INTRODUCTION

The behavior of noble metals in igneous systems is poorly understood, and their extraction from an evolving magma has been related to a variety of processes, including the separation of PGE alloys (e.g., Keays, 1982), chromite (e.g., Agiorgitis and Wolf, 1978; Stockman and Hlava, 1984), an immiscible sulfide phase (e.g., Keays and Crocket, 1970; Naldrett et al., 1979; Barnes and Naldrett, 1985; Brugmann et al., 1987), and silicates, particularly olivine (Crocket, 1981; Brugmann et al., 1987; Fryer and Greenough, in press). This lack of understanding of their behavior reflects, in large part, the difficulty that has traditionally been associated with accurately determining the abundance of these elements in basaltic rocks where concentration levels are in the low-to-sub-ppb range. The recent development of an analytical procedure using inductively coupled plasma-mass spectrometry (ICP-MS; Jackson et al., 1990) allows routine analysis with mean instrumental detection limits in the sub-ppb range.

This paper outlines the concentration of the noble metals Pt, Pd, Rh, Ir, and Au in intraoceanic basaltic volcanic rocks of the Lau Basin, and evaluates their abundance with respect to the supra-subduction signature of the source magma. The paper provides the first accurate data on noble metal concentrations in basalts and andesites from a modern backarc basin.

The Lau Basin forms part of the Tonga Arc system and is an ideal natural laboratory to study arc-backarc magmatism and the role of slab-controlled, crust-mantle interaction in governing noble metal abundances. The arc lies within a zone of complexly interacting microplates that have developed along the boundary between the Indo-Pacific and Atlantic plates. Plate boundaries within this region have evolved rapidly since the commencement of subduction in the Eocene, resulting in the progressive movement of the southwestern boundary of the Pacific Plate away from the Australian continent.

The Tonga Arc records a relatively simple evolutionary picture. It is divisible into three tectonic elements. Extending westward from the Tonga Trench, these are (1) the active magmatic arc and forearc (Tonga Ridge), (2) the active backarc basin (Lau Ridge), and (3) the remnant arc (Lau Ridge). Before the initiation of seafloor spreading in the Lau Basin in the Miocene, the Lau and Tonga ridges constituted a single magmatic arc-forearc succession. The Lau Basin was the first backarc basin in which active seafloor spreading was documented (Karig, 1970), and it provides one of the best developed records of spreading processes. On Leg 135, a transect across the backarc investigated the process of crustal generation. Sites drilled range from Miocene-age basalts adjacent to the Lau Ridge, which were generated during the early stages of arc rifting and backarc basin development (Site 834), to approximately 1 Ma crust near the main spreading ridge for the Lau Basin (Site 836; Fig. 1).

The geochemical data available for the Lau Basin (e.g., Frenzel et al., 1990; Hawkins and Melchior, 1985; Jenner et al., 1987; Vallier et al., 1991; Volpe et al., 1988) indicate a range of compositional groups (and mantle-source components), from basaltic and andesitic rocks showing a strong arc (slab-derived) signature to basaltic rocks showing an N-MORB signature.

Recent cruises to the Valu Fa Ridge, the active spreading ridge within the southeastern position of the basin, have recognized active hydrothermal vents (Fouquet et al., 1990, 1991) and reported high concentrations of gold and visible gold in the sulfides (Herzig et al., in press). This is the first reported occurrence of visible, primary gold in sulfides from active seafloor vents. Establishing the noble metal concentrations within the Lau Basin basaltic flows will allow a direct measurement of gold levels in potential source basaltic to hydrothermal system.

SAMPLE DESCRIPTION

Samples for noble metal analysis were selected on the basis of petrographic descriptions and major and trace element analyses to provide a range of lithologic and compositional types. Samples analyzed from Sites 834, 835, 836, and 839 include both phric and aphyric basalts and hyaloclastic glass, and represent a wide composi-
Figure 1. Distribution of major morphotectonic elements within the Tonga arc system and location of Leg 135 drill sites.
tional range, on the basis of Mg# and the Cr and Ni contents. The following sample descriptions were derived from shipboard data (Parsons, Hawkins, Allain, et al., 1992).

The locations of the Leg 135 drill sites (834, 835, 836, and 839) are given in Figure 1. Sites are located in the western and central Lau Basin, and all sites lie west of the youngest crust in the basin, which is forming at the Central (CLSC) and Eastern Lau (ELSC) Spreading Centers (Vala Fa). Sites 834 and 835 lie in small basins east of the Lau Ridge, which formed through rifting of older arc lithosphere, whereas Sites 836 and 839 lie close to the ELSC on true oceanic lithosphere related to organized back arc basin spreading.

Site 834 is located along the northwest margin of the Lau Basin, 100 km east of the axis of the Lau Ridge, and represents the oldest crust drilled on Leg 135. Igneous basement consists of a series of flows and sills. Seven samples from Units 2, 5, 6, and 7 were analyzed from this site and, on the basis of the biostratigraphy of sediments intercalated with the flows, range in age from lower Miocene (4 Ma) to upper Miocene (6 Ma). Unit 2 is an aphyric, massive vesicular basalt with a groundmass of plagioclase, clinopyroxene, and minor interstitial magnetite and olivine. Unit 3 is a moderately altered aphyric, vesicular diabase. The groundmass is dominated by plagioclase microlites and larger clinopyroxene oikocrysts rich in plagioclase laths with lesser amounts of intergrown clinopyroxene. Magnetite is the main opaque oxide phase, but ilmenite, Cr-spinels, and sulfide globules are also present in minor to trace amounts. Glassy, interstitial phases of the groundmass are altered to clay minerals. Unit 6 consists of aphyric to sparsely phyric, vesicular basalt. A glomeroporphyritic texture of plagioclase, clinopyroxene, and rare olivine is locally developed in a groundmass of plagioclase, clinopyroxene, olivine, magnetite, and a mesostasis that is typically altered to clays. Unit 7 is a highly phryic plagioclase basalt. At 76 m, it is the thickest unit delineated at Site 834, and it consists of a series of pillowed flows. Vesicles are absent or extremely rare. Phenocrysts are glomeroporphyritic and consist mainly of euhedral plagioclase but also include minor olivine. Groundmass mineral phases in approximate decreasing order of abundance are plagioclase, clinopyroxene, magnetite, Cr-spinel, and ilmenite. The mesostasis shows minor alteration to clays.

Site 835 is located 200 km east of the axis of the Lau Ridge. Igneous basement is constrained to be of late Pliocene age (3.4 Ma) on the basis of biostratigraphic data obtained from the immediately overlying sediments. A vesicular, plagioclase-clinopyroxene-olivine basalt is the only lithologic unit intersected at this site. Plagioclase is the main mineral phase followed by clinopyroxene. Olivine is a minor phase and magnetite is present in trace amounts. A continuous size gradation occurs from phenocryst to groundmass phases. The mesostasis shows minor alteration to clays.

Site 836 is situated 50 km west of the active ELSC (Vala Fa), which constitutes the main spreading center within the basin. Igneous basement at Site 836 may represent the oldest lithosphere generated at the ELSC. Five rock units were recognized in the basement, which on the basis of biostratigraphic data from intercalated sediments has a maximum age of middle Pleistocene (<0.8 Ma). Units 1 and 2 are hyaloclastites and consist of vesicular, sparsely plagioclase phytic glassy andesitic gravel. Samples analyzed from these units were repeatedly cleaned ultrasonically with distilled, deionized water before being crushed, and effectively represent clean glass samples. Unit 4 is divisible into two subunits of aphyric basalt separated by a thin sediment intercalation. It consists of plagioclase and clinopyroxene with minor olivine, magnetite, and interstitial mesostasis, variably altered to clays.

Site 839 is located some 75 km west of the ELSC (Vala Fa). Igneous basement was divided into eight lithologic units. Overlying sediments are of late Pliocene age (2 Ma). The single sample analyzed for noble metal abundances from this site occurs within Unit 1, a vesicular, sparsely to moderately phryic olivine basalt. The holocrystalline groundmass consists of plagioclase and clinopyroxene with minor olivine and magnetite.

**ANALYTICAL METHODS**

Gold, Pd, Pt, Rh, and Ir were determined by ICP-MS on 10% aqua regia solutions after NiS fire assay preconcentration of 10 g of rock powder. Complete method blanks, for this study, are 0.54, 0.82, 1.34, 0.16, and 0.05 ppb (rock), respectively, for the above elements; these were subtracted from the raw data. The standard deviations of the method blanks (N = 9) are 0.35, 0.42, 0.13, 0.02 and 0.02 ppb (rock) for Au, Pd, Pt, Rh, and Ir, respectively; they provide the best estimate of the limits of detection for this method. It should be recognized that the complete method blanks include all sources of uncertainty such as reagent blanks, instrument memory, and backgrounds, and it is a much more reliable (and conservative) indicator of data reliability for analyses near “detection limits” than the more traditional 3 sigma above background. The reader is referred to Jackson et al. (1990) for details of the complete procedure. Replicate analyses of the reference material SARM-7 have given 245, 1452, 3923, 227, and 77 ppb Au, Pd, Pt, Rh, and Ir, respectively, with relative standard deviations of 7% or better for all elements. All PGE values are within 10% of the certified concentrations. Replicate analyses of our in-house komatiite standard (BT-4) give relative deviations of 10% or better for concentrations (1–10 ppb) more closely approximating the rock compositions of this study.

**RESULTS**

Chemical data for the Leg 135 volcanic rocks are presented in Table 1. Major, trace, and rare earth element (REE) data are taken from Hawkins and Allain (this volume). The gold contents of all samples are surprisingly high, ranging from 1.0 to 11.4 ppb. Rhodium and Ir contents are very low, ranging from below detection (approximately 0.02 ppb) to 0.08 ppb. The Pd and Pt contents range from 0.3 to 4 ppb, which is relatively high compared with other oceanic basalts (Table 2). Gold appears to be anomalously enriched relative to both the PGE and Cu. Within this range of noble metal abundances, two groups are recognized. Sites 836 and 839 generally have higher abundances of Rh, Pd, and Pt than do Sites 834 and 835. Gold and Ir values cover similar ranges at all sites.

In general terms, the major and trace element chemistry of the samples ranges from arc-like to more MORB-like in the sequence from Sites 839, 836, 835, and 834, respectively (e.g., Fig. 2). The highest noble metal contents are found in the sites (836 and 839) that have the strongest arc signatures. At least for Au, Pd, and Pt, noble metal abundances are not related to the degree of “fractionation” of the lavas, as expressed by Mg# (Fig. 3). The bulk of the samples show a restricted range of Mg# (60–70), although the two fresh andesitic glass samples (Hole 836A) are distinctly lower (35). The only significant trend between noble metal content and major or trace elements is with Cu (Fig. 4). Lavas from Sites 839 and 836 have distinctly higher average Cu contents than those from Sites 835 and 834 (123 vs. 74 ppm), which parallels their much higher Pd and Pt contents (3.42 and 1.79 ppb vs. 0.78 and 0.52 ppb, respectively). Although both groups have a similar overall range of Au values, mean Au abundances at Sites 836 and 839 are also much higher (7.7 ppb) compared with Sites 834 and 835 (3.9 ppb). Within the high-Cu group, however, Pd and Pt contents decrease with increasing Cu contents. This trend corresponds with increasing SiO2 contents. These relationships suggest that the compositional differences between the two groups cannot be ascribed to fractionation processes from similar primary magma sources.

Primitive mantle-normalized, noble metal distributions for the Lau Basin samples (Fig. 5) demonstrate the anomalous Au concentrations for all samples relative to the PGE and Cu, as well as the strong fractionation within the PGE. In particular, Ir is strongly depleted relative to Pd, with a mean Pd/Ir ratio of approximately 90 for Sites 834 and 835, and almost 100 for Sites 836 and 839 (Table 2).
observed sulfide blebs have similar noble metal contents to other units coexisting basaltic liquids; however, the Leg 135 rock units with immiscible sulfide phase could effectively remove noble metals from them as well as the composition of the subducting slab and the relative metal abundances in oceanic basalts (the products of mantle melting) is compromised by the very limited data currently available on noble metal distributions observed are more likely related to mantle source processes such as Mg# (Fig. 3) and Ni and Cr contents suggests that the noble metal contents with low-pressure magmatic differentiation indicate that this is the case. Indeed, the negative correlation of Pt and Pd with Cu at Sites 836 and 839 mitigates against a significant role for sulfides in these magmas. The overall lack of correlation of noble metal contents with low-pressure magmatic differentiation indicates such as Mg# (Fig. 3) and Ni and Cr contents suggests that the noble metal distributions observed are more likely related to mantle source processes rather than to liquid fractionation.

Within supra-subduction zone environments, potential source reservoirs for any erupturn magma include variably depleted and enriched mantle, as well as components released from the downgoing slab into the overlying mantle-wedge. Evaluation of the relative contribution of each of these possible sources to Leg 135 volcanic rocks is compromised by the very limited data currently available on noble metal abundances in oceanic basalts (the products of mantle melting) as well as the composition of the subducting slab and the relative behavior of possible components released into the mantle-wedge. Table 2 compares the noble metal and Cu abundances and ratios of the two Leg 135 groups with the best available data for a variety of mantle melts, as well as MOR-sulfide data. It should be recognized from the leg. Furthermore, sulfide separation will not fractionate the noble metals from each other because of the similar distribution coefficients (Kd's). Although we cannot rule out the possibility that the noble metal concentration differences between the two groups of lavas is a result of the previous separation of an immiscible sulfide liquid from the Site 834 and 835 magmas, no compelling data exists to indicate that this is the case. Indeed, the negative correlation of Pt and Pd with Cu at Sites 836 and 839 mitigates against a significant role for sulfides in these magmas. The overall lack of correlation of noble metal contents with low-pressure magmatic differentiation indicates such as Mg# (Fig. 3) and Ni and Cr contents suggests that the noble metal distributions observed are more likely related to mantle source processes rather than to liquid fractionation.

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Table 2. Comparison of average noble metal (ppb) and Cu abundances (ppm) from Leg 135 with oceanic basalts, boninites, komatities, and hydrothermal seafloor sulfide.

<table>
<thead>
<tr>
<th></th>
<th>Cu</th>
<th>Au</th>
<th>Rh</th>
<th>Pt</th>
<th>Ru</th>
<th>Os</th>
<th>Ir</th>
<th>Pd/Ir</th>
<th>Au/Pd</th>
<th>Au/Ir</th>
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<tr>
<td>834-835</td>
<td>74</td>
<td>3.9</td>
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<td>0.52</td>
<td></td>
<td></td>
<td>0.01</td>
<td>89</td>
<td>5.04</td>
<td>390</td>
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<tr>
<td>836-839</td>
<td>123</td>
<td>7.75</td>
<td>0.06</td>
<td>1.79</td>
<td></td>
<td></td>
<td>0.04</td>
<td>98</td>
<td>2.26</td>
<td>194</td>
</tr>
<tr>
<td>MORB</td>
<td>115</td>
<td>3.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.12</td>
<td>48</td>
<td>0.61</td>
<td>30</td>
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<tr>
<td>Indian MORB</td>
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<td>2.2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.06</td>
<td>283</td>
<td>0.13</td>
<td>37</td>
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<tr>
<td>Leg 115</td>
<td>147</td>
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<td>0.31</td>
<td>7.3</td>
<td>0.22</td>
<td>0.11</td>
<td>74</td>
<td>0.4</td>
<td>29</td>
<td></td>
</tr>
<tr>
<td>Ocean Island tholeiite</td>
<td>2</td>
<td>2.3</td>
<td></td>
<td></td>
<td></td>
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<td>0.25</td>
<td>8.8</td>
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<td>Ocean Island alk.</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>0.30</td>
<td>5.0</td>
<td>0.33</td>
<td>1.67</td>
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<td>Komatites</td>
<td>69</td>
<td>2.4</td>
<td></td>
<td>8.2</td>
<td>5.8</td>
<td>2.4</td>
<td>1.9</td>
<td>4.8</td>
<td>0.26</td>
<td>1.26</td>
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<tr>
<td>MOR sulfide</td>
<td>1,138</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:
1. Average of Sites 834 and 835 from Table 1.
2. Average of Sites 836 and 839 from Table 1.
3. Average of DSDP Leg 37, Hole 332B-12 (13 samples); Mid-Atlantic Ridge, Vema Fracture Zone (1 sample); Mid-Atlantic Ridge 22°S (1 sample); Mid-Atlantic Ridge, FAMOUS area (1 sample); South Atlantic, Bouvet Triple Junction (1 sample); South West Triple Junction, Indian Ocean (1 sample). Each sampling location given equal weight. Sources: Leg 37 (Au, Pd, Ir) = Crocket and Teruta (1977); all other data from Hertogen et al. (1980).
4. Average of 2 samples of Indian Ocean MORB from DSDP Leg 24, Site 238 (Hertogen et al., 1980).
5. Average of 3 groups of modern boninites (Bonin Island, Cape Vogel, Mariana forearc; DSDP Site 458 = Hamlyn et al., 1985).
7. Au = average of 26 Hawaiian basalts (Gottfried et al., 1972); Pd = average of data for 5 samples from Hawaii, Maui (Hawaii), James Island (Galapagos), Albermarle Island (Galapagos), and Réunion (Crocker and Skippen, 1966); Ir = average of 10 samples from Hawaii (Gottfried and Greenland, 1972).
8. Au and Ir = average of 4 samples, Tahiti (Crocket et al., 1973); Pd = average of 4 samples from Tutuila (Samoa), Tahiti, St. Michael (Azores), and Ascension Island (Crocket and Skippen, 1966), 4 samples from Tahiti (Crocket et al., 1973).
9. Composite average of 3 most primitive Gorgona, 3 most primitive Alex flow (Au, Pd, Ru, Os, and Ir = Brugmann et al., 1987), and 42 spinifex-textured komatities from western Australia and Munro Township, Canada (Keays, 1982).
10. Single massive sulfide analysis, Middle Valley, northern Juan de Fuca Ridge (Crocker, 1990). Additional data: average Au and Ir abundances and Au/Ir ratio for MORB sulfide globules are 10,000 ppb, 1,200 ppb, and 16; massive sulfides from the Mid-Atlantic Ridge TAG area are 2,990 ppb, 0.50 ppb, and 6,000 (Crocker, 1990).

Figure 2. Plot of Ti vs. V for Leg 135 samples. Field for arc and ocean-floor basalts from Shervais (1982). Open circle = Site 834, open square = Site 835, solid circle = Site 836, solid square = Site 839. For Samples 135-834B-13R-1 and 135-836A-3H-3, major and trace element values used in data analysis were taken from Samples 134-834B-12R-3 and 134-836A-3H-4, respectively, which on the basis of REE (Table 1) and other data are geochemically similar. Sample 135-836B-5R2-65-74C is a duplicate sample split of Sample 135-836B-5R2-65-74 cm, and major and trace values from this latter sample were used in plotting the former.
that the comparative data is of variable quality and often is only available for a limited suite of the noble metals.

Figure 6 compares average Leg 135 data with selected igneous rock groups. The relatively strong fractionation of the Leg 135 PGE is similar to that observed in Leg 115 Indian Ocean hotspot basalts (Fig. 6A) and MORB (Fig. 6B). However, the distinctive enrichment of Au with respect to both Cu and Pd in Leg 135 volcanic rocks is not shown by other mantle melts such as MORB, ocean-island basalts, and komatiites, all of which have Au/Pd ratios = 1 (Table 2 and Fig. 7). Anomalous primitive mantle-normalized Au values were observed in some supra-subduction zone settings (e.g., boninites; Fig. 6B), but these are depletions in gold relative to Pd (Au/Pd = 0.13; see Table 2). The only potential source component for Leg 135 samples that is characterized by high Au/Pd ratios is hydrothermal fluids and their derived precipitates (e.g., sediments and sulfides; Table 2 and Fig. 7). Gold concentrations in these precipitates are 2–3 orders of magnitude higher than for other oceanic crustal components. Even small additions of this component from the downgoing oceanic crust to the mantle melt zone

Figure 3. Plots of the noble metals Pt, Pd, Au, and Pd+Pt vs. Mg# for Leg 135 lavas. Symbols are as in Figure 2.

Figure 4. Plots of the noble metals Au, Pd, Pt, and Pd+Pt vs. Cu for Leg 135 lavas. Symbols are as in Figure 2.
Noble Metal Abundances in Lau Basin

Figure 5. Primitive mantle normalized distribution patterns of Cu and the noble metals Au, Pd, Pt, Rh, Ru, and Ir for Leg 135 Sites 834 and 835 (A) and Sites 836 and 839 (B). Mantle normalizing values for Cu (28 ppm) are from Sun (1982) and for Au, Pd, Pt, Rh, and Ir (1.2, 4.4, 8.3, 1.6, and 4.4 ppb) are 0.15 chondritic values as reported by Naldrett (1981) and summarized in Barnes et al. (1988). The order of elements has been reversed from normal to show increasing compatibility from left to right as is standard in petrogenetic studies.

Figure 6. Primitive mantle normalized noble metal concentrations of Leg 135 lavas compared to average (A) komatiite and Leg 115 hotspot basalts, and (B) MORB and boninite. Data from Table 2. Open rectangle = Sites 834 and 835, and filled rectangle = average of Sites 836 and 839.

could account for the high mean Au/Pd ratios of the Leg 135 samples (Sites 834 and 835, Au/Pd = 5.04; Sites 836 and 839, Au/Pd = 2.26). Comparison of Leg 135 data with other supra-subduction zone rock types is limited to boninites, for which only Au, Pd, and Ir data are available (Hamlyn et al., 1985). Boninites are characterized by extreme PGE fractionation (Pd/Ir ratio = 280; see Table 2) and strong depletion in both Au and Cu relative to Pd (Fig. 6). Recent suggestions that boninites represent melting of a young, hot subducted slab that lacked a sedimentary cover (J.A. Pearce, pers. comm., 1992) are consistent with our suggestion that the high Au/Pd ratio of Leg 135 basalts reflects a sediment signature from the downgoing slab.

The high concentrations of gold and visible gold-bearing phases in sulfides from the Valu Fa Ridge (Eos, 1990; Herzig et al., in press) may be related to anomalous Au contents like those contained in the Leg 135 samples. Geochemical and isotopic analyses of Valu Fa lavas have previously established a strong slab-signature in their mantle source (Jenner et al., 1987; Valu Fa et al., 1991). However, Au concentrations of Lau Basin lavas are less than an order of magnitude greater than MORB, whereas Valu Fa hydrothermal precipitates are enriched by several orders of magnitude relative to any potential basaltic source, thereby indicating that source-rock composition alone cannot explain the presence of Au-bearing phases along the spreading ridge. Herzig et al. (in press) have suggested that efficient precipitation processes are crucial in ultimately determining the presence or absence of gold-bearing hydrothermal phases from the Valu Fa Ridge.

CONCLUSIONS

Leg 135 backarc basin lavas are characterized by anomalously high Au contents and strongly fractionated relative PGE abundances. The Au signature may reflect separation of Au from the PGE within an oceanic hydrothermal system and subsequent release of this component to the mantle wedge during subduction of the downgoing slab. However, a detailed analysis of the relative contribution from other mantle sources (e.g., variably depleted or enriched mantle) to the Leg 135 magmas is limited by the lack of data on magmas derived from such sources.
ACKNOWLEDGMENTS

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REFERENCES


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Figure 7. Plot of Pd vs. Au for Sites 834 and 835 (shaded square), Sites 836 and 839 (filled square), average MORB (asterisk), Leg 115 basalts (larger open circle), Indian Ocean MORB (filled circle), komatiite (open diamond), boninite (plus sign), and hydrothermal sulfide (smaller filled circle).