Tracing the Indian Ocean Mantle Domain Through Time: Isotopic Results from Old West Indian, East Tethyan, and South Pacific Seafloor

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The isotopic difference between modern Indian Ocean and Pacific or North Atlantic Ocean ridge mantle (e.g. variably lower 206Pb/ ^{204}Pb for a given ε_{Nd} and $^{208}Pb/^{204}Pb)$ could reflect processes that occurred within a few tens of millions of years preceding the initial breakup of Gondwana. Alternatively, the Indian Ocean isotopic signature could be a much more ancient upper-mantle feature inherited from the asthenosphere of the eastern Tethyan Ocean, which formerly occupied much of the present Indian Ocean region. Age-corrected Nd, Pb, and Sr isotopic data for 46–150 Ma seafloor lavas from sites in the western Indian Ocean and ocean-ridge-type Tethyan ophiolites (Masirah, Yarlung-Zangpo) reveal the presence of both Indian-Ocean-type compositions and essentially Pacific-North Atlantic-type signatures. In comparison, Jurassic South Pacific ridge basalts from Alexander Island, Antarctica, possess normal Pacific-North Atlantic-type isotopic ratios. Despite the very sparse sampling of old seafloor, the age-corrected $\varepsilon_{M}(t)$ values of the old Indian Ocean basalts cover a greater range than seen for the much more thoroughly sampled present-day spreading axes and islands within the Indian Ocean (e.g. 18 ε_{Nd} units for basalts in the 60–80 Ma range vs 15 ε_{Nd} units for 0–10 Ma ones). The implications of these results are that the upper mantle in the Indian Ocean region is becoming increasingly well mixed through time, and

that the Indian Ocean mantle domain may not greatly pre-date the age of earliest spreading in the Indian Ocean.

KEY WORDS: mantle geochemistry; old Indian Ocean; Tethyan crust

INTRODUCTION

Isotopic studies of MORB (mid-ocean ridge basalts) have established the existence of a vast mantle domain in the Indian Ocean distinct from the sources of Pacific and North Atlantic MORB. Along the present Indian Ocean spreading axes, this domain includes the entire Central Indian and Carlsberg ridges and most of the Southeast and Southwest Indian ridges, stretching from about 126°E on the Southeast Indian Ridge (Klein *et al.*, 1988; Pyle *et al.*, 1992) to about 26°E on the Southwest Indian Ridge (Mahoney *et al.*, 1992) and northward into the Red Sea (e.g. Schilling *et al.*, 1992; Volker *et al.*, 1993). Indian MORB are characterized, in particular, by lower

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values of 206 Pb/ 204 Pb relative to ϵ_{Nd} and 208 Pb/ 204 Pb than Pacific and North Atlantic MORB, and also tend to have comparatively high ⁸⁷Sr/⁸⁶Sr (e.g. Hedge et al., 1979; Dupré & Allègre, 1983; Hamelin et al., 1986; Michard et al., 1986; Price et al., 1986; Dosso et al., 1988; Mahoney et al., 1989, 1992; Hall et al., 1995). When isotopic data for samples from the fringes of the Indian Ocean domain are removed from consideration, there is remarkably little overlap of the Indian MORB data set with the isotopic field defined by >95% of published Pacific and North Atlantic MORB data in either the ε_{Nd} vs ²⁰⁶Pb/ ²⁰⁴Pb or ²⁰⁸Pb/²⁰⁴Pb vs ²⁰⁶Pb/²⁰⁴Pb diagrams (Fig. 1). Furthermore, recent studies of Western Pacific back-arc and marginal basin lavas (e.g. Hochstaedter et al., 1990; Loock et al., 1990; Hickey-Vargas, 1991, 1998; Tu et al., 1992; Crawford et al., 1995; Hickey-Vargas et al., 1995; Spadea et al., 1996), and of island-arc lavas in the Philippines (Mukasa et al., 1987; Castillo, 1996), show that isotopically Indian-MORB-like asthenosphere also underlies this region and thus appears to extend far to the east of the Indian Ocean proper. However, despite its great size, the history and origins of this domain (i.e. why it is different from Pacific and North-Atlantic-type mantle) are understood only poorly.

Two general classes of hypotheses have been proposed to account for the Indian Ocean asthenospheric domain. One is that it was created shortly before and during the breakup of Gondwana in the processes that formed the Indian Ocean itself. Possible causes involve either upwelling of deep, isotopically unusual plume-related mantle or widespread introduction of small amounts of continental lithospheric or old, shallowly subducted sedimentary material into the MORB source mantle (promoted by Gondwanan rifting, subduction-erosion, and/or the erosive action of the Kerguelen, Marion, Crozet, and Bouvet starting-plume heads) (e.g. Castillo, 1988; Klein et al., 1988; le Roex et al., 1989; Mahoney et al., 1989, 1992; Storey et al., 1989, 1992; Pyle et al., 1992, 1995; Weis et al., 1992; Hickey-Vargas et al., 1995; Rehkämper & Hofmann, 1997). In either case, this material is postulated to have been dispersed to its present extent along asthenospheric flow paths as the Indian Ocean opened. In this class of hypotheses, older Indian MORB would be expected to show more variable isotopic signatures than modern ones because less time would have been available for intermixing of 'normal' (i.e. Pacific-North Atlantic type) and contaminated asthenosphere. Depending upon their location and age, and the nature of asthenospheric dispersal patterns, some older lavas might not have Indian-MORB-type isotopic signatures at all; the same would probably be true of seafloor formed north of Greater India in the eastern Tethyan Ocean (Mahoney et al., 1992), the Mesozoic ocean that existed in much of the same region now occupied by the Indian Ocean before opening of the latter. An alternative

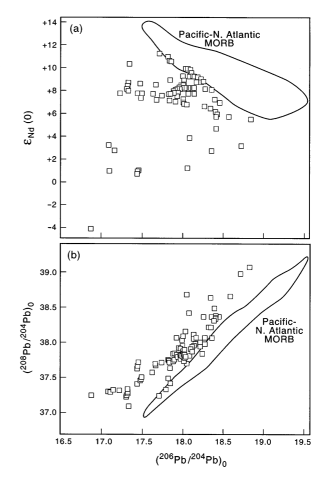


Fig. 1. Plot of ε_{Nd} (a) and ${}^{208}Pb/{}^{204}Pb$ (b) vs ${}^{206}Pb/{}^{204}Pb$ for Indian MORB (\Box) compared with the field defined by >95% of published high-quality data for Pacific and North Atlantic MORB. Data for the edges of the modern Indian Ocean mantle domain, where some transitional compositions occur, are excluded (i.e. in the Australian–Antarctic Discordance, along the western Southwest Indian Ridge, and Red Sea–Gulf of Aden). Principal MORB data sources include those cited in the text, plus Ito *et al.* (1987), White *et al.* (1987), Hanan & Schilling (1989), Dosso *et al.* (1993), Bach *et al.* (1994), Mahoney *et al.* (1994), and other recent studies cited therein.

hypothesis is that the Indian MORB mantle domain is a much longer-lived asthenospheric feature that existed well before the Indian Ocean started opening up (e.g. Hart, 1984; Crawford *et al.*, 1995). In this case, the characteristic Indian-Ocean-type isotopic signature would be expected to be typical of both old Indian MORB and seafloor erupted in a widespread region of the Tethyan Ocean.

Recently, Lanyon (1995), Pyle *et al.* (1995), and Weis & Frey (1996) have studied old seafloor basalts from drill and dredge sites in the eastern Indian Ocean east of about 90°E. Their results indicate that lavas with ages ranging from 15 to 125 Ma exhibit rather typical Indian-MORB-type isotopic signatures; that is, values lying to

the low-²⁰⁶Pb/²⁰⁴Pb side of the Pacific–North Atlantic MORB fields in Fig. 1. However, ~150 Ma samples from Deep Sea Drilling Project Site 261 (off the northwestern corner of Australia) lack clear Indian-MORB-type characteristics, and the same may be true of similar-age MORB from nearby Site 765 (see Weis & Frey, 1996). Although preserved within the northeastern corner of the Indian Ocean, these basalts were formed around 30–35°S at a Tethyan spreading center some 15 my before significant spreading began in the eastern Indian Ocean between Greater Indo-Madagascar and Australia–Antarctica (e.g. Ogg *et al.*, 1992).

In this paper, we present results for 46–140 Ma lavas from drill sites in the western Indian Ocean. Also, although the eastern Tethyan Ocean no longer exists, fragments of Tethyan seafloor are preserved in MORBtype ophiolites along the Tethyan suture belt in southern Asia (Fig. 2). Here, we discuss results for two such suites: 110 Ma basalts from the Yarlung–Zangpo suture of Tibet, and a group of 150 Ma and 120 Ma rocks from the Masirah ophiolite off the Arabian peninsula. As a comparison, we also present isotopic data for Jurassic Pacific MORB from Alexander Island, Antarctica.

METHODS

Several of the samples studied were fairly fresh but most were affected by seawater-mediated alteration, ranging from mild 'brownschist' to zeolite, prehnite-pumpellyite or, in some cases, lower greenschist facies (see Davies et al., 1974; Fisher et al., 1974; Simpson et al., 1974; Whitmarsh et al., 1974; Moseley & Abbotts, 1979; Abbotts, 1981; Girardeau et al., 1985; Pearce & Deng, 1988; Doubleday et al., 1994). Rarely, we were able to pick enough fresh glass or clear-looking plagioclase for isotopic and isotope-dilution analysis; glass and plagioclase separates were cleaned ultrasonically in ultrapure, 1 M HCl and water before dissolution and further processing. For other samples, we followed a preparation procedure closely similar to that used in our previous isotopic studies of old, non-glassy submarine basalts. Chips from the least-altered interior portions of samples were broken to pieces of 3-5 mm in size, which were handpicked under a microscope to avoid visible alteration products (veins, vesicle fillings, and more altered patches of groundmass). The pieces selected were briefly cleaned ultrasonically in ultrapure, weak HF-HNO₃ and H₂O (in sequence) and then broken into smaller (~2 mm) pieces, after which the picking and cleaning procedure was repeated. The pieces chosen were ground in a boron carbide mortar, dissolved, and analyzed for isotopic ratios of Nd, Pb, and Sr and isotope-dilution abundances of Nd, Sm, Pb, Th, U, Sr, and Rb at the University of Hawaii. Several of the samples from Masirah (those lacking MSX or MA

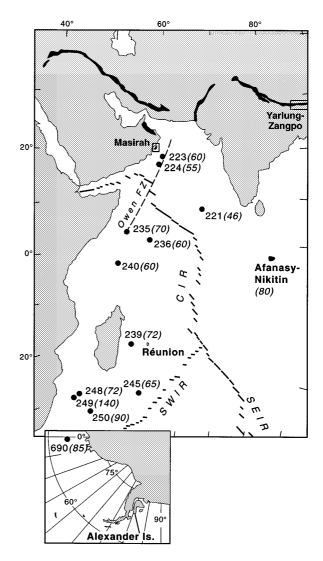


Fig. 2. Map of the western Indian Ocean showing locations of the drill sites (with basement ages or age estimates in parentheses), Masirah, and Yarlung–Zangpo basalts. The belt of Tethyan ophiolites and colored mélanges in southern Asia is depicted in black (after Coleman, 1981). Inset shows locations of Site 690 and Alexander Island. CIR, Central Indian Ridge; SWIR, Southwest Indian Ridge; SEIR, Southeast Indian Ridge. It should be noted that the large difference in age between Site 249 and nearby Site 248 reflects a major boundary in seafloor age provinces (e.g. Simpson *et al.*, 1974).

prefixes in the tables) were analyzed for Nd and Pb isotopes and Nd, Sm, Pb, and U abundances at the University of Bern following a generally similar preparation procedure; Th was analyzed for these samples at the University of Hawaii. In addition, isotopic ratios and parent–daughter element abundances were determined for some samples on splits of powder subjected to a multistep, HCl-dominated acid-leaching procedure effective at removing low-temperature alteration phases (carbonates, clays, chlorite, phosphate, zeolites, ferromanganese oxides; some fresh material is also removed in the process; see Mahoney, 1987; Mahoney & Spencer, 1991). It should be noted that because the picking procedure, as well as acid leaching, when employed, variably modifies a sample's mineralogical composition relative to that of the bulk rock, the isotope-dilution data do not strictly represent bulk-rock elemental abundances and are used here for isotopic age-corrections only. The results are given in Tables 1 and 2.

Very few trace element data have been published for the western Indian Ocean drillhole lavas. Thus, we analyzed a subset of the bulk-rock samples for a broad suite of trace elements; we also analyzed several Masirah samples. Slabs of fresher portions of rock with a minimum of veins and amygdules were chosen (note that for some of the smaller drillhole samples, the freshest looking material had already been reserved for isotopic work). To avoid possible drilling- and/or handling-related contamination, the slabs (typically $5-30 \text{ cm}^3$) were taken from sample interiors, lapped with SiC, cleaned briefly (~5 min) in ultrapure, weak HF–HNO₃ (each ~0.2 M) and water in an ultrasonic bath and powdered in alumina; experience has shown that this procedure does not significantly modify bulk-basalt compositions for the elements analyzed. The resulting bulk-rock powders were prepared and analyzed by inductively coupled plasmamass spectrometry at the University of Hawaii following techniques similar to those described by Jain & Neal (1996). The data appear in Table 3.

A note on effects of alteration on isotopic ratios

Because seawater has a fairly high concentration of Sr (~8 ppm; e.g. Li, 1991) and high ⁸⁷Sr/⁸⁶Sr (today ~0.709) relative to oceanic mantle, ⁸⁷Sr/⁸⁶Sr values in basalts altered by seawater-derived solutions are typically elevated above values in pristine samples; in constrast, Nd and Sm abundances are extremely low in seawater (4 \times 10^{-6} and 8 \times 10^{-7} ppm, respectively) and Nd isotopes are resistant to modification by even rather high amounts of alteration (e.g. McCulloch et al., 1981; Staudigel et al., 1995). Like Nd, Pb abundances in seawater are very low $(2 \times 10^{-6} \text{ ppm})$, so that Pb isotope ratios are affected little by seawater interaction. Pb can be mobile in hydrothermal systems, but redeposition of Pb from one part of a volcanic system to another will normally not be isotopically distinguishable (unless the system possesses significant local-scale isotopic heterogeneity). However, unlike ¹⁴⁷Sm/¹⁴⁴Nd values, ²³⁸U/²⁰⁴Pb ratios (and to a lesser extent, ²³²Th/²⁰⁴Pb) can be affected markedly by mobility of U and uptake of U from seawater (3.2 \times 10⁻³ ppm), in particular, as well as Pb (and sometimes Th) mobility (e.g. Tatsumoto, 1978; Macdougall et al., 1979; Chen & Pallister, 1981). If alteration of an oceanic basalt occurs within several million years after eruption, as typically appears to be the case (e.g. Staudigel et al., 1981), and the rock remains a nearly closed system thereafter, then age-correction of Pb isotope ratios will result in values close to the initial magmatic values even for an old specimen. Indeed, previous studies have demonstrated that good initial-Pb isotope information can be obtained on a range of crystalline magmatic rocks with hydrothermal overprints as high as greenschist facies (e.g. Chen & Pallister, 1981; Göpel et al., 1984). However, if alteration of parent-daughter ratios occurs long after eruption, or repeatedly over many millions of years, then age-adjusted Pb isotopic values will be erroneous. As a hypothetical illustration, let us consider a basalt erupted at 100 Ma whose ²³⁸U/²⁰⁴Pb ratio has been elevated from an original value of, say, 10 to 40 by some very recent seawater-alteration process; using the measured value of 40 to calculate an 'initial' 206 Pb/204 Pb ratio leads to a substantial overcorrection of 0.47.

RESULTS Old Pacific MORB

The basalts of Alexander Island represent faulted slices of Jurassic seafloor formed by spreading between the Pacific and Phoenix plates, and preserved in an accretionary wedge complex (Doubleday *et al.*, 1994). Ages are well determined at 150 Ma in one location (Sullivan Glacier; samples KG 3513-27 and KG 3513-4) and relatively poorly known at a second (Herschel Heights), where a 150 Ma age is assumed; a third area (Lully Foothills) is well dated at Early Jurassic (~200 Ma). Chemically, the basalts include both normal-type and incompatible-element-enriched MORB (N- and E-MORB); ocean-island-type compositions are also present (Doubleday *et al.*, 1994). The samples we analyzed isotopically are altered to zeolite and prehnite–pumpellyite facies.

In an initial $\varepsilon_{Nd}(t)$ vs (⁸⁷Sr/⁸⁶Sr)_t diagram (Fig. 3), most of the Alexander Island data lie to the high-⁸⁷Sr/⁸⁶Sr side of the MORB field: although $\varepsilon_{Nd}(t)$ is between +8.6 and +5.4, within the range of modern N- and E-MORB, the (⁸⁷Sr/⁸⁶Sr)_t of unleached splits varies from 0.70295 to 0.70439. In contrast to many of the drillhole and Masirah samples (see below), acid-leaching of the two Alexander Island samples whose unleached splits had the highest age-adjusted Sr isotope ratios produced only modest decreases in (⁸⁷Sr/⁸⁶Sr)_t, not enough to move their data points into the MORB field in Fig. 3. This result probably reflects the fact that the main repository of Sr, plagioclase (which along with clinopyroxene typically makes up most of the residue for leached tholeiites; e.g. Mahoney, 1987), was largely replaced by secondary feldspar in these rocks

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Table 1: \mathcal{N}

Sample		Age (Ma)	Nd (ppm)	Sm	Sr	Rb	¹⁴⁷ Sm/ ¹⁴⁴ Nd	⁸⁷ Rb/ ⁸⁶ Sr	(1 ⁴³ Nd/ ¹⁴⁴ Nd)	(¹⁴³ Nd/ ¹⁴⁴ Nd) ₀ (⁸⁷ Sr/ ⁹⁶ Sr) ₀	ε _{Nd} (t)	(⁸⁷ Sr/ ⁸⁶ Sr) _t
Old Pacific MORB Alexander I.	æ											
KG 2988-7E	D	200	14-94	4-034	282.0	43.1	0.1632	0-4414	0.512930	0.70421	+ 6-5	0.70295
KG 3513-4	D	150	10-20	3.138	151-9	5.07	0.1859	0.0965	0.512975	0.70431	+6.7	0.70410
KG 3513-27	_	150	2.463	1.003	184-4	16.0	0.2463	0.2504	0.513091	0.70472	+7.8	0.70419
	D		7-674	2.611	156-5	10-8	0.2056	0.2000	0.513052	0.70482	+7.9	0.70439
KG 3981-2E	_	150			84.60	13.8		0.4722		0.70509		0.70408
	D		17-96	5-064	82.24	15.6	0.1704	0.5474	0.512893	0.70528	+5-4	0.70411
KG 3999-2	р	150	8-045	3-077	45-99	11.0	0.2312	0-6944	0-513117	0.70531	+ 8.6	0.70383
Old Western Indian Ocean	ian Ocean											
Drill sites												
249-33-1(31)	D	140	8-502	2.522	108-8	4.27	0.1793	0.1134	0.512825	0.70412	+ 3.9	0.70389
249-33-3(115)	D	140	2.611	0.8938	99.43	0.508	0.2069	0.0148	0.512876	0.70368	+ 4-4	0.70365
250A-25-3(128)	D	06	6-970	2.496	126.8	8.20	0.2165	0.1871	0.513092	0.70316	+8.6	0.70292
250A-26-5(64)	D	06	9.796	2.608	236-4	1.12	0.1609	0.0137	0.512938	0.70301	+6.2	0.70299
690C-24-1(109)	D	85	77-56	12.93	1724	47.2	0.1008	0.0792	0-512557	0.70386	9.0-	0.70376
239-20-1(127)	_	72	2.966	1.489	111.3	4.25	0.3035	0.1105	0.513190	0.70295	+ 9.7	0.70284
	Ъ		8-624	3.224	118-8	6-46	0.2259	0.1574	0.513129	0.70348	+ 9.3	0.70332
248-17-2(63)	_	72	4-026	1.377	187-0	1-02	0-2067	0.0157	0-512799	0.70432	+ 3.0	0.70431
	D		17.76	4.999	196-6	3-08	0.1702	0.0545	0-512768	0.70438	+2.7	0.70432
235-20-5(73)	D	70	4.604	1.747	99.97	1.00	0.2293	0.0290	0-513074	0.70305	+8:2	0.70302
245-19-1(01)	_	65	1-864	1.122	36-25	2.86	0.3639	0.2283	0.513153	0.70309	+8.6	0.70287
	D		6.208	2.534	68-96	7.99	0-2468	0.3349	0.513104	0.70362	+8.6	0.70331
236-33-3(116)	Glass	60	3.117	1.190	55-64	1.25	0.2308	0.0651	0.513030	0.70345	+7-4	0.70339
	D		2.879	1.106	55.16	1.14	0-2322	0.0599	0.513016	0.70352	+7.1	0.70347
236-34-2(123)	_	60	1.718	0.7133	62.81	1.33	0.2510	0.0613	0.513013	0.70350	+ 6.9	0.70345
	D				69-66	1-37		0.0567		0-70404		0.70399
240-7-1(60)	D	60	4.750	1-816	72.80	5-65	0.2310	0.2245	0.513153	0.70350	+ 9.8	0.70331
240-7-1(97)	_	60	2.130	1-040	66-20	4.59	0.2951	0.2003	0-513206	0.70294	+ 10-3	0.70277
	Ъ		3.590	1.427	68-61	1.93	0.2402	0.0815	0.513197	0.70337	+ 10-5	0.70330
223-40-2(48)	_	60	3.189	1.408	81-03	18.1	0.2669	0.6453	0-513049	0.70391	+7-4	0.70336
	D		3.334	1.238	110.1	22.2	0.2241	0.5828	0-513039	0.70446	+7.6	0.70396
224-11-cc	Ъ	55	33-32	6-777	621-4	12.8	0.1229	0.0597	0.512783	0.70362	+3.3	0.70357

MAHONEY *et al.* INDIAN OCEAN MANTLE DOMAIN THROUGH TIME

Sample		Age (Ma)	(mqq) bN	Sm	Sr	Rb	¹⁴⁷ Sm/ ¹⁴⁴ Nd	⁸⁷ Rb/ ⁸⁶ Sr	(¹⁴³ Nd/ ¹⁴⁴ Nd	(¹⁴³ Nd/ ¹⁴⁴ Nd) ₀ (⁸⁷ Sr/ ⁶⁶ Sr) ₀	$\varepsilon_{\rm Nd}(t)$	(⁸⁷ Sr/ ⁸⁶ Sr) _t
Tethyan basalts Masirah	<i>(</i>											
MSX-75	_	150	3.731	1.576	32.39	0.115	0.2552	0.0103	0.513161	0.70305	+ 9.0	0.70302
	D		4.310	1.699	46-47	0.118	0.2382	0.0074	0.513158	0.70455	+ 9.3	0.70453
MSX-134	_	150	7.168	2.308	159.1	2.18	0.1946	0-0397	0.513071	0.70318	+8.4	0.70310
	D		10-25	3-089	202.4	2.47	0.1821	0-0353	0-513054	0.70352	+8.4	0.70345
MSX-158	_	150			108-9	0.193		0.0051		0.70291		0.70289
	D		6.787	2.171	145-5	0.273	0.1933	0.0054	0-512966	0.70334	+ 6-4	0.70333
MSX-171	_	150			39-67	0.111		0.0081		0.70307		0.70305
	D		4-567	1.783	60.71	0.128	0.2362	0.0061	0-513132	0.70467	+8.8	0.70466
MSX-239	L	150			121-0	2.00		0.0479		0.70318		0.70308
	D		7.040	2.453	167.8	2.16	0-2106	0-0372	0-513043	0.70420	+7.6	0.70412
MA-401	L	150	5-554	2.042	158-6	20.7	0-2222	0.3768	0-513033	0.70431	+7.2	0.70351
	Ъ		6-302	2.238	156-5	24-6	0-2146	0.4543	0-513035	0.70451	+ 7.4	0.70354
MSX-71g	Plag	150	0.2057	0.0574	264-5	0.308	0.1687	0.0034	0-513076	0.70336	0.6+	0.70335
P-20	D	150	10-69	3.267			0.1847		0-513023		+7.7	
P-24	С	150	9.170	2.960			0.1947		0-513041		+7.9	
F-23g	С	150	0.3280	0-0734			0.1353		0-513022		+8.6	
F-24g	D	150	0.2584	0.0639			0.1494		0-513133		+10.5	
MSX-29E	_	120	25.39	5.605	330-0	20-3	0.1335	0.1777	0-512897	0.70314	+ 6.0	0.70284
	С		25-36	5-451	318-9	26-6	0.1300	0.2411	0-512921	0.70345	+ 6-5	0.70304
MSX-219E	_	120	15.45	4.364	67-69	0-690	0.1708	0.0295	0-512881	0.70435	+ 5-1	0.70430
	D		49.34	9.797	88-32	0-977	0-1200	0.0320	0-512850	0.70450	+ 5.3	0.70445
MSX-22E	L	120			698-8	105		0.4342		0.70455		0.70381
	Ъ		23.59	4.786	565-5	41-4	0-1226	0.2116	0-512827	0.70459	+ 4.8	0.70423
F-54E	D	120	20.38	2.786			0-0826		0-512700		+2.9	
F-70E	D	120	24-96	5.240			0.1269		0-512845		+ 5-1	
I-669E	С	120	40-24	6.610			0-0993		0-512819		+ 5.0	
J-53E	⊃	120	70-56	13-97			0.1196		0-512732		+3.0	
J-179E	⊃	120	28-66	5.770			0-1217		0-512873		+ 5-7	
Z-114E	∍	120	26-02	5.261			0.1222		0.512814		+ 4-5	

1290

Sample		Age (Ma)	(mqq) bN	Sm	Sr	Rb	¹⁴⁷ Sm/ ¹⁴⁴ Nd	⁸⁷ Rb/ ⁸⁶ Sr	(¹⁴³ Nd/ ¹⁴⁴ Nd	(¹⁴³ Nd/ ¹⁴⁴ Nd)₀ (⁸⁷ Sr/ ⁸⁶ Sr)₀	ε _{Nd} (t)	(⁸⁷ Sr/ ⁸⁶ Sr),
Yarlung–Zan,	Yarlung-Zangpo (Xigaze)	_										
YZS-1	Ч	110			182.2	13.2		0.2088		0.70412		0.70380
	D		4.005	1-515	118-9	5-53	0.2287	0.1344	0.513095	0.70418	+8-4	0.70397
YZS-2	_	110			133-1	3.89		0.0845		0.70419		0.70406
	D		6.246	2.274	105-8	4-14	0.2200	0.1132	0.513075	0.70428	+8.2	0.70410
YZS-3	_	110			153-4	13.7		0.2579		0.70432		0.70391
	⊃		4.224	1.603	130-9	11-2	0.2294	0.2485	0.513098	0-70446	+8.5	0.70407
YZS-6	_	110			140-0	4-11		0-0850		0.70408		0.70395
	D		9.214	3.173	130-3	4-07	0.2082	0.0905	0.513059	0.70410	+8.0	0.70395
YZS-7	_	110			130-9	6-61		0.1461		0.70406		0.70383
	⊃		7.922	2.793	109-5	6-56	0.2131	0.1734	0.513066	0.70421	+8.1	0.70394
YZS-11	_	110			129.1	0-622		0.0139		0.70407		0.70405
			8.160	2-846	116-3	7.66	0.2148	0.1904	0.513075	0.70425	+ 8.2	0.70395
U, unleach U, unleach ccean-islar relative to ±0.00002 (or equal to abundance & Frei (199 calculated samples ar number [e.	ed split, bu nd-type cor a value of last 2 year the extern s are estim of atta fo assuming e from the g. 249-33-3	t chips han mposition. I: ¹⁴³ Nd/ ¹⁴⁴ Nd ¹⁴³ Nd/ ¹⁴⁴ Nd ¹⁴³ Nd/ ¹⁴⁴ Nd s); for La Jc s); for La Jc ated at <0.2 r the Site 2 r the Site 2 r flos Site 2 r (115) is Site	U, unleached split, but chips hand-picked and acid-cleaned; L, stre ocean-island-type composition. Isotopic fractionation corrections relative to a value of ¹⁴³ Nd/ ⁴⁴ Nd = 0.511850 for the La Jolla Nd s \pm 0.00002 (last 2 years); for La Jolla Nd it is \pm 0.000011 (0.2 ϵ unit or equal to the external uncertainties on these standards. Total bi abundances are estimated at <0.2%, on Sr, <0.5%, and on Rb, ~1% & Frei (1997); data for the Site 221 glass are from Mahoney <i>et</i> a calculated assuming ¹⁴⁷ Sm/ ¹⁴⁴ Nd = 0.1967. Ages of Alexander Isi samples are from the relevant Deep Sea Drilling Project or Ocea number [e.g. 249-33-3(115) is Site 249, core 33, section 3, 115 cm]	acid-cleanec nation corri r the La Jo 2000011 (0: standards. 1%, and on I rom Mahon so of Alexar ng Project section 3, 5	t; L, strongly ections are ¹ lla Nd stand. 2 ε units) in 1 Rb, ~1% for 1 Rb, ~1% for 1 lev <i>et al</i> . (19 nder Island, 1 or Ocean Dri 115 cm].	acid-leache ⁴⁸ NdO/ ¹⁴ Nd ard, and ⁸⁷ S Hawaii and -660 pg for Hawaii data 89). $\varepsilon_{Nd}(t)$ an Masirah, an Masirah, an	d powder; Plat 0 = 0.242436 1^{68} Sr = 0.7102 ± 0.000025 in 1 Sr, <15 pg for Sr, <15 pg for 1^{60} Bern data = 0 today corrat d Yarlung-Zan am volume for	g, plagiocla (1 ¹⁴ Nd/ ¹⁴⁴ Nd 24 for NBS Bern. Within Nd in Hawe Nd in Hawe esponds to gpo sample each site.	se separate; (= 0.241572 987 Sr. The t n-run errors of an a <170 f un a <170 f se are from r ss are from r Site number	 gabbro. E ; gabbro. E ; eSt/^mSr = otal range m on the isotopi on Bern. Ul- is 0.512640; e 0.512640; e ferences cite s are indicate 	suffix indic 0.1194. D easured for easured for ic data abo ic data abo ic data abo ic data abo in text; $M_{\rm of}$ ($n = 0$. ad in text; ad at begi	acid-cleaned; L, strongly acid-leached powder; Plag, plagioclase separate; g, gabbro. E suffix indicates E-MORB or onation corrections are ¹⁴⁸ NdO/ ¹⁴⁴ NdO = 0.241572), ⁸⁸ Sr/ ⁸⁸ Sr = 0.1194. Data are reported for the La Jolla Nd standard, and ⁸⁷ Sr/ ⁸⁶ Sr = 0.71028 for NBS 987 Sr. The total range measured for NBS 987 Sr is -0.000011 (0.2 ε units) in Hawaii and ± 0.00025 in Bern. Within-run errors on the isotopic data above are less than 5.5 , and on Rb, ~1% for Hawaii and ± 0.00025 in Bern. Within-run errors on the isotopic data above are less than 5.5 , and on Rb, ~1% for Hawaii data; for Bern data, ¹⁴⁷ Sm/ ¹⁴⁴ Nd uncertainty is -0.3% . Data for F.24 are from Nägler from Mahoney <i>et al.</i> (1989). $\varepsilon_{\rm Ne}$ (t) = 0 today corresponds are from references cited in text; ages of drill-site ling Project or Ocean Drilling Program volume for each site. Site numbers are indicated at beginning of sample sation 3, 115 cm].

1291

undances	
utios and isotope-dilution ab	•
Table 2: Pb isotopic rat	4

		(Ma)												
Old Pacific MORB Alexander I.	ß													
KG 2988-7E	D	200	0.7713	0.218	0.692	18-5	60.6	3.17	19.771	15-625	39.324	19.187	15-596	38.721
KG 3513-4	D	150	0-5738	0-0814	0.136	9.16	15-9	1.68	19-354	15-616	38.788	19.138	15-605	38-670
KG 3513-27	_	150	0-1112	0-0358	0.0866	18-7	46.7	2.42	19-418	15-574	38-794	18-978	15-552	38-446
	С		0.3097	0.0980	0.294	20.5	63.4	3.00	19.389	15-593	38-866	18-907	15-569	38.393
KG 3981-2E	D	150	0-8495	0.397	1.28	30.6	102	3.23	19.735	15.582	39-487	19-015	15-547	38-726
KG 3999-2	D	150	0.1211	0-0703	0.169	37.9	93.9	2-40	19-819	15-589	39-097	18-928	15-545	38-398
Old Western Indian Ocean	ian Ocea	c												
Drill sites														
249-33-1(31)	D	140	1.390	0.330	1.42	15.4	66.7	4.29	18.188	15-553	38-606	17.858	15-537	38.142
249-33(115)	∍	140	1.005	0.310	1.00	19-5	65.3	3.24	18.128	15-554	38-553	17.700	15-533	38-099
250A-25-3(128)	D	06	0.1975	0.0868	0.1202	27.9	39.9	1.38	18.696	15-514	38.169	18-304	15-495	37.991
250A-26-5(64)	D	06	0.7651	0.113	0.756	9.59	66.3	6.69	19.322	15-585	39.195	19.187	15-579	38-899
690C-24-1(109)	D	85	660.6	1.04	1.90	7.03	13.3	1.83	17.491	15.496	37-501	17.398	15.492	37-445
239-20-1(127)	Ļ	72	0.1175	0.0075	0.0060	3.94	3.23	0.79	17.593	15.428	37.348	17-549	15-426	37.336
	D		0.3776	0.0580	0.0921	9-51	15.6	1.59	17.759	15.452	37.468	17.652	15-447	37-412
248-17-2(63)	L	72	0-4975	0.136	0.370	17.6	49.4	2.72	18-596	15.658	39.141	18-400	15.649	38-965
	D		1-997	0.251	0.993	8.06	33.0	3.96	18-577	15-630	38-957	18-487	15.626	38-839
235-20-5(73)	D	70	0.2533	0.0326	0.0890	8-07	22.7	2.73	18-131	15.456	37.876	18-043	15.452	37.797
245-19-1(01)	Ļ	65	0.0560	0.0154	0.0106	17.2	12.2	0.69	18.353	15.467	37.587	18-178	15.459	37 ·548
	D		0.2408	0-0547	0.129	14-4	35.0	2.36	18-586	15-577	38.107	18-440	15-570	37.994
236-33-3(116)	Glass	60	0.3704	0.0496	0.154	8.67	27.8	3.11	19.364	15.627	38-893	19.283	15.623	38-810
	D		0-5434	0-0454	0.140	5.39	17.2	3.08	19.191	15-630	38-854	19.141	15.628	38-803
236-34-2(123)	Ļ	60	0.1480	0.0103	0.0232	4-47	10.3	2.24	18-662	15-591	38.770	18-620	15-589	38-739
	D		0.6857	0.0290	0.114	2.70	10.9	3.92	18-601	15-573	38.689	18-576	15-572	38-656
240-7-1(60)	D	60	0.2210	0.0536	0.117	15.3	34.5	2.18	18-457	15-514	38.191	18-314	15-507	38-080
240-7-1(97)	L	60	0.0440	0-0369	0.0117	53.5	17.6	0.32	19-045	15-543	38-219	18-545	15-519	38.167
	D		0.2294	0.126	0.0468	34.8	13.4	0.37	18.732	15-520	38-214	18-406	15-505	38-174
223-40-2(48)	Ļ	60	0.1862	0-0863	0.0716	29.6	25.4	0.83	18-726	15-551	38.659	18-449	15-538	38-584
	D		0.9523	0.146	0.182	9.74	12.6	1.25	18-625	15-532	38-500	18-534	15-528	38-463
224-11-cc	D	55	1-977	0.782	3·00	25.3	100	3.84	18-675	15-504	38-819	18-458	15.494	38-546
221-19-2(81)	Glass	46	0.3387	0.0493	0.185	9.14	35.6	2 T.C	007 07	7 7 1 V				000

TativalianMasimiMasimi35.73.1118.80715.54438.443MasimiNSX-134L1500.09680.00120.037211.135.73.1118.80715.57940.400NSX-134L1500.09680.06410.43964.53114.6619.51915.61539.72NSX-134L1500.09680.06410.43964.53114.6619.51915.61539.40NSX-134U1500.01200.01210.07230.1310.5574.42015.61538.40NSX-134U1500.01220.1230.37383.74224.926.6119.1641.30NSX-136U1500.01200.012110.573203615.64338.40NSX-136U1500.01200.01211150.57320.6615.64338.40NSX-136U1500.01200.01211150.57320.6615.64338.50NSX-136U1500.01200.00111150.52315.63138.473P.24U1500.01200.02366.171144.1915.63338.473P.24U1500.0120.02366.171144.9015.64338.473P.24U1500.0280.0230.0231144.1915.64338.473P.24U150 </th <th>Sample</th> <th>< .</th> <th>Age I (Ma)</th> <th>Pb (ppm)</th> <th>D</th> <th>Тh</th> <th>²³⁸U/²⁰⁴Pb</th> <th>²³²Th/²⁰⁴Pb Th/U</th> <th>Th/U</th> <th>(²⁰⁶Pb/²⁰⁴Pb)₀</th> <th>(²⁰⁷Pb/²⁰⁴Pb)₀</th> <th>(²⁰⁸Pb/²⁰⁴Pb)₆</th> <th>(²⁰⁶Pb/²⁰⁴Pb)₀ (²⁰⁷Pb/²⁰⁴Pb)₀ (²⁰⁸Pb/²⁰⁴Pb)₀ (²⁰⁶Pb/²⁰⁴Pb)₁ (²⁰⁷Pb/²⁰⁴Pb)₁ (²⁰⁸Pb/²⁰⁴Pb)₁</th> <th>(²⁰⁷Pb/²⁰⁴Pb),</th> <th>(²⁰⁸Pb/²⁰⁴Pb)_t</th>	Sample	< .	Age I (Ma)	Pb (ppm)	D	Тh	²³⁸ U/ ²⁰⁴ Pb	²³² Th/ ²⁰⁴ Pb Th/U	Th/U	(²⁰⁶ Pb/ ²⁰⁴ Pb) ₀	(²⁰⁷ Pb/ ²⁰⁴ Pb) ₀	(²⁰⁸ Pb/ ²⁰⁴ Pb) ₆	(²⁰⁶ Pb/ ²⁰⁴ Pb) ₀ (²⁰⁷ Pb/ ²⁰⁴ Pb) ₀ (²⁰⁸ Pb/ ²⁰⁴ Pb) ₀ (²⁰⁶ Pb/ ²⁰⁴ Pb) ₁ (²⁰⁷ Pb/ ²⁰⁴ Pb) ₁ (²⁰⁸ Pb/ ²⁰⁴ Pb) ₁	(²⁰⁷ Pb/ ²⁰⁴ Pb),	(²⁰⁸ Pb/ ²⁰⁴ Pb) _t
0 150 0.0686 0.0120 0.037 11.1 35.7 3.11 18.807 15.544 1 1 150 0.0969 0.0941 0.439 64.5 311 4.66 19.870 15.549 1 1 150 0.0969 0.0941 0.439 64.5 311 4.66 19.870 15.549 1 1 150 0.0612 0.0753 0.370 83.7 425 4.92 20.608 15.619 1 1 150 0.0120 0.0131 0.0388 12.2 37.3 2.95 18.773 15.541 1 1 150 0.01560 0.0131 0.0388 12.2 37.3 2.956 15.541 1 1 15 0.0130 0.0131 1.15 0.0288 15.541 15.541 1 1 1 1 1 1 1 1 15.51 15.54 1 1 0.2033	Tethyan basalts Masirah														
1 1 150 0.0969 0.0941 0.430 64.5 311 4.66 19.870 15.79 1 1 0 19.60 0.0182 0.121 0.557 44.2 209 4.59 19.519 15.615 8 1 150 0.0612 0.0753 0.370 83.7 425 4.92 20.608 15.619 1 1 150 0.0129 0.0729 0.0381 12.2 37.3 2.95 18.773 15.519 9 1 150 0.0131 0.0381 0.272 0.175 37.3 2.95 18.773 15.519 1 1 150 0.0130 0.0720 0.0153 30.8 6.45 3.00 15.619 15.561 1 1 150 0.718 0.0203 0.0036 0.113 10.523 15.569 15.569 1 1 150 0.718 0.250 0.715 15.46 15.43 15.569 </td <td>MSX-75</td> <td>U L</td> <td></td> <td>0.0686</td> <td>0.0120</td> <td>0-0372</td> <td>11.1</td> <td>35.7</td> <td>3.11</td> <td>18-807</td> <td>15.544</td> <td>38-443</td> <td>18-545</td> <td>15-531</td> <td>38-177</td>	MSX-75	U L		0.0686	0.0120	0-0372	11.1	35.7	3.11	18-807	15.544	38-443	18-545	15-531	38-177
	MSX-134	L 1		0-0969	0-0941	0.439	64-5	311	4.66	19-870	15-579	40.400	18.352	15-505	38-084
1 150 0.0612 0.0753 0.370 8.37 4.25 4.92 20.608 15-645 1 1 150 0.0684 0.0131 0.0388 12.2 37.3 2.95 18.773 15-519 39 1 150 0.0729 0.0429 0.197 38.3 182 4.60 19.18 15-51 1 1 150 0.1560 0.0752 0.153 30.8 64.6 1.91 18.911 15-53 1 1 1 150 0.118 0.0200 0.011 11.5 0.66 0.06 15-53 1 1 150 0.718 0.0201 11.5 0.66 11.6 15-53 1 150 0.718 0.726 0.728 0.728 15-53 1 150 0.718 0.728 0.768 15-53 1 1 1 1 1 1<1		D	-	0.1802	0.121	0-557	44.2	209	4.59	19-519	15.615	39.752	18.479	15-564	38.192
1 1 10 150 0.084 0.0131 0.0381 12.2 37.3 2.95 18.773 15.519 30 1 150 0.0729 0.0429 0.197 38.3 182 4.60 19.268 15.591 1 1 1 0.0729 0.0752 0.156 0.075 0.197 38.3 182 4.60 19.268 15.534 10 150 0.118 0.0200 0.011 11.5 0.668 0.207 17.5 54.2 3.00 18.486 15.534 10 150 0.118 0.0200 0.011 11.5 0.668 17.7 14.14 5.38 18.694 15.539 10 150 0.164 0.013 0.028 0.0075 3.08 4.11 4.19 15.433 15.534 10 150 0.164 0.013 0.013 0.013 15.33	MSX-158	U 1		0.0612	0.0753	0-370	83.7	425	4.92	20.608	15-645	41.382	18.638	15-548	38-214
39 U 150 0.0729 0.197 38.3 182 4.60 19-268 15-591 1 L 150 0.1560 0.0752 0.153 30.8 64.6 1.91 18-911 15-524 10 U 0.2503 0.0688 0.207 17.5 54.2 3.00 18.649 15-53 10 150 0.1118 0.0200 0.0011 11.5 0.66 0.66 19.148 15-53 10 150 0.361 0.119 0.499 9.48 41.1 4.19 15-43 15-53 10 150 0.718 0.0078 0.019 0.499 9.48 41.1 4.19 15-43 15-53 10 150 0.716 0.725 3.08 3.00 0.943 15-33 15-515 11 11 11 11 11 11 11 15-32 15-515 11 11 11 11 11 11	MSX-171	U 1		0.0684	0.0131	0.0388	12.2	37.3	2.95	18.773	15-519	38-400	18.485	15-505	38.122
I L 150 0.1560 0.0752 0.153 30.8 64.6 1.91 18.911 15.24 I U 0 0.2503 0.0688 0.207 17.5 54.2 3.00 18.649 15.539 I U 150 0.1118 0.0200 0.0011 11.5 0.66 0.06 19.148 15.639 U 150 0.361 0.126 0.627 22.2 114 5.38 18.649 15.639 U 150 0.798 0.119 0.2078 0.6172 22.2 114 5.38 18.643 15.631 U 150 0.150 0.195 0.019 0.0396 6.17 13.3 2.08 18.643 15.631 U 150 0.156 0.019 0.0078 6.17 13.3 2.08 18.643 15.631 U 150 0.156 0.019 0.0078 6.17 <td>MSX-239</td> <td>U 1</td> <td></td> <td>0.0729</td> <td>0.0429</td> <td>0.197</td> <td>38.3</td> <td>182</td> <td>4-60</td> <td>19.268</td> <td>15-591</td> <td>39-420</td> <td>18.367</td> <td>15-547</td> <td>38-066</td>	MSX-239	U 1		0.0729	0.0429	0.197	38.3	182	4-60	19.268	15-591	39-420	18.367	15-547	38-066
U 0.2503 0.0688 0.207 17.5 54.2 3.00 18.649 15.539 1 15 0.1118 0.0200 0.0011 11.5 0.66 0.66 19.148 15.539 1 1 150 0.361 0.126 0.627 2.2 114 5.38 18.694 15.631 1 1 150 0.361 0.126 0.627 2.2 114 5.38 18.694 15.631 1 1 150 0.163 0.019 0.0365 3.08 3.10 0.943 15.53 1 150 0.1508 0.019 0.0356 6.17 13.3 2.08 18.488 15.511 1 1 10 1.508 0.872 4.32 38.6 177 4.44 19.576 15.581 1 1 10 1 10 1.43 38.6 177 4.44 19.709 15.581 1 1 1	MA-401	-		0.1560	0.0752	0.153	30.8	64.6	1.91	18.911	15-524	38-501	18.187	15.488	38-019
Ig Plag 150 0:1118 0:0200 0:0011 11-5 0:66 0:66 19:148 15-689 U 150 0:351 0:126 0:627 2:2 114 5:38 16-631 15-631 U 150 0:798 0:119 0:627 2:2 114 5:38 16-532 U 150 0:798 0:199 0:499 9:48 41·1 4·19 18:332 15-515 U 150 0:195 0:019 0:0075 3:08 3:00 0:943 15-515 U 150 1-508 0:717 13·3 2:08 15-515 U 120 0:798 0:872 4:32 38·0 19·10 15-514 U 120 2:299 1·13 3:39 32·1 9/4 19·10 15-514 U 120 2:299 1·13 3:30 3:31 9/4 19·10 15-514 U 120		D	-	0.2503	0.0688	0-207	17.5	54.2	3.00	18-649	15-539	38-356	18.238	15.519	37.952
U 150 0.361 0.126 0.627 22.2 114 5.38 18.694 15.493 U 150 0.798 0.119 0.499 9.48 41.1 4.19 18.694 15.531 U 150 0.798 0.119 0.0078 9.48 41.1 4.19 18.543 15.531 U 150 0.195 0.019 0.0076 3.08 3.00 0.943 18.392 15.511 U 150 0.195 0.019 0.0396 6.17 13.3 2.08 18.488 15.511 U 120 1-508 0.872 4.32 38.6 177 4.44 19.575 15.581 U 120 2.320 2.322 11.8 67.2 354 5.09 19.768 15.561 U 120 2.298 1.118 73.0 413 5.09 19.568 15.561 U 120 2.298 1.118 73.0 413				0.1118	0.0200	0-0011	11.5	0.66	0.06	19.148	15-589	38-565	18-878	15-577	38-560
U 150 0.798 0.119 0.499 9.48 41.1 4.19 18.543 15.531 U 150 0.164 0.0078 0.0075 3.00 0.943 18.543 15.515 U 150 0.164 0.0078 0.0075 3.08 3.00 0.943 18.543 15.515 U 150 0.195 0.019 0.0396 6.17 13.3 2.08 18.488 15.511 U 120 1.517 1.01 4.49 38.6 177 4.44 19.575 15.581 U 120 2.320 1.13 3.39 32.1 99.4 3.00 19.575 15.581 U 120 2.239 1.13 3.39 32.1 99.4 3.00 19.563 15.581 U 120 2.239 1.13 3.30 20.168 15.561 15.561 U 120	P-20	U 1		0.361	0.126	0.627	22.2	114	5.38	18-694	15-493	38-513	18-171	15.467	37.661
	P-24	U 1		0.798	0.119	0.499	9.48	41.1	4.19	18-543	15-531	38-474	18-320	15.520	38.168
	F-23g	U 1		0.164	0.0078	0.0075	3.08	3.00	0.943	18.392	15-515	38-134	18-320	15-511	38.112
E L 120 1-508 0-872 4-32 38-0 195 4-96 19-709 15-592 U 1.717 1.01 4-49 38-6 177 4-44 19-575 15-581 19E U 120 2.320 2.32 11-8 67/2 354 5.09 20-158 15-581 19E U 120 2.320 2.32 11-8 67/2 354 5.09 20-158 15-581 22E U 120 2.299 11-13 3.33 32-1 99-4 3.00 19-508 15-581 10 120 2-299 1-13 3.33 32-1 99-4 3.00 19-508 15-581 10 120 2-168 1-33 2-71 47/2 162 3:33 19-838 15-602 10 120 2-168 1-33 2-71 47/2 162 3:33 19-838 15-602 10 120 2-	F-24g	U 1		0.195	0.019	0.0396	6.17	13.3	2.08	18.488	15-511	38.299	18-343	15-504	38-200
U 1.717 1.01 4.49 38.6 177 4.44 19.575 15-581 19E U 120 2.320 2.32 11.8 67.2 35.4 5.09 20.158 15.629 22E U 120 2.320 1.13 3.39 32.1 99.4 3.00 19.508 15.514 U 120 2.041 3.41 18.7 73.0 413 5.48 19.189 15.514 U 120 3.041 3.41 18.7 73.0 413 5.48 19.189 15.514 U 120 1.121 0.813 2.71 47.2 162 3.33 19.838 15.602 U 120 2.168 1.33 6.69 40.0 208 5.04 19.433 15.61 U 120 2.461 1.19 7.15 31.6 196 6.00 19.427 15.561 U 120 0.829 0.834 4.	MSX-29E	-		1.508	0-872	4.32	38-0	195	4.96	19.709	15-592	39-842	18-995	15-557	38-683
I9E U 120 2.320 2.32 11.8 67.2 354 5.09 20.158 15.629 22E U 120 2.299 1.13 3.39 32.1 99.4 3.00 19.508 15.631 U 120 3.041 3.41 18.7 73.0 413 5.48 19.189 15.61 U 120 3.041 3.41 18.7 73.0 413 5.48 19.189 15.603 U 120 1.121 0.813 2.711 47.2 162 3.33 19.838 15.602 U 120 2.168 1.33 6.69 40.0 208 5.04 19.622 15.599 U 120 2.461 1.19 7.15 31.6 196 6.00 19.427 15.561 U 120 0.829 0.834 4.45 65.3 367 5.45 18.939 15.561 U 120 0.829 0.3		D		1.717	1-01	4-49	38-6	177	4-44	19-575	15-581	39.689	18-850	15-546	38-634
22E U 120 2.299 1.13 3.39 32.1 99.4 3.00 19.608 15.581 U 120 3.041 3.41 18.7 73.0 413 5.48 19.189 15.514 U 120 3.041 3.41 18.7 73.0 413 5.48 19.189 15.514 U 120 1.121 0.813 2.71 47.2 162 3.33 19.638 15.602 U 120 2.168 1.33 6.69 40.0 208 5.04 19.622 15.599 U 120 2.461 1.19 7.15 31.6 196 6.00 19.427 15.561 U 120 0.829 0.834 4.45 65.3 367 5.45 18.939 15.561 U 120 1.238 0.929 3.38 48.5 182 3.64 19.403 15.561	MSX-219E	U 1		2.320	2.32	11-8	67.2	354	5.09	20.158	15.629	40.903	18-895	15-568	38-797
U 120 3.041 3.41 18.7 73.0 413 5.48 19.189 15.514 U 120 1.121 0.813 2.71 47.2 162 3.33 19.838 15.602 U 120 1.121 0.813 2.71 47.2 162 3.33 19.838 15.602 U 120 2.168 1.33 6.69 40.0 208 5.04 19.562 15.599 U 120 2.461 1.19 7.15 31.6 196 6.00 19.427 15.561 U 120 0.829 0.834 4.45 65.3 367 5.45 18.939 15.518 U 120 1.238 0.929 3.38 48.5 182 3.64 19.403 15.561	MSX-222E	U 1		2.299	1.13	3.39	32.1	99.4	3.00	19.508	15-581	39-403	18-905	15-552	38-811
U 120 1-121 0-813 2-71 47-2 162 3-33 19-838 15-602 U 120 2-168 1-33 6-69 40.0 208 5-04 19-562 15-599 U 120 2-461 1-19 7-15 31-6 196 6-00 19-427 15-561 U 120 0-829 0-834 445 65-3 367 5-45 18-939 15-561 U 120 1-238 0-929 3-38 48-5 182 3-64 19-403 15-561	F-54E	U 1		3-041	3.41	18.7	73-0	413	5.48	19.189	15-514	39.662	17.818	15.448	37.200
U 120 2.168 1.33 6.69 40.0 208 5.04 19.662 15.599 U 120 2.461 1.19 7.15 31.6 196 6.00 19.427 15.561 U 120 0.829 0.834 4.45 65.3 367 5.45 18.939 15.518 U 120 1.238 0.929 3.38 48.5 182 3.64 19.403 15.561	F-70E	U 1		1.121	0.813	2.71	47.2	162	3.33	19-838	15-602	39.268	18-951	15-559	38-303
U 120 2.461 1.19 7.15 31.6 196 6.00 19.427 15.561 U 120 0.829 0.834 4.45 65.3 367 5.45 18.939 15.518 U 120 1.238 0.929 3.38 48.5 182 3.64 19.403 15.561	I-669E	U 1		2.168	1.33	6.69	40-0	208	5.04	19-562	15-599	39.742	18-810	15-563	38-503
U 120 0.829 0.834 4.45 65.3 367 5.45 18.939 15.518 . U 120 1.238 0.929 3.38 48.5 182 3.64 19.403 15.561 .	J-53E	U 1		2-461	1.19	7.15	31-6	196	6.00	19-427	15-561	39.506	18-833	15-532	38-340
U 120 1-238 0-929 3-38 48-5 182 3-64 19-403 15-561	J-179E	U 1		0-829	0.834	4-45	65-3	367	5.45	18-939	15-518	39-639	17.712	15.459	37.451
	Z-114E	U 1		1.238	0.929	3.38	48.5	182	3.64	19-403	15-561	39.253	18-492	15-517	38-169
Z131E U 120 1-457 3-10 13-3 142 632 4-30 20-404 15-610 40-409	Z131E	U 1	20	1.457	3.10	13.3	142	632	4.30	20-404	15-610	40.409	17 <i>.</i> 728	15.480	36-645

MAHONEY *et al.* INDIAN OCEAN MANTLE DOMAIN THROUGH TIME

	Tabi	Table 2: continued	ntinued											
Sample location		Age (Ma)	U (mqq) dA	Þ	۲	²³⁸ U/ ²⁰⁴ Pb	²³² Th/ ²⁰⁴ Pb Th/U	Th/U	(²⁰⁶ Pb/ ²⁰⁴ Pb)	(²⁰⁶ Pb/ ²⁰⁴ Pb)。(²⁰⁷ Pb/ ²⁰⁴ Pb) ₀ (²⁰⁸ Pb/ ²⁰⁴ Pb) ₀ (²⁰⁶ Pb/ ²⁰⁴ Pb) ₁ (²⁰⁷ Pb/ ²⁰⁴ Pb) ₁ (²⁰⁸ Pb/ ²⁰⁴ Pb)	(²⁰⁸ Pb/ ²⁰⁴ Pb) ₀	(²⁰⁶ Pb/ ²⁰⁴ Pb) _t	(²⁰⁷ Pb/ ²⁰⁴ Pb) _t	(²⁰⁸ Pb/ ²⁰⁴ Pb) _t
Yarlung–Zangpo (Xigaze)	o (Xigaze	(é												
YZS-1	∍	110	0.3837	0.0853	0-0375	13-8	6.25	0.44	17.771	15.427	37-417	17.534	15.416	37.383
YZS-2	∍	110	0.6177	0.108	0-0763	10.7	7.88	0.71	17.704	15-411	37.351	17-519	15.402	37.308
YZS-3	⊃	110	0.4667	0.0654	0.0506	8-67	6.93	0.77	17-697	15.426	37.421	17.548	15.419	37.383
9-SZY	⊃	110	0.2712	0.0644	0.102	14.7	23.9	1-58	17.726	15.413	37.432	17.473	15-401	37.301
YZS 7	⊃	110	0.2450	0.0807	0.0827	20.4	21.6	1-02	17.765	15.409	37.388	17-415	15.392	37.270
YZS-11	⊃	110	0.2994	0-0546	0-0894	11-2	19-0	1.64	17-630	15-404	37.369	17.436	15.395	37.265
Pb isotope ratios are corrected for fractionation using the NBS 981 standard values of Todt <i>et al.</i> (1996); the total ranges measured for NBS 981 are \pm 0.011 for ²⁰⁶ Pb/ ²⁰⁴ Pb, \pm 0.010 for ²⁰⁷ Pb/ ²⁰⁴ Pb, and \pm 0.032 for ²⁰⁶ Pb/ ²⁰⁴ Pb (last 2 years). Within-run uncertainties (2 SE) on the isotopic data above are less than these values. Estimated uncertainty on Pb abundances ~0.5%, on U ~1%, and on Th <2%. Propagation of errors on age-corrected Pb isotope ratios: for a 150 Ma sample with ²³⁸ U/ ²⁰⁴ Pb = 30 and ²³² Th/ ²⁰⁴ Pb = 100, a maximum error on parent-daughter ratios translates to an additional error of \pm 0.011 on (²⁰⁶ Pb/ ²⁰⁴ Pb), \pm 0.005 on (²⁰⁷ Pb), and \pm 0.019 on (²⁰⁹ Pb/ ²⁰⁴ Pb) = 0.011 on (²⁰⁶ Pb), total procedural blanks: Pb 4-34 pg in Hawaii and <70 pg in Bern; U <5 pg in Hawaii and <30 pg in Bern; Th <3 pg.	tios are 0.010 fc certaint 30 and ² 0.019 o	t correct or ²⁰⁷ Pb/ ⁴ y on Pb ³² Th/ ²⁰⁴ P	ted for fract ²⁰⁴ Pb, and <u>⊥</u> abundance 'b = 100, a / ²⁰⁴ Pb) _t . Tota	tionation 1 ± 0.032 for ss ~0.5%, maximur al procedu	using the - ∞Bb/∞4F on U ~1% n error or ıral blankı	NBS 981 s bb (last 2 yt 6, and on T n parent-dé s: Pb 4-34	tandard ve sars). With h <2%. Prc tughter rat pg in Hawe	alues of T in-run un pagation ios transl aii and <7	odt <i>et al.</i> (15 certainties (2 of errors or lates to an a '0 pg in Berr	tion using the NBS 981 standard values of Todt <i>et al.</i> (1996); the total ranges measured for NBS 981 are \pm 0.011 for 52 for ²⁰⁸ Pb/ ²⁰⁴ Pb (last 2 years). Within-run uncertainties (2 SE) on the isotopic data above are less than these values. 5%, on U ~1%, and on Th <2%. Propagation of errors on age-corrected Pb isotope ratios: for a 150 Ma sample with timum error on parent-daughter ratios translates to an additional error of \pm 0.011 on (²⁰⁸ Pb/ ²⁰⁴ Pb), \pm 0.005 on (²⁰⁷ Pb/ ²⁰⁴ Pb) to be added to be above at a sample with timum error on parent-daughter ratios translates to an additional error of \pm 0.011 on (²⁰⁸ Pb/ ²⁰⁴ Pb), \pm 0.005 on (²⁰⁷ Pb/ ²⁰⁴ Pb) to be added to be added to be above at a sample with the error on parent-daughter ratios translates to an additional error of \pm 0.011 on (²⁰⁸ Pb/ ²⁰⁴ Pb), \pm 0.005 on (²⁰⁷ Pb/ ²⁰⁴ Pb) to be added	I ranges me isotopic data ed Pb isotop or of ±0.011 Hawaii and	asured for N a above are to ratios: for on (²⁰⁶ Pb/ ²⁰ <30 pg in B	JBS 981 are less than th less than th a 150 Ma si 4b),, ±0.000 ern; Th <3 p	± 0.011 for se values. imple with on (²⁰⁷ Pb/ J.

during alteration (Doubleday *et al.*, 1994) and that little material with pristine ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ remains.

Unlike Sr isotopes, the age-corrected Nd and Pb isotopic data all fall in a restricted field within the Pacific-North Atlantic MORB mantle array in Fig. 4c and d, particularly when the modern array is adjusted to the approximate position it would have occupied at 150 Ma (assuming the only changes have been caused by radioactive decay of parent nuclides in the MORB source mantle). The data plot in the low- $\varepsilon_{\rm Nd}$, high-²⁰⁶Pb/²⁰⁴Pb, high-208Pb/204Pb region of this array. Values of (207Pb/ ²⁰⁴Pb)_t lie within the range for MORB and oceanic islands as well (Fig. 5a). The one sample for which we determined Nd and Pb isotopes on both acid-leached and unleached splits (KG 3513-27) has identical $\varepsilon_{Nd}(t)$, within errors, for each split, at +7.8 and +7.9; leaching also produced only small changes in the age-adjusted Pb isotope ratios of this sample [e.g. $({}^{206}\text{Pb}/{}^{204}\text{Pb})_t$ of 18.98 and 18.91; see linked light and dark squares in Fig. 4]. Because the agecorrected Pb isotopic values of all the samples (1) plot in a small field within the narrow Pacific-North Atlantic MORB array in both panels c and d of Fig. 4 and (2) have a lesser total spread in ²⁰⁸Pb/²⁰⁴Pb and ²⁰⁶Pb/²⁰⁴Pb relative to the measured, present-day range (0.33 vs 0.70and 0.28 vs 0.47, respectively; the range in ²⁰⁷Pb/²⁰⁴Pb is the same within analytical error), and because (3) values for the leached and unleached splits of KG 3513-27 agree well with each other, it appears that alteration affecting U/Pb or Th/Pb ratios mainly occurred within a few million years after eruption. The same appears to be true of most samples from the other localities studied (see below).

Western Indian Ocean drill sites

The drillhole samples display a wide range in elemental composition. Primitive-mantle-normalized incompatible element patterns of several samples are illustrated in Fig. 6, and can be seen to vary from typical N-MORB type (sloping generally downward to the left; e.g. Site 235-20-5) to E-MORB type (moderate upward slope to the left; e.g. Site 250A-26-5). The basalt from Site 690C (on Maud Rise) has an ocean-island-like pattern resembling those of Gough Island lavas (South Atlantic), including a spike at Ba and trough at Th and U; this rock is an unusual xenocryst-bearing alkalic basalt that Schandl et al. (1990) concluded originated from a hydrous source containing small amounts of phlogopite and apatite. The Site 224 sample shows somewhat similar, but less extreme, ocean-island-like features. In contrast, the pattern for a Site 249 lava has a sizeable trough at Nb and Ta (as well as a larger than usual peak at Pb), a characteristic commonly seen in lavas influenced by continental lithosphere, and in arc-related basalts, but

Sample Rb Ba	ЧT			Та	[a	e C	à	ů		T N	, ,	2 0	L	č	ſ	;			I	2	-
		,	qN		ì	3	-	ō		DN	- -	=	E	5	à	~	РЧ	Ъ	E	٩	Lu
Old Western Indian Ocean																					
249-33-3(115) 7-8 80	1-35	5 0.24	3.5	0.19	7-36	15.3	2.11	136	1.12	9-80	85	2.90	0.99	4.13	4-44	28-0	0.92	2.70	0.37	2.83	0.37
250A-26-5(64) 13 193	3 1.71	1 0-47	24	1.35	14-4	31.7	3.62	I	1.16	15.7	123	4.19	1.43	4.94	4.53	25-5	0.86	2.76	0.33	2.68	0.32
239-20-1(127) 6.3 9	9.7 0.10	0 0.04	1.1	0.08	2.33	7.60	1-40	I	0.39	8-03	87	3.13	1.16	4-81	5.56	33.8	1-07	3.64	0.46	3.71	0.50
248-17-2(65) 3.0 68	3 0-86	6 0-20	8.6	0.45	8-47	23.3	3.15	207	1.25	16-6	123	4.66	1.63	5-09	5.36	28.7	1-02	2.83	0.37	2.55	0.35
235-20-5(73) 2.0 4	4-5 0-04	4 0.03	:	0.06	2.04	6.39	1.12	135	0:30	6.38	68	2.27	0.95	3.25	4.20	25-5	0.82	2.53	0.34	2.60	0.35
245-19-1(01) 6-8 5	5.1 0.04	4 0.09	0-87		1.46	5.18	1.00	86	0.23	6.43	57	2.57	0-97	3.79	5.11	32.0	1.07	3.28	0.47	3.23	0-44
236-33-3(117) 5-5 7	7.8 0.14	4 0.05	1.9	0.13	1.84	4.69	0.70	103	0.20	3.96	39	1.26	0.60	2.07	2.81	16.1	0.58	1.85	0.25	1.75	0.24
240-7-1(97) 18 16	§ 0-06	6 0-04	1.5	0.09	1.96	5.41	0.91	89	0.37	5.48	54	1.88	0-85	2.91	3.74	24.3	0.78	2.57	0.33	2.58	0.35
223-40-2(48) 17 82	2 0-27	7 0.11	4.4	0.25	4-96	12.6	1.75	269	0.71	9.52	78	2.58	1.05	3.63	4.26	24.9	0.86	2.44	0.36	2.29	0.33
224-11-cc 28 692	2 4.28	8 1.12	47	3.31	52.6	98.1	11.1	750	2.66	44-8	268	8.69	3.04	9.58	6.36	32.7	1.10	3.03	0.37	2.50	0.36
Masirah																					
MSX-75 0.15 4	4.1 0.07	7 0.03	0-91	90·00	1.21	4.00	0.78	65	0.18	4-64	47	1.95	0.65	2-80	4.06	21.3	0.76	2.49	0.35	2.55	0.37
MSX-134 2.3 50	0.58	8 0.18	8-4	0-55	6-56	14.9	2.10	200	0.36	10-6	78	3.41	1.15	3.88	4.82	24.4	0.91	2.75	0.37	2.41	0.32
MSX-158 0.35 23	3 0.38	8 0.11	5.2	0.34	4-21	9.61	1.42	175	0.21	6-97	50	2.23	0.85	2.73	3.42	18.1	0.69	1.92	0.26	1.89	0.25
MSX-239 2.0 46	3 0.17	7 0.06	3.3 3	0.22	3-22	8-53	1.41	Ι	0.22	7.08	60	2.55	0.98	3-50	3.94	23-5	0-86	2.53	0.36	2.28	0.33
MA-401 22 280	0.25	5 0.09	2.7	0.19	2.63	6-99	1.15	200	0.43	6-64	57	2.31	0-84	3.14	3.80	21.2	0.77	2.32	0.36	2.15	0.32
MSX-29E 30 404	1 4.30	0 0.89	48	2.87	31-3	57.4	6-55	299	2.07	27.2	180	6.04	1.91	6.25	5.89	32.4	1.12	3.17	0.43	3.01	0.40
MSX-219E 1.7 34	1 8-24	4 1-93	75	4.38	52.8	96-5	11.1	70	2.32	41.0	300	8.68	1.59	8.34	8.43	45.8	1.62	5.14	0.74	5.10	0.70
Standard																					
BHVO-1 meas. 9-5 136	3 1-09	9 0-41	18-6	1.11	15-7	40.3	5.21	416	2.13	24.7	181	6.17	2.14	6-51	5.24	25-8	0.85	2.48	0.30	2.05	0.28
BHVO-1 rec. 9.7 133	3 1.233	33 0-41	19-0	1.16	15-8	37.8	5.40	420	2.051	24.8	179	6.10	1.98	6.40	5.20	27.1	66-0	2.40	0.33	2.02	0.29

Table 3: Bulk-rock incompatible element abundances (ppm)

1295

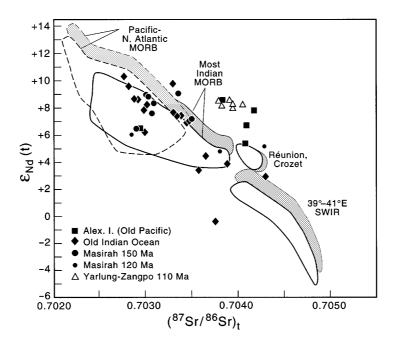


Fig. 3. Age-corrected $\epsilon_{Nd}(t)$ vs (⁸⁷Sr/⁸⁶Sr)₁. It should be noted that not all samples were acid-leached and that for samples with data on both unleached and acid-leached or glass splits, only values for leached or glass splits are shown. Present-day fields for Pacific–North Atlantic MORB, Indian MORB [including the 39°–41°E section of the Southwest Indian Ridge (SWIR)], and Réunion and Crozet hotspot volcanoes are shaded; the adjacent unshaded fields are positioned for 150 Ma, assuming the modern MORB and Réunion–Crozet mantle sources have average ¹⁴⁷Sm/¹⁴⁴Nd of 0·24 and 0·17, and ⁸⁷Rb/⁸⁶Sr 0·02 and 0·10, respectively (see Peng & Mahoney, 1995). Data sources for the fields are as in Fig. 1 and, for Réunion and Crozet, W. M. White (unpublished data, 1993) and Mahoney *et al.* (1996, and references therein).

rarely in either fresh or altered oceanic basalts (compare Site 248 pattern). Patterns of the visibly fresher N- and E-MORB samples (e.g. Site 236-33-3, Site 250A-26-5) are relatively smooth, and even the more altered samples lack the pronounced spikes or troughs seen for some elements in patterns of highly altered basalts (Bienvenu *et al.*, 1990; Staudigel *et al.*, 1995; Jochum & Verma, 1996). Alteration effects are most evident in a marked elevation of Rb in the N-MORB lavas; Ba is also elevated significantly in many of these lavas (e.g. the Site 223 sample). Small peaks or troughs, which may reflect alteration, are present at Pb in several patterns, whereas U peaks and/or low Th/U ratios indicate significant U uptake in several samples (e.g. Site 245-19-1).

We determined Sr and Pb isotopes on both unleached and leached (or glass) splits of seven of the drill-core basalts studied and Nd isotopes on six pairs. The (⁸⁷Sr/ ⁸⁶Sr)_t values of all but one of the leached splits are significantly lower (by as much as 0.0006) than those of the unleached splits, whereas only negligible differences are observed in $\varepsilon_{Nd}(t)$ (0–0.4 epsilon units). The differences in age-corrected Pb isotope ratios for the members of each unleached–leached (or glass) pair are also relatively small, with one exception (see linked symbols in Fig. 4a and b). For example, the difference in (²⁰⁶Pb/²⁰⁴Pb)_t ranges from 0.04 to 0.14, except for sample 245-19-1, which shows a difference of 0.26, by far the largest observed for any of the samples in our study. As with Nd isotopes, the age-corrected Pb isotope ratios of an unleached split can be either slightly higher or lower than for the corresponding leached residue or glass separate. Values of (²⁰⁷Pb/²⁰⁴Pb), are all within the range for MORB and ocean islands (Fig. 5a), except for sample 248-17-2, for which both leached and unleached splits have high (207Pb/ ²⁰⁴Pb), (15.65, 15.63) relative to (²⁰⁶Pb/²⁰⁴Pb), (18.40, 18·49). Abundances of Nd, Sm, Pb, U, Th, and Rb typically dropped substantially with leaching. Although the ¹⁴⁷Sm/¹⁴⁴Nd ratios of leached splits tend to be higher than for their unleached counterparts, consistent with a relative enrichment of clinopyroxene in most of these residues (e.g. Mahoney, 1987), the 238 U/ 204 Pb and 232 Th/ ²⁰⁴Pb values can be either higher or lower, probably depending on the particular combination of altered and unaltered phases remaining in the leached residue. It is important to note that, except in the glasses, neither set of values necessarily corresponds to those in the pristine rock.

The total range in $\varepsilon_{Nd}(t)$ (unless otherwise specified, in the text hereafter we use isotopic values for leached residues or glass, when available) is considerable: +10·3 to -0·6. That for (⁸⁷Sr/⁸⁶Sr)_t is also large, from 0·70277 to 0·70431. In contrast to the Alexander Island basalts, most of the age-adjusted drillhole-basalt data plot within or very close to the MORB fields in the Nd–Sr isotope

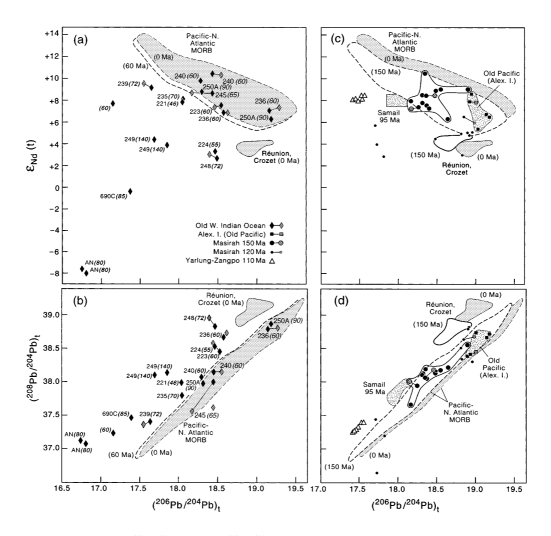


Fig. 4. Age-corrected $\varepsilon_{Nd}(t)$ (a, c) and $(^{208}\text{Pb}/^{204}\text{Pb})_i$ (b, d) vs $(^{206}\text{Pb}/^{204}\text{Pb})_i$. Acid-leached (or glass) and unleached pairs are indicated by light and dark symbols, respectively, linked by tie lines. As in Fig. 3, the field for the modern Pacific–North Atlantic MORB source at 150 Ma (c, d) and 60 Ma (a, b) is unshaded, and additionally assumes an average ${}^{238}\text{U}/{}^{204}\text{Pb} = 5$ and ${}^{232}\text{Th}/{}^{238}\text{U} = 2\cdot3$ in the source mantle (White, 1993); that for the Réunion–Crozet field at 150 Ma (c, d) is also unshaded and assumes source values of 12 and 3·3, respectively (Peng & Mahoney, 1995). Site numbers are shown adjacent to data points in (a) and (b), with ages in parentheses. Diamonds labeled AN at $\varepsilon_{Nd}(t) \sim -8$ are for Afanasy-Nikitin Seamount (Mahoney *et al.*, 1996); that at $({}^{206}\text{Pb}/{}^{204}\text{Pb})_t \sim 17\cdot 2$ is for a seamount southwest of Afanasy-Nikitin estimated at 60 Ma (our unpublished data, 1997). Field for Samail ophiolite is from data of Chen & Pallister (1981) (for Pb isotopes) and McCulloch *et al.* (1981) [for $\varepsilon_{Nd}(t)$].

diagram (Fig. 3). An interesting exception is the chemically and petrographically unusual alkalic lava from Site 690C (Schandl *et al.*, 1990) on Maud Rise, which may have formed in association with a jump of the ancestral Southwest Indian Ridge toward the Bouvet hotspot in the 85–100 Ma period (e.g. Barker *et al.*, 1990); this sample has anomalously low (87 Sr/ 86 Sr)_t (0.70376) for its low $\varepsilon_{Nd}(t)$ (–0.6).

Pb isotopes also exhibit a wide range, with (²⁰⁶Pb/ ²⁰⁴Pb)_t, varying from 17·40 (the Site 690C sample) to 19·28 (the Site 236 glass). Many of the old western Indian Ocean basalts resemble modern Indian MORB and hotspot islands in that their age-corrected Pb and Nd isotopic ratios place them on the low-²⁰⁶Pb/²⁰⁴Pb side of the Pacific–North Atlantic MORB-source field in Fig. 4a and b. The oldest lavas, from Site 249, were erupted at ~140 Ma during early spreading between Africa and southern Greater Indo-Madagascar, and display a strong Indian-Ocean-type signature in both Fig. 4a and b. Samples from Sites 690C, 239, 248, 235, 224, and 221, with ages between ~85 and 46 Ma, also have clear Indian-Ocean-type isotopic signatures. In addition, dredged lavas from Afanasy-Nikitin Seamount (Fig. 2), erupted on or near the western Southeast Indian Ridge at ~80 Ma in a location far from continental landmasses, recently have been shown to possess lower $\epsilon_{\rm Nd}(t)$ (–8) and (²⁰⁶Pb/²⁰⁴Pb)_t

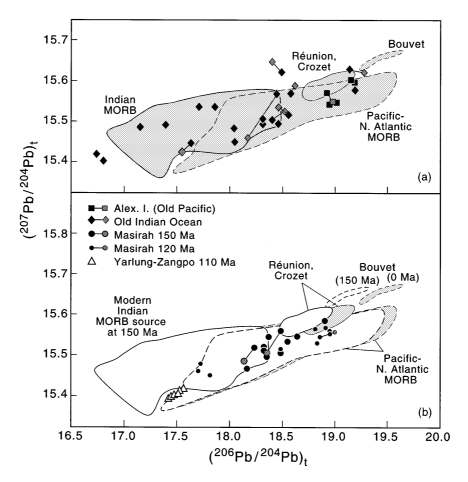


Fig. 5. Age-corrected $(^{207}Pb/^{204}Pb)_i$ vs $(^{206}Pb/^{204}Pb)_i$. Symbols, fields, and data sources are as in Figs 3 and 4, plus Sun (1980) for Bouvet.

(16.77) values than any modern Indian Ocean (or any other oceanic) lavas (see Fig. 4). Thus, Indian-MORB-type isotopic compositions clearly were present in the old western Indian Ocean mantle, in good agreement with results for old lavas from the eastern Indian Ocean (Lanyon, 1995; Pyle *et al.*, 1995; Weis & Frey, 1996).

However, data for several sites overlap the Pacific– North Atlantic MORB-source array. As noted earlier, such characteristics appear to be very rare within the main part of the Indian Ocean domain today (Fig. 1). The Site 250A samples (including an N-MORB and an E-MORB), the two flows we analyzed from Site 240, and the upper of two petrographically distinct units (Fisher *et al.*, 1974) at Site 236 have values that fall within the Pacific–North Atlantic MORB-source field in both Fig. 4a and b. The same is true for the Site 245 basalt, despite the significant difference in age-corrected Pb isotopic values between the leached and unleached splits of this sample. Moreover, the Site 250A-26-5 (90 Ma) E-MORB and the Site 236-33-3 glass–crystalline-rock pair (60 Ma) have age-corrected ($^{206}Pb/_{204}Pb)_t > 19$,

significantly greater than seen for any modern Indian MORB (most of which have ²⁰⁶Pb/²⁰⁴Pb <18·4). [Although it was not age-corrected, an even higher presentday ²⁰⁶Pb/²⁰⁴Pb value of 19.580 was reported by Hart (1988) for a Site 250A lava deeper in the core than our E-MORB sample, with a value of 19.322.] Some modern Indian Ocean island lavas have values around 19, but they also have markedly lower $\epsilon_{\! Nd}\,(\sim +4 \mbox{ or less vs } +6{\cdot}2$ and +7.4), higher relative ²⁰⁸Pb/²⁰⁴Pb, and usually higher ⁸⁷Sr/⁸⁶Sr (e.g. ~0.704 for Réunion and Crozet). Two samples display notably ambiguous 'mixed' isotopic characteristics, one from the lower unit at Site 236 and one from Site 223 (both \sim 60 Ma). In Fig. 4a, values for both the unleached and leached splits of these samples fall near the edge of but within the estimated Pacific-North Atlantic MORB-source array for 60 Ma. However, in Fig. 4b, data for both samples lie well above this field. Along the present-day spreading ridges, qualitatively similar mixed signatures have been found only in some lavas at the fringes of the Indian MORB domain (Mahoney et al., 1992; Pyle et al., 1992; Volker & et al., 1993).

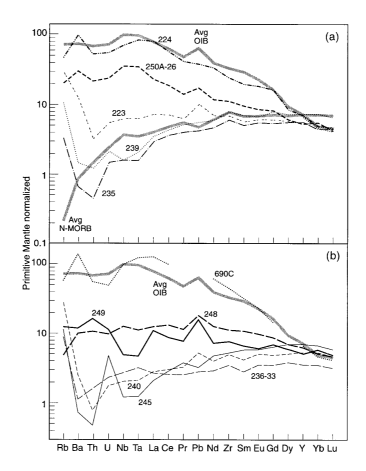


Fig. 6. Incompatible element patterns of several western Indian Ocean drillhole lavas. Shown for comparison are average N-MORB and oceanic island basalt (OIB) patterns. Arrangement into two panels is for clarity only. Primitive-mantle normalizing values and N-MORB and OIB averages are Sun & McDonough's (1989). The Site 690C pattern is from data of Schandl *et al.* (1990). The Ta value for the Site 245 pattern is inferred from the measured Nb value assuming Nb/Ta = 17.

Masirah

Masirah is an island off the coast of Oman that contains well-preserved exposures of uplifted abyssal oceanic crust. An older suite of MORB-type magmatic and ultramafic rocks is present, as well as a younger group of magmatic rocks, principally alkalic basalts and their differentiates but also amphibole-clinopyroxene gabbros and rare oceanic granites (e.g. Moseley & Abbotts, 1979; Abbotts, 1981; Moseley, 1990; Smewing et al., 1991; Gnos & Perrin, 1996; Nägler & Frei, 1997). Recent dating reveals that the MORB-type suite is ~150 Ma, whereas the younger suite is ~120 Ma (Smewing et al., 1991; Immenhauser, 1996; Nägler & Frei, 1997). The Masiran seafloor appears to have formed on the slow-spreading, transform-fault-dominated ridge system (e.g. Fisher et al., 1986) linking the main Tethyan and early western Indian Ocean (northeastern Somali Basin) spreading centers in the narrow basin between northwestern Greater Indo-Madagascar and the northeastern corner of Arabia-Africa (see Fig. 7) (e.g. Mountain & Prell, 1990; Smewing et al., 1991; G. Mountain, personal communication, 1993). (Note that the similar-age basement at Site 249 was formed in the Mozambique Basin farther southwest, to the southwest of Madagascar.) The cause of the later magmatism at ~120 Ma is uncertain but may be related to lithospheric fracturing and passage of the region near a hotspot (Meyer *et al.*, 1996; Nägler & Frei, 1997).

Primitive-mantle-normalized element patterns of representative tholeiitic and alkalic Masiran lavas (the latter represented by an E suffix in the tables) are shown in Fig. 8a and b. Low, MORB-like abundances of incompatible elements characterize the 150 Ma tholeiitic basalts, and their patterns range from typical N-MORB type to transitional-MORB type (relatively flat). The patterns of the 120 Ma alkalic lavas slope generally upward to the left and broadly resemble those of many oceanic island basalts [compare the average OIB pattern in Fig. 8; see also Meyer *et al.* (1996)]. Overall, the patterns are relatively smooth. However, Rb and Ba can be mildly to dramatically enriched or depleted relative

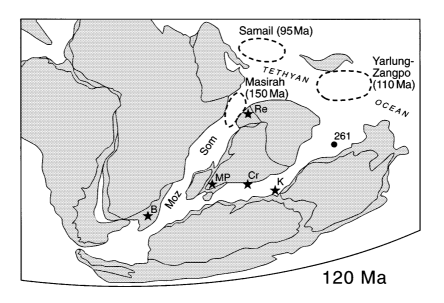


Fig. 7. Reconstruction at 120 Ma (after Lawver & Gahagan, 1993). Approximate 120 Ma location of Masiran crust is shown, along with that of Site 261 and positions of the present Réunion (Re), Bouvet (B), Crozet (Cr), Marion–Prince Edward (MP), and Kerguelen (K) hotspots. Rough indications of the location of the Yarlung–Zangpo and Samail crust are also shown. Moz, Mozambique Basin; Som, Somali Basin.

to Th in the visibly more altered samples analyzed (e.g. MA-401, MSX-219E). Small to moderate troughs or peaks at Pb are present in some patterns as well, and a substantial negative Eu anomaly can be seen in the pattern for MSX-219E, a trachytic lava.

As with the drillhole samples, acid-leaching of the Masiran rocks reduced their (87Sr/86Sr)t values while causing negligible changes in $\varepsilon_{Nd}(t)$ and only modest ones in age-corrected Pb isotope ratios. In several cases, a large reduction in (⁸⁷Sr/⁸⁶Sr), occurred: for example, from 0.70466 to 0.70305 for MSX-171. In the Nd-Sr isotope diagram (Fig. 3), the data for most of the Masiran rocks plot in or very close to the estimated 150 Ma MORBsource field (in the broad area of this diagram where Pacific-North Atlantic and Indian fields overlap), although leaching failed to bring the Sr isotope values of MA-401 or MSX-219E into this field. The plagioclase separate-which we did not leach-of a gabbro (MSX-71g) also yielded higher $({}^{87}\text{Sr}/{}^{86}\text{Sr})_t$ (0.70335) than the leached splits of basalts MSX-75 and MSX-171 [with Sr isotopic values of ~0.7030 at similar $\varepsilon_{Nd}(t)$], evidently indicating that the plagioclase was somewhat affected by interaction with seawater.

For the 150 Ma samples, $\varepsilon_{Nd}(t)$ ranges from +10.5 to +6.4, indicative of intrinsic heterogeneity in the mantle source; however, eight of the 11 samples have values between +9.0 and +7.6. Values of $(^{206}Pb/^{204}Pb)_t$ vary from 18.17 to 18.88, with eight of the samples having ratios between 18.32 and 18.64. Moreover, most of the Pb isotope data define good positive correlations close to 150 Ma reference isochrons in plots of present-day $^{206}Pb/^{204}Pb$ vs $^{238}U/^{204}Pb$ and $^{208}Pb/^{204}Pb$ vs $^{232}Th/$

²⁰⁴Pb (Fig. 9a,b), consistent with alteration largely occurring within a few million years after eruption for most of these samples. In both Fig. 4c and d, data for the 150 Ma rocks lie within the Pacific–North Atlantic MORBsource field, except for the leached split of sample MA-401, which falls slightly to the left of this field.

The 120 Ma alkalic lavas have lower $\varepsilon_{Nd}(t)$ than the 150 Ma rocks, from +6.0 to +2.9. Six of the ten samples show little variation in age-corrected $(^{206}\text{Pb}/^{204}\text{Pb})_{t}$ ratios, which are between 18.81 and 19.00. In Fig. 4c, data points for these six samples plot toward the low- $\varepsilon_{Nd}(t)$ end of the Pacific-North Atlantic MORB-source array and in or near the Réunion-Crozet source field. In Fig. 4d, the data for these alkalic lavas lie beneath this field, and straddle the Pacific-North Atlantic MORBsource array. The remaining four samples analyzed have significantly lower $(^{206}\text{Pb}/^{204}\text{Pb})_t$, between 18.49 and 17.71, yet have $\varepsilon_{Nd}(t)$ in exactly the same range as the other alkalic lavas. In Fig. 4d, the data point for one of these four samples falls well below the Pacific-North Atlantic MORB field, unlike any modern ridge or oceanic island basalts, and in Fig. 9c and d data for these samples lie far from the array defined by the others. We infer that relatively recent alteration (including variable loss of Pb) has seriously disturbed the Pb isotope systematics of these four alkalic lavas.

In addition to our work, Nägler & Frei (1997) have recently analyzed 120 Ma Masiran oceanic granites and amphibole-bearing gabbros, as well as several 150 Ma samples, for Nd and Pb isotopic ratios and U, Pb, Nd, and Sm abundances. Their age-corrected $\epsilon_{Nd}(t)$ and $(^{206}\text{Pb}/^{204}\text{Pb})_t$ values are similar to ours in that their data

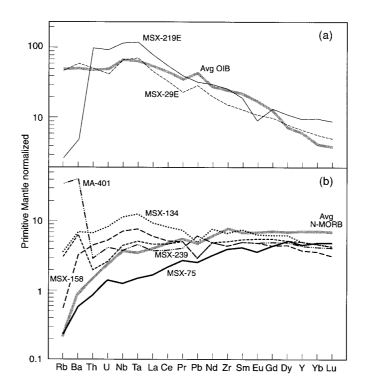


Fig. 8. Incompatible element patterns of selected Masirah 120 Ma alkalic lavas (a) and 150 Ma tholeiites (b).

fall largely within the estimated 150 Ma Pacific–North Atlantic MORB-source field of Fig. 4c or, for one granite, close to the Réunion–Crozet source field; also, as with our alkalic lavas, their 120 Ma gabbros and granites tend to have lower $\varepsilon_{Nd}(t)$ values than those of the 150 Ma rocks.

Yarlung-Zangpo suture zone

The samples from the Yarlung–Zangpo suture zone were collected from along the length of basalt outcrops to the southwest of Lhasa in the Xigaze area. Elemental and petrographic analyses of lavas from several of the same general areas show them to be chemically N-MORB (Pearce & Deng, 1988) altered to prehnite-pumpellyite or lower greenschist facies, with plagioclase replaced by albite and with abundant secondary groundmass actinolite and chlorite (Girardeau et al., 1985). The lavas were erupted at an eastern Tethyan spreading center north of Greater Indo-Madagascar and to the south of the Tibetan block (see Fig. 7; Pozzi et al., 1984). An age of 110 Ma was determined by Marcoux et al. (1982) from Radiolaria in cherts interbedded conformably with pillow basalts. Previous isotopic work on magmatic rocks consisted of Pb isotope and Pb and U (but not Th) abundance measurements by Göpel et al. (1984). Their results revealed a rough U–Pb whole-rock isochron (120 ± 10 Ma) which gave nearly the same age as the paleontologically derived age, indicating that the alteration affecting these rocks (including their U/Pb ratios) occurred fairly soon after eruption and that the rocks had remained nearly closed systems thereafter.

As with the Alexander Island basalts, Sr isotope ratios of the Yarlung-Zangpo samples are elevated relative to $\varepsilon_{\rm Nd}(t)$, and acid-leaching yielded only modest reductions in $({}^{87}\text{Sr}/{}^{86}\text{Sr})_t$ (to 0.70380–0.70406; Fig. 3). These results are consistent with the similar level of alteration in the two suites, specifically with the extensive replacement of original plagioclase by albite (in the Yarlung-Zangpo basalts, clinopyroxene is also partly replaced with secondary phases). Values of $\varepsilon_{Nd}(t)$ show very little variation in the Yarlung–Zangpo samples, all being between +8.0and +8.5, indicating a nearly homogeneous mantle source. The age-corrected Pb isotope ratios also vary only slightly: the spread in $(^{206}\mathrm{Pb}/^{204}\mathrm{Pb})_{t},$ for example, is only 17.42-17.55 (in the same range as for Göpel et al.'s samples) and only 37·27–37·38 in $(^{208}\text{Pb}/^{204}\text{Pb})_{t}.$ Our results confirm the good overall positive correlation of present-day 206Pb/204Pb with 238U/204Pb as well. Most important for our present purposes, the Yarlung-Zangpo basalts define a very small field with a clear Indian-MORB-type signature in both Fig. 4c and d.

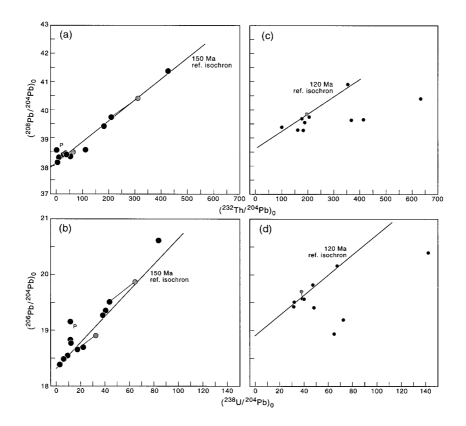


Fig. 9. Present-day ${}^{208}\text{Pb}/{}^{204}\text{Pb}$ vs ${}^{232}\text{Th}/{}^{204}\text{Pb}$ and ${}^{206}\text{Pb}/{}^{204}\text{Pb}$ vs ${}^{238}\text{U}/{}^{204}\text{Pb}$ for 150 Ma Masiran rocks (a, b) and for 120 Ma lavas (c, d); symbols as in Fig. 4. Also plotted are 150 Ma and 120 Ma reference isochrons. P is plagioclase separate.

DISCUSSION AND CONCLUSIONS

Consistent with the results for the Jurassic South Pacific lavas from Alexander Island, basalts of roughly similar age from several drill sites in the North Pacific recently have been shown to also possess Pacific–North-Atlantictype age-adjusted isotopic signatures (Janney & Castillo, 1997). Allowing for relatively small changes in ε_{Nd} and Pb isotope ratios resulting from radiogenic ingrowth in the source mantle during the last 150 my, the available data thus imply that the Pacific MORB mantle in the Jurassic and earliest Cretaceous was isotopically very similar to that of today.

In the western Indian Ocean, the ~140 Ma Site 249 basalts reveal that Indian-Ocean-type isotopic compositions were present from almost the very beginning of the ocean itself—at least in some locations. However, the ~150 Ma rocks of Masirah and Site 261 (Weis & Frey, 1996) essentially lack normal Indian-MORB-type signatures. The crust at both sites formed near the southern boundary of the Tethyan Ocean, Masirah on the northwest and Site 261 on the northeast side of what would later become Greater India (see Fig. 7). Thus, the data for these sites provide no evidence that the Indian Ocean isotopic signature was inherited from Tethyan asthenosphere or, indeed, that the Indian Ocean mantle domain existed in anything like its present form north of East Gondwana in the Late Jurassic.

On the other hand, the Yarlung-Zangpo basalts demonstrate the existence of Indian-MORB-type mantle in at least a part of the equatorial Tethys by 110 Ma. In addition, some high-quality, age-corrected Pb and Nd isotopic data have been published for the Samail ophiolite of Oman, the crust of which appears to have formed at a low northern latitude (Perrin et al., 1994) at ~95 Ma (e.g. Tilton et al., 1981), to the west of that preserved in the Yarlung-Zangpo suture (see Fig. 7). In Fig. 4d, the Samail Pb isotope data occupy a restricted field above the Pacific-North Atlantic MORB-source array, and in Fig. 4c the combined Nd and Pb isotopic results [which were obtained on different samples (Chen & Pallister, 1981; McCulloch et al., 1981)] define a rectangle that again largely falls outside the Pacific-North Atlantic MORB-source field (adjusted to a 95 Ma position; not shown in figure). Thus, the available data indicate an essentially Indian-Ocean-type mantle source for Samail crust at 95 Ma [see also Benoit (1997)].

Although very few locations have been studied as yet, the differences between the older and younger Tethyan sites suggest the possibility of a temporal change in asthenospheric composition. A period of continental lithospheric thinning was followed by spreading in the eastern Indian Ocean at ~135 Ma (e.g. Powell et al., 1988), and one possibility is that some Indian-MORBtype asthenosphere flowed northward out of the widening rift between Greater Indo-Madagascar and Australia-Antarctica, into parts of the Tethyan region. On the west side of Greater Indo-Madagascar, spreading and linkage with the Tethys began around 160–170 Ma (e.g. Lawver & Gahagan, 1993) (slightly before the 150 Ma Masiran rocks were formed), and northward flow of Indian-Oceantype mantle may have begun earlier there. If so, to reach the paleolatitude of the Samail crust by 95 Ma, such asthenospheric flow must have occurred at rates of ~70-100 mm/yr; in comparison, rates of 25-40 mm/yr are indicated for westward flow of Pacific-type asthenosphere into the southeastern Indian Ocean between Antarctica and Australia since 43 Ma (Pyle et al., 1992, 1995). Possible driving forces for significant asthenospheric outflow from the young, narrow Indian Ocean include (1) the ascent into the upper mantle of the large starting-plume heads of the Kerguelen, Marion, Bouvet, and Crozet hotspots [all of which may have reached the base of the lithosphere in the 200-120 Ma period; e.g. Storey (1995) and references therein], and (2) the upward advection of the 660 km boundary between upper and lower mantle proposed to have occurred beneath East Gondwana early in the Cretaceous in response to accelerated circum-Pacific subduction of slabs into the lower mantle (Larson & Kincaid, 1996). However, dispersion of stringers of Indian-Ocean-type mantle into the Tethys is not a unique explanation of the existing isotopic data. Although evidence is lacking for a widespread, pre-110 Ma Tethyan upper mantle possessing Indian-MORB-type characteristics, an alternative possibility is that the Tethyan asthenosphere may have contained pockets of both Pacific-North-Atlantic-type and Indian-MORB-type compositions (the latter presumably generated by the same types of processes as those acting in the Indian Ocean mantle) as far back as 150 Ma or even earlier. Study of Tethyan basalts from other locations along the southern Asian suture belt is required to evaluate these possibilities.

The 120 Ma alkalic lavas of Masirah have oceanisland-like elemental signatures, and six have isotopic compositions rather similar to those of modern Réunion (21°S) and Crozet (46°S) hotspot basalts, particularly in Fig. 4c. Recent plate reconstructions in the hotspot reference frame (e.g. Curray & Munasinghe, 1991; Lawver & Gahagan, 1993; Müller *et al.*, 1993) differ somewhat but suggest the Masiran region was situated between about 15°S and 25°S at 120 Ma, and thus appear to permit a Réunion connection of some sort (e.g. see Fig. 7) while ruling out an association with the Crozet hotspot, other than perhaps a 'far-field' effect related to dispersion of Crozet plume-head material. Meyer et al. (1996) recently suggested that the Marion (Prince Edward) hotspot, located to the west of Crozet, could have been the source of the alkalic magmatism; however, like Crozet, the Marion hotspot is located at \sim 46°S. Also, the volcanoes of the Marion hotspot have different isotopic compositions (e.g. $\varepsilon_{Nd} = +5.7$ to +7.4, 206 Pb/ 204 Pb = 18.5–18.6; Hart, 1988; Mahoney *et al.*, 1992) from those of Réunion and Crozet. However, the origin of the 120 Ma Masiran rocks remains problematic, because although the isotopic signature of the main component in the Réunion plume appears to have changed little in the last 66 my (White et al., 1990; Peng & Mahoney, 1995), the Réunion hotspot is believed by most workers to have appeared only shortly before 66 Ma, the Deccan Traps event being interpreted as the hotspot's initial, plume-head phase [e.g. Basu et al. (1993) and references therein].

Both Indian-Ocean-type and some Pacific-North-Atlantic-type isotopic signatures are preserved in the old western Indian Ocean drill sites. Moreover, when results for the drillholes are combined with those for Afanasy-Nikitin Seamount, a 3 $\varepsilon_{\rm Nd}$ unit wider total spread of $\varepsilon_{\rm Nd}(t)$ is encompassed than is found for present-day Indian MORB and oceanic islands (compare Figs 1a and 4a). This comparison excludes the Early Cretaceous Kerguelen and Naturaliste plateaux, which reach even lower $\varepsilon_{Nd}(t)$ values than Afanasy-Nikitin but, unlike Afanasy-Nikitin, probably contain blocks of continental lithosphere [e.g. see Mahoney et al. (1996) and references therein]; it also excludes recent lavas (with less extreme values; e.g. Weis et al., 1992) erupted through the thick lithosphere of the Kerguelen Plateau. As such, the total range in $\varepsilon_{Nd}(t)$ for modern (0–10 Ma) Indian Ocean basalts is -4.0 to +11.3, whereas that for the old lavas is -8.0 to +10.3, with both maximum and minimum values seen in lavas formed at 60-80 Ma (note that Site 261 in the northeastern Indian Ocean, although formed at a Tethyan ridge, yielded an even higher value of +14.4; Weis & Frey, 1996). This difference is much greater than achievable by plausible isotopic evolution (i.e. 'aging') in the source mantle since the Cretaceous.

The larger isotopic range observed for the old basalts is remarkable in view of the very sparse sampling of old Indian Ocean crust relative to sampling of the presentday spreading centers and islands; further, it is unlikely that the few existing old sites have fortuitously sampled the full isotopic range present in old Indian Ocean seafloor. Admittedly, substantial sections of the modern spreading system remain unsampled or have only recently been dredged, and could potentially harbor more extreme isotopic compositions than found elsewhere. However, continuing work on the largest previously unsampled stretch of the system, the Southeast Indian Ridge between the Australian-Antarctic Discordance and St Paul Island, thus far reveals isotopic values that are all well within the previous range for modern Indian MORB (Hall et al., 1995; unpublished data, 1997). Nor does it appear that the difference in isotopic ranges can be ascribed in any simple way to differences in spreading rate. Greater isotopic heterogeneity along ridges generally is associated with slower spreading rates-for example, both the highest and lowest $\epsilon_{\scriptscriptstyle Nd}$ values observed in the modern Indian Ocean $(+11\cdot3 \text{ and } -4\cdot0)$ are found on the central part of the very slowly spreading Southwest Indian Ridge (Mahoney et al., 1992). However, the old Indian Ocean samples with both the highest and lowest ϵ_{Nd} values were formed (also on-ridge or very near-ridge) in the 60-80 Ma period of super-fast spreading (e.g. Fisher & Sclater, 1983). The most straightforward interpretation of the existing data is therefore that the Indian Ocean asthenosphere was isotopically more heterogeneous in the past and is gradually becoming better mixed on a time scale of tens of millions of years. In turn, greater heterogeneity in the not too distant past, including the presence in some locations of Pacific-North-Atlantic-type compositions, is most consistent with a relatively young origin for the Indian Ocean mantle domain; that is, one not considerably older than the age of the Indian Ocean itself.

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