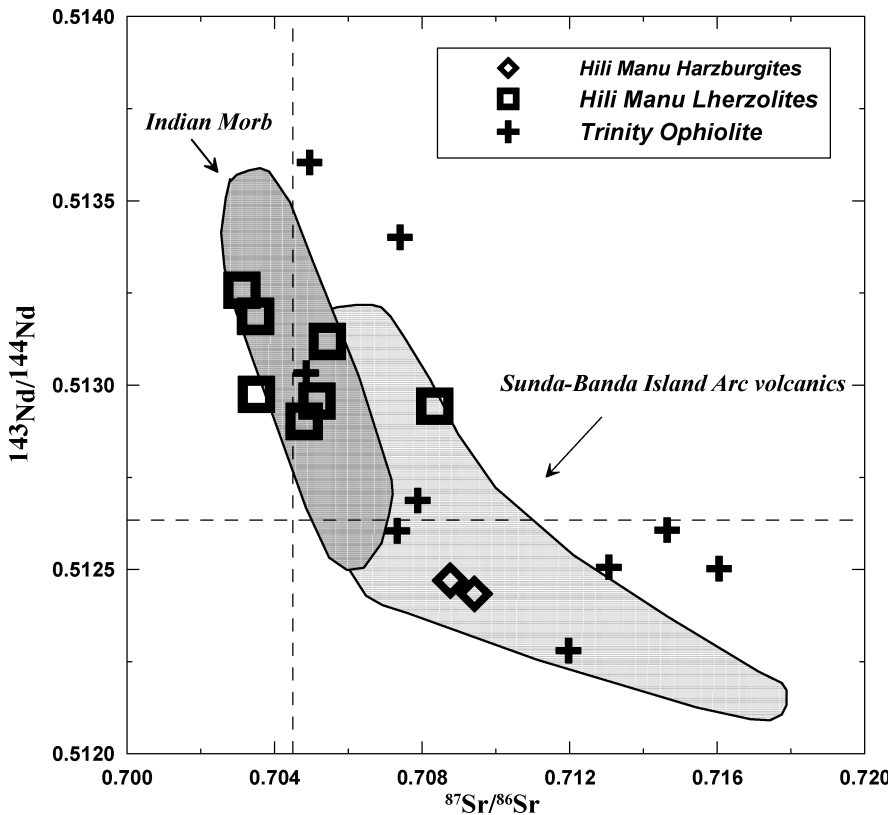


Figure 6 Nd and Sr isotopic composition of Hili Manu peridotite samples compared with glasses from Indian Ocean spreading centres, Sunda-Banda Island Arc volcanics, and peridotites from the Trinity ophiolite (Gruau *et al.* 1998). Data source for Indian Ocean MORB glass data (<http://petdb.ldeo.columbia.edu/petdb/query.asp>); for Sunda-Banda Island Arc volcanics (<http://georoc.mpch-mainz.gwdg.de/georoc/>).

Timor peridotite bodies were derived from Australian subcontinental lithosphere and exhumed during collision. The only exception to this conclusion is the Ocussi peridotite body (Figure 1) which has a very refractory major-element composition similar to

peridotites from supra-subduction settings. The Ocussi peridotite body is also associated with volcanic rocks that have clear island-arc basalt affinities (Harris 1992). Therefore, at least one Timor peridotite body is clearly related to the Banda arc upper plate of the collision

zone. The question is whether any of the other Timor peridotite bodies have a supra-subduction zone geochemical signature.

The Hili Manu peridotite, as discussed by Berry (1981) and Harris and Long (2000), has a range in major-element composition intermediate between fertile, relatively undepleted, continental peridotite bodies and more-refractory abyssal peridotites. The geochemistry of the Hili Manu peridotite suggests that it is not a simple partial melting residue and that it has undergone some other additional processes to produce the U-shaped REE patterns in the most depleted harzburgites, and the range in Nd and Sr isotope compositions. We argued above that similar features are displayed by peridotite

bodies from ophiolites which have a clear supra-subduction setting.

Supra-subduction zone ophiolites form two broad groups in terms of their structure, tectonics and magmatic features (Moores *et al.* 2000; Beccaluva *et al.* 2004). ‘Tethyan’ complexes (e.g. the Oman ophiolite) consist of complete and extensive volcanic, dyke, plutonic and mantle sections that have been obducted as relatively intact slabs onto passive continental margins during collisions (Beccaluva *et al.* 2004). ‘Cordilleran’ complexes (e.g. the Trinity ophiolite) mostly represent dismembered sections of arc volcanic, plutonic and mantle sequences associated with metamorphic rocks and tectonically emplaced, often with no

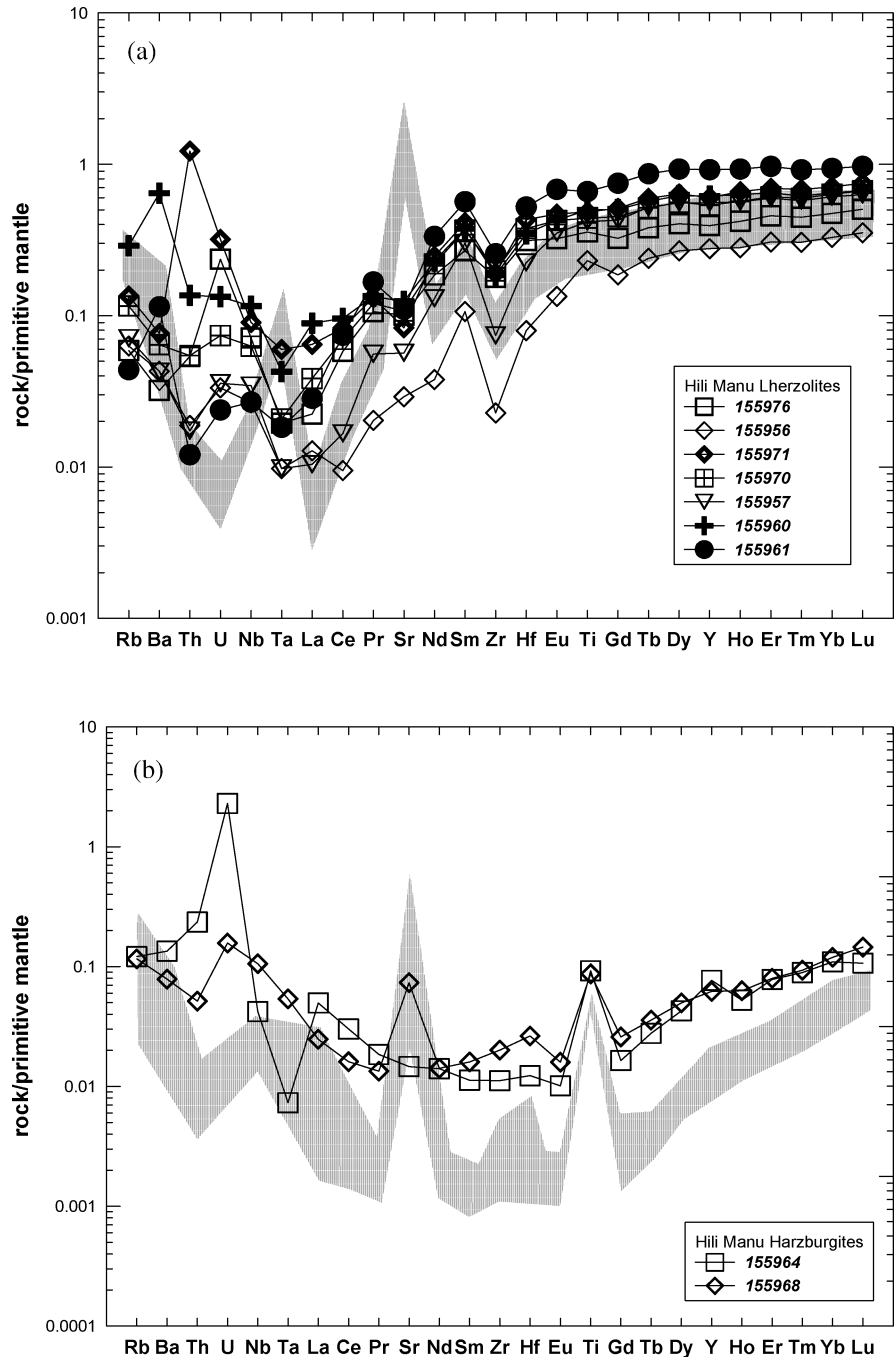


Figure 7 Primitive mantle-normalised abundance patterns for Hili Manu lherzolites (a) and harzburgites (b). Shaded field in (a) is the range in values for Type II lherzolites from Oman (Takazawa *et al.* 2003), and in (b) the range in values for harzburgites from the Western Zone of the Wuqbah peridotite, Oman (Girardeau *et al.* 2002). Normalising values for primitive mantle are from Sun and McDonough (1989).

clear relationships, onto or against a collisional continental margin (Beccaluva *et al.* 2004). The Oman ophiolite has a clear subduction zone influence, reflected in the chemistry of its associated lavas (Pearce *et al.* 1981), but the scale of the ophiolite is such that it records many features characteristic of major mid-ocean ridge spreading centres (MacLeod & Rothery 1992). In contrast, the dismembered Trinity ophiolite is a polygenetic supra-subduction zone ophiolite that formed in a trench-proximal extensional environment (Brouxel & Lapierre 1988; Metcalf *et al.* 2000).

The peridotite from the Trinity ophiolite (Figure 6) shows a range in Nd and Sr isotopic compositions even more extreme than the Hili Manu peridotite (Gruau *et al.* 1998). In the case of the Oman ophiolite, Sharma and Wasserburg (1996) reported a very enriched $^{143}\text{Nd}/^{144}\text{Nd}$ value of 0.511997 for a harzburgite with a U-shaped normalised REE pattern. As can be seen from Figure 7, the trace-element-normalised abundance patterns of lherzolite and harzburgite samples from the Hili Manu peridotite are very close to similar lithologies from the Oman ophiolite (Godard *et al.* 2000; Girardeau *et al.* 2002; Takazawa *et al.* 2003). These comparisons are interpreted here as evidence that the Hili Manu peridotite was derived from a supra-subduction zone setting rather than a continental setting.

Gruau *et al.* (1998) reported a detailed study of peridotites from the Trinity ophiolite. They argued on the basis of oxygen, carbon and hydrogen stable-isotope values in serpentinised peridotites that the enrichment in LREE and isotopic values of the Trinity ophiolite was more likely to be related to a late-stage crustal-derived fluid that had pervaded the peridotite protolith during emplacement onto the North American continental margin. The alternative hypothesis is that the enrichment in LREE and isotope values occurred through a high-temperature interaction of melts/fluids derived

from a subducted crust when the peridotite protolith was located in a supra-subduction setting.

One possible way to distinguish between these two models is to consider the variation of Nb, Ta, Zr and Hf. Experimental data on partitioning of HFSE between solid and fluids indicate that fluids derived from a slab or a continental source are unlikely to carry significant HFSE (Brenan *et al.* 1994; Stolz *et al.* 1996; Audétat & Keppler 2005; Kessel *et al.* 2005; Tropper & Manning 2005). In contrast, experimental studies demonstrate that these ratios can be fractionated by solid/melt equilibria (Foley *et al.* 1999; Green *et al.* 2000; Tiepolo *et al.* 2000). Thus, any changes in Nb/Ta or Zr/Hf ratios must be attributed to addition or removal from a melt. In Figure 8, the Nb/Ta and Zr/Hf values of the Hili Manu samples are plotted with whole-rock peridotite values from alpine (Balumucci, Ronda, Horoman) and abyssal peridotites, and supra-subduction peridotites (Oman ophiolite, Tongan Forearc–Trench).

Experimental studies combined with studies of terrestrial rocks (Godard *et al.* 2000; Rudnick *et al.* 2000; Takazawa *et al.* 2000; Weyer *et al.* 2002, 2003) suggest that the bulk distribution coefficients between solid/melt are in the following order $D_{\text{Nb}} < D_{\text{Ta}}$ and $D_{\text{Zr}} < D_{\text{Hf}}$. Therefore, partial melts will have super-chondritic ratios (chondritic Zr/Hf ~ 36.3 and Nb/Ta ~ 17.57), but residues will be highly depleted in Nb, Ta, Zr and Hf with substantially sub-chondritic Nb/Ta and Zr/Hf ratios. The predicted trends for residues and melts are shown on Figure 8. The process of melting combined with subsequent melt/rock reaction best explains the range in Zr/Hf and Nb/Ta values in both alpine and abyssal peridotites (Weyer *et al.* 2003; Takazawa *et al.* 2003; Niu 2004). Although peridotites from Hili Manu, Oman and Tongan Forearc–Trench samples have all undergone partial melting and melt/rock reactions, their relatively high Nb/Ta ratios

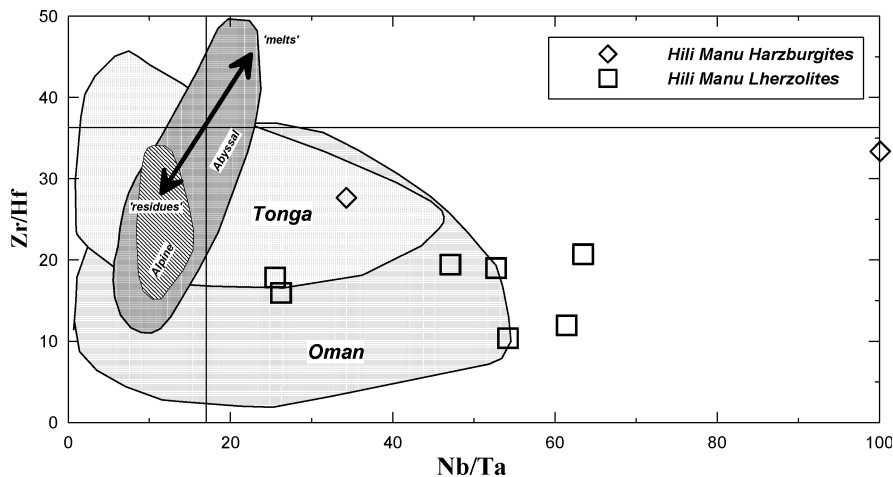


Figure 8 Variation in Nb/Ta and Zr/Hf values in the Hili Manu peridotite compared with values from abyssal peridotites [field encompasses average values for the following fracture zone samples: Udinstev, Vulcan, Islas Orcadas, Prince Edward, Atlantis II, Andrew Basin, Garrett, Bullard, Discovery, Central Indian Ridge (Niu 2004); and Hess Deep (T. J. Falloon unpubl. data)], alpine peridotites [field encompasses the average values for the following bodies: Balmuccia (Weyer *et al.* 2003), Horoman (Takazawa *et al.* 2000) and Ronda (Lenoir *et al.* 2001)], the Oman ophiolite (Godard *et al.* 2000; Takazawa *et al.* 2003) and peridotites from the Tonga Forearc–Trench (T. J. Falloon unpubl. data). Straight lines delineate chondritic values. Arrows indicate the expected trend for residues and melts during mantle-melting processes (see text for further explanation).

suggest that the melt/rock reaction process involved a melt composition distinct from that which was involved in the petrogenesis of alpine and abyssal peridotite protoliths. Stolz *et al.* (1996) suggested that high Nb/Ta ratios are produced by partial melts of subducted oceanic crust/subducted sediment. Therefore, the high Nb/Ta ratios of the Hili Manu peridotite are consistent with a supra-subduction origin for the original peridotite protolith, which was subsequently faulted against the Australian continental margin.

CONCLUSIONS AND IMPLICATIONS FOR TIMOR PERIDOTITES

Most models for the tectonic evolution of the Outer Banda Arc interpret peridotite bodies on Timor, such as Hili Manu peridotite, as fragments of oceanic lithosphere derived from forearc (Hamilton 1979; Sopaheluwakan *et al.* 1989). However, Harris and Long (2000), using major-element geochemistry, considered most of these peridotite bodies to be derived from the Australian subcontinental lithosphere and that no Oman-style ophiolite emplacement took place. Our geochemical study of the Hili Manu peridotite body supports the model that the peridotite bodies associated with the outer Banda Arc represent pieces of oceanic lithosphere from a supra-subduction setting from either a forearc or backarc position. In particular, the Nd and Sr isotopic composition and HFSE ratios such as Nb/Ta indicate that the Hili Manu peridotite was not derived from a continental or mid-ocean ridge environment. This result, along with the clear supra-subduction setting of the Ocuzzi peridotite and associated volcanics in West Timor, gives support to the model that the Miocene collision between the Banda Arc and the Australian continental margin is an analogue for 'Cordilleran'-style ophiolites and their emplacement (Beccaluva *et al.* 2004).

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SUPPLEMENTARY PAPER

Appendix 1 Comparison of literature values for international peridotite PCC-1 and values obtained in this study.