30. CHARACTERIZATION AND CORRELATION OF MEGASCOPIC TEPHRAS IN SITE 792
CORES FROM THE IZU-OGASAWARA FOREARC BASIN (JAPAN) BY TRACE ELEMENTS
AND $^{87}$Sr/$^{86}$Sr AND $^{143}$Nd/$^{144}$Nd ISOTOPES

Per Kristian Egeberg, Arild O. Brunfelt, and Arne S. Stabel

ABSTRACT

Glass shards from 26 megascopic tephras (1–29 cm thick) were separated and analyzed for 26 elements and $^{87}$Sr/$^{86}$Sr and $^{143}$Nd/$^{144}$Nd isotope ratios. The isotope data allowed us to assess the degree of alteration of the samples. The separates contain less than 10% of palagonitized glass or secondary alteration minerals. Correlation diagrams and multivariate analyses have been used to characterize the glass shards in a tectonic setting and group the tephras according to their magmatic affinity. Four tholeiitic groups, two calc-alkaline groups, and one group with glass of shoshonitic affinity can be distinguished. The tholeiitic groups form a time series; however, judging from the rare-earth-element patterns and the temporal variations of the Ce/Yb ratios, the evolution is not unidirectional from strong to weak tholeiitic affinity.

The tephras have been correlated with volcanism along the Izu-Ogasawara and Ryukyu arcs by means of trace element distribution, $^{87}$Sr/$^{86}$Sr ratios, and their magmatic affinity. The tholeiitic layers were derived from volcanoes on the Izu-Ogasawara Arc, most probably from the volcanic island Hachijojima. Only one group of the calc-alkaline layers can have derived from the Izu-Ogasawara Arc. The other calc-alkaline layers and the two layers with shoshonitic affinity probably derive from volcanism along the Ryukyu Arc.

INTRODUCTION

The western Pacific has been characterized by extensive explosive volcanism throughout the Neogene. Documentation of the volcanic record has been obtained through the efforts of several Deep Sea Drilling Project/Ocean Drilling Program (DSDP/ODP) drilling in the Philippine Sea, along the Palau Kyushu Ridge, in the Nankai Trough and Shikoku Basin, and across the Mariana and Izu-Ogasawara arcs. Studies of ash layers from this area have focused mainly on sedimentological and volcano-chronological aspects, and several determinations of the major element composition of volcanic glasses have been reported (e.g., Furuta and Arat, 1980; Migdisov et al., 1981; Packham and Williams, 1982; Fujioka et al., 1986; Poullet et al., 1986). However, only a very limited amount of data on trace elements and rare-earth elements (REEs) are available (e.g., Migdisov et al., 1981; Fujioka et al., 1986), and none of these discuss the degree of alteration of their samples. There are no published data on Sr isotopes of submarine volcanic ash layers from this region.

The main objectives of this study were to evaluate the extent of seawater alteration of the isolated volcanic glasses, to characterize their composition in a tectonic context, and to correlate the ash layers with volcanism along the Izu-Ogasawara Arc by means of trace element concentrations, REE patterns, and $^{87}$Sr/$^{86}$Sr and $^{143}$Nd/$^{144}$Nd isotope ratios.

MATERIALS AND METHODS

Site 792 is situated in the Izu-Ogasawara forearc basin southeast of the volcanic island Aogashima (Fig. 1). The geological setting has been described elsewhere (Taylor, Fujioka, et al., 1990). At Site 792 the Shipboard Scientific Party registered 200 ash layers from Hole 792A. In this study we have included 23 of these and an additional 3 from Hole 792E (Table 1). The ash layers were not indurated. We have numbered the layers from 1 to 26, and for reference purposes Table 1 also includes the numbers assigned to the ash layers from Hole 792A by the Shipboard Scientific Party (Taylor, Fujioka, et al., 1990).

Isolation of Glass

The samples (about 10 cm$^3$ of wet sediment) were dispersed ultrasonically in distilled water, and the 63–125 µm size fraction was isolated by wet sieving. In some layers no glass particles were greater than 63 µm. Because the risk of contamination and alteration effects increase with decreasing particle size, these samples were discarded. After being cleaned with distilled water, the 63–125 µm size fractions were dried for 1 week at 160°C. The density values of five samples that, by microscopic examination, proved to be clean samples of clear glass were determined, using a 0.7-g sample and a 5.000-cm$^3$ pycnometer. The results are listed in Table 2.

The average density of 2.35 g/cm$^3$ is similar to that of cristobalite. Based on these results, two solutions of sodium polytungstate of density 2.25 and 2.45 g/cm$^3$ were prepared for centrifugation-assisted, gravitational separation of phillipsite and biogenic opal (density 1.7–2.2 g/cm$^3$) and crystalline silicates, aluminosilicates, and carbonates (density greater than 2.5 g/cm$^3$). Following heavy liquid separation, the glass separates were kept in the centrifuge tubes and rinsed with 10 volumes (10–20 cm$^3$) of distilled water, two volumes of 1N supra pure HCl, and finally with 10 volumes of distilled water.

Instrumental Neutron Activation Analyses

After drying at 160°C, two subsamples of ~40 mg from each layer were weighed in aluminum foil and irradiated for 24 hr at 1.2×$10^{13}$ n/cm$^2$ x s (at the Institute for Energy and Technology, Kjeller, Norway). Half of the subsamples were only exposed to epithermal irradiation by shielding with Cd (Steinnes, 1971). The samples were allowed to cool for 7 days and were counted four times during the following 1.5 months using the coaxial Ge(Li) detector, the planar HpGe detector, and a multichannel analyzer (at the Mineralogical and Geological Museum, Oslo, Norway). The United States Geological Survey standards BCR-1, BHVO-1, and MAG-1 were run with the samples. In addition, five splits of Sample 26 were run to determine the precision of the analyses (reported as relative standard deviation [Rel. std. %] in Table 1).
Rb-Sr and Sm-Nd Mass Spectrometry

The separation of Rb, Sr, Nd, and Sm from about 20-mg samples and mass spectrometric analyses were performed at the Geological Museum, University of Oslo, using standard ion exchange techniques and a Finnigan MAT 262 mass spectrometer in static mode (Nd), and a VG 354 machine in multicollector-dynamic mode (Sm, Sr, and Rb). The isotope dilution measurements were performed with the VG 354 machine in single-collector, peak-switching mode (Rb, Sr, Nd, and Sm). The techniques used in Oslo are described in detail by, for example, Mearns (1986) and Neumann et al. (1988). During the period of analysis, mean measured standard values were $^{87}\text{Sr}/^{86}\text{Sr}$: NBS987Sr $= 0.71024 \pm 0.00010$; $^{143}\text{Nd}/^{144}\text{Nd}$: Johnson and Matthey C 321 $= 0.511108 \pm 0.000002$; and La Jolla Nd $= 0.511851 \pm 0.000002$. The concentrations of Sr, Nd, Sm, and Rb determined by isotope dilution are superior to the results obtained by INAA (standard deviations in Table 1).

RESULTS AND DISCUSSION

The number of chemical analyses of unaltered tephra is small compared with the enormous volume of published analyses of lavas. This is because of (1) the ease by which the composition of tephra is changed by diagenetic alteration, (2) the difficulty of recognizing mixtures in tephras, and (3) the problem of sorting of the tephra components during transport. For these reasons, tephra composition has been used to a very limited extent for deciphering magma evolution. Yet pyroclastic fragments such as glass shards are unique because they quench almost instantly on eruption and are not subjected to crystal fractionation processes occurring in slowly cooling flows. The potential for high-resolution dating of marine tephra should make these very attractive targets for studies of short-wave temporal variations in magma composition of a given tectonic segment.

This discussion will focus on three aspects of the tephra geochemistry: (1) the degree of alteration of the glasses, (2) chemical characterization in a tectonic context, and (3) correlation of the tephra with volcanism along the Izu-Ogasawara Arc.

The analytical results are listed in Table 1, which also includes the relative standard deviation (Rel. std. %), the depth of each layer in meters below seafloor (mbsf), and the approximate age (determined from the sediment accumulation curves prepared by the Shipboard Scientific Party on the basis of nannofossil and palaeomagnetic stratigraphy (Taylor, Fujioka, et al., 1990). No sample contained more than 41.6 ppm W and no element correlated significantly (at the 95%
信心水平）与 W。这表明该清理过程
减少的放射性强度来自于重的液体 (指重辐射物质) 完全
对所有元素解释了 except W。该指标
Fujimura et al. (this volume) 和 Rodolfo et al. (this volume) 辨认
区别出清晰的玻璃片和粒状玻璃片。该分选
分离程序有效地分离了清晰的玻璃片，未采样
含有超过 5%的玻璃片和粒状玻璃片。

### Degree of Alteration of the Glasses

火山玻璃是热力学不稳定的，并分解
为更近似于附近所有相联的阶段。火山玻璃
玻璃在不明显的不整合处被氧化，因此
存在可能出现的观点变化的主要原因
这个研究。

### Alteration of basaltic glass under marine conditions is fairly well documented (e.g., Furnes, 1980; Ailin-Byzuk and Sommer, 1981; Staedtigel and Hart, 1983), and includes conservation of Fe and Ti (Furnes observed loss of Ti) and loss of other major elements except K. Some elements, in particular Rb and Cs, show strong enrichment during alteration. The alteration of silicate glass differs from the alteration of basaltic glass in many ways. The initial stage involves diffusion-controlled hydration and alkali ion exchange but only minor overall chemical changes. This is commonly followed by a stage of glass matrix destruction and precipitation of secondary phases (Fisher and Schmincke, 1984). The clay-altered silicate glass is also enriched in Fe and Ti; in contrast to altered basaltic glass, it is depleted in Cs, K, and Rb, however. Given this background, the positive correlation between TiO$_2$ and Fe$_2$O$_3$ (Fig. 2) need not be an igneous trend; rather, it could be the result of seawater alteration and/or inclusion of secondary alteration products.

Seawater/basaltic glass interactions gave rise to a considerable exchange of Sr (Staudigel and Hart, 1983), and the alteration products formed from silicate glass were enriched in Sr (Zielinski, 1982). The $^{87}$Sr/$^{86}$Sr ratio of arc-derived volcanic material (0.703-0.705) is quite different from that of Quaternary seawater (0.7090-0.7092). The Sr/Nd concentration ratio of seawater is about 2.0 $\times$ 10$^5$ times greater than in arc-derived volcanic products. Thus, seawater alteration reactions
will have a strong impact on the $^{143}$Nd/$^{144}$Nd ratio but will leave the $^{143}$Nd/$^{144}$Nd ratio of the glass samples virtually unchanged.

During exchange with seawater, the evolution of the $^{143}$Nd/$^{144}$Nd isotope ratio of the glass is related to the evolution of the $^{143}$Nd/$^{144}$Nd isotope ratio of the glass through Equation 1:

$$\frac{^{143}Nd}{^{144}Nd} = \frac{^{143}Nd^{87}Sr}{^{86}Sr} - \left( ^{143}Nd^{87}Sr\right)_{eq} \cdot \left( ^{143}Nd\right)_{eq} \cdot \left( ^{143}Nd\right)_{eq}$$

where $g$, $gi$, and $sw$ denote altered glass, initial glass, and seawater respectively. On a $^{143}$Nd/$^{144}$Nd vs. $^{87}$Sr/$^{86}$Sr diagram, all our observations plot inside or very close to the mantle array (Fig. 3). The isotope ratios of the glasses are similar to the values reported for basalts from the same area. We take this as an indication that the separates do not contain appreciable amounts of palagonitized glass or secondary alteration products. However, the seawater alteration line does allow for about 10% exchange of Sr for a few samples. The fact that the samples define a linear parallel to the mantle array indicates that even the samples with the most radiogenic Sr-isotope ratios (0.7050-0.7052) do not contain significant amounts of palagonitized glass or secondary alteration products. Figure 3 indicates the presence of at least three different glass populations. This will be discussed further below.

### Chemical Characterization of the Glasses

Several different schemes have been used to classify and discriminate between magma types using element and trace element concentrations. The La/Ta ratios of the samples (Fig. 4) are close to that found for island-arc tholeiites and calc-alkaline lavas from Japan (Jorin and Treuil, 1977; Wood et al., 1979) and are quite distinct from that of N- and E-type mid-ocean ridge basalts (MORBs). On the Th/Hf-Ta diagram (Fig. 5), all samples plot in the field of destructive plate-margin basalts and differentiates as defined by Wood (1980). These observations further support our contention that the glasses represent unaltered samples of island-arc volcanism. The samples span the entire calc-alkaline and tholeiitic
fields, and there are also indications of several other populations, one of intermediate composition and several of tholeiitic affinity.  

Identification of those ash layers that may derive from the same source is interesting both for corollary purposes and because it can provide a clue to how arc-magmas evolve. In an attempt to identify those layers that have the most similar source, we have run a cluster analysis. Because we are mainly interested in primary compositional signatures given in Table 1. 

If the maturation of the arc was synchronous along the segment of the volcano or a suite of volcanoes fed by magma of similar composition (these volcanoes may be distributed along the arc) and, (2) that each layer is characterized by low Ta/Yb and Th/Yb ratios. We do not have full major element analyses to confirm the classification of the layers into shoshonitic, calc-alkaline, and tholeiitic groups; nevertheless, for the purpose of reference and simplification of the following discussion, we will refer to the layers as shoshonitic, calc-alkaline, and tholeiitic with the subdivisions shown in Figures 6 and 7 and Table 3. 

Comparing Figures 6 and 7 reveals the presence of the same four tholeiitic groups (Cluster 3, 4, 5, and 7); and the single ungrouped layer is characterized by low Ta/Yb and Th/Yb ratios. We do not have full major element analyses to confirm the classification of the layers into shoshonitic, calc-alkaline, and tholeiitic groups; nevertheless, for the purpose of reference and simplification of the following discussion, we will refer to the layers as shoshonitic, calc-alkaline, and tholeiitic with the subdivisions shown in Figures 6 and 7 and Table 3. 

Comparing Figures 6 and 7 reveals the presence of the same groups. When the full “more-HYG” signatures are compared, calc-alkaline group C2 has a larger affinity toward the tholeiitic groups than toward the other calc-alkaline group. 

There are two different interpretations of the grouping of the layers: (1) that each group represents ash layers derived from one volcano or a suite of volcanoes fed by magma of similar composition (these volcanoes may be distributed along the arc) and, (2) that each group represents a particular stage of arc magma evolution and that the layers in each group may derive from several different volcanoes. If the maturation of the arc was synchronous along the segment of the arc that fed tephras to Site 792, then the second interpretation predicts temporal separation of the groups. Figure 8 shows that, indeed, except for one layer in T1, the tholeiitic groups are separated in time. This
but the material was insufficient to repeat the analyses. Data from the Izu-
Ogasawara, Volcano, and Mariana arcs were taken from Lin et al. (1990) and
Hochstaedter et al. (1990). The seawater alteration line (divided into fraction
of Sr exchanged) was estimated with Equation 1 (see text) and the following
conditions for seawater: Sr = 8 ppm, Nd = 2.6 ± 10^{-6} ppm, \(^{87}\)Sr/\(^{86}\)Sr = 0.70906,
and \(^{143}\)Nd/\(^{144}\)Nd = 0.51245. The following conditions apply for glass: Sr = 100
ppm, Nd = 15 ppm, \(^{87}\)Sr/\(^{86}\)Sr = 0.704, and \(^{143}\)Nd/\(^{144}\)Nd = 0.51288.

Figure 3. Isotopic ratios of Nd and Sr in glass shards isolated from tephras in
Holes 792A and 792E. The point marked with "?" is regarded as suspicious,
but the material was insufficient to repeat the analyses. Data from the Izu-
Ogasawara, Volcano, and Mariana arcs were taken from Lin et al. (1990) and
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Figure 4. La vs. Ta in glass shards isolated from tephras in Holes 792A and
792E. Lines for Japanese island-arc tholeiites and calc-alkaline lavas as well
as N- and E-type MORBs are taken from Wood et al. (1979).

indicates that the second interpretation is correct but does not exclude
the first interpretation. During the last 1 m.y., the composition of the
tholeiitic arc volcanism might have changed from T1 through T4 and
T3 and into T2 (Fig. 8).

The large-ion-lithophile–high-field-strength (LIL-HFS) diagram (Fig. 9A) shows that the indicated evolution from the oldest (T1) to the
most recent group (T2) is unidirectional only for Tb and Yb. For the
remaining elements the evolution appears to be better described as a cyclic
process. The same picture emerges from the rare-earth patterns (Fig.
10A). The two temporally most distant groups (T2 and T1) are more
similar to each other than to any of the intermediate groups.

The view is generally held (e.g., Jakes and White, 1972) that early
arc volcanism is characterized by tholeiitic series that grade into
calc-alkaline series with increasing age. This shift in magma affinity
takes tens of million of years. Based on the temporal distribution of
the groups (Fig. 8), the rare-earth patterns (Fig. 10A) represent an
evolution from less tholeiitic (T1) to more tholeiitic (T4 and T3)
volcanism and then a return to less tholeiitic volcanism (T2). This
apparent discrepancy between the generally accepted linear model
and the cyclic evolution found here may be taken as evidence that the
T2 group layers do not belong to the same line of evolution as the T1,
T3, and T4 group layers. A more detailed look at the evolution of the
Ce/Yb-ratio (Fig. 11) may be taken as a support of this hypothesis.
The rate and direction of inferred magma evolution is quite constant
from 1 to 0.2 m.y. when an abrupt reversal of the trend occurred.
However, in a study of ash layers from North Pacific arcs, Scheideg-
ger et al. (1980) observed rapid transitions between calc-alkaline and
tholeiitic affinities during the last 2 m.y. These short time fluctuations
may represent noise superimposed on the long-term transition from
tholeiitic toward calc-alkaline volcanism. A comprehensive discus-
sion of the temporal trends is beyond the scope of this presentation.

The strong Ti depletion (Figs. 9A and 9B) is similar for all groups and
underlines the importance of the crystallization of Fe and Ti
oxides, hence the positive correlation between Fe and Ti (Fig. 2).
However, the strong Ta depletion is unique for the tholeiitic layers
and may indicate different modes of Ti removal in the calc-alkaline
and shoshonitic systems.

The calc-alkaline and shoshonitic groups exhibit strong light-rare-
earth-element (LREE) enrichment (Fig. 10B) with Ce/Yb ratios in-
creasing systematically from 5.5 to 17.0 as one moves through the
calc-alkaline field and into the shoshonitic field (Fig. 7).

Source Correlation

As revealed by the distribution of ash layers (e.g., Kennett, 1981;
Machida, 1981), the atmospheric circulation over Japan and the
surrounding oceans was dominated by westerly winds throughout the
Quaternary. Site 792 is situated on the western flank of the volcanic
front; thus, most probably the layers are derived from volcanoes on
the arc itself. However, considering the widespread tephras produced
by some volcanoes (Machida, 1981), derivation from the mainland
or even the Ryukyu Arc cannot be excluded. The probability of local
derivation increases with increasing layer thickness. The average
thickness of the layers included in this study is about 10 cm; only two
layers are less than 5 cm, and the thinnest is 1 cm (Layer 16, Table 1).
Also, increasing particle size favors local derivation. We have only
included layers containing more than about 100 mg/10 cm\(^2\) (wet
sample) of glass particles in the 63–125 \(\mu\)m size range.

The layers cover the entire range of high SiO\(_2\) island-arc volcanism
from tholeiitic to calc-alkaline and shoshonitic series. The view is
generally held (e.g., Jakes and White, 1972) that these series are
related both laterally and stratigraphically. The tholeiitic series repre-
sent immature island-arc volcanism that grades into calc-alkaline and
shoshonitic series with increasing age and distance from the volcanic
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17 30
19 14 9 7 5
4 3

Figure 6. Dendogram showing the result of a clustering analysis with Cs/Rb, Ba/Rb, La/Rb, Ce/Rb, Ta/Rb, Th/Rb, and U/Rb as primary data. The data were normalized by subtracting the average element ratio and dividing this difference by the element concentration. The clusters are numbered sequentially from one to seven. The numbers identifying the layers refer to Tables 1 and 3.

Figure 7. Th/Yb vs. Ta/Yb in glass shards isolated from tephras in Holes 792A and 792E (after Pearce et al., 1981). MORB = mid-ocean-ridge basalt, WPB = within-plate basalt, VAB = volcanic-arc basalt, TH = tholeiitic, TR = transitional, ALK = alkaline, CA = calc-alkaline, SHO = shoshonitic, and IAT = island-arc tholeiite. The grouping indicated by the enveloping lines is the same as in Figure 6 (see Table 3 for key).

The volcanoes of the Izu Islands may be classified into two groups (Onuma et al., 1983; Notsu et al., 1983). The first group consists of the volcanoes situated on the volcanic front (Oshima, Omurodashi, Miyakejima, Mikurajima, Hachijojima, Aogushima, Torishima, and Sofugan). The second group consists of the volcanoes situated behind the volcanic front (Toshima, Udonejima, Niijima, Shikinejima, Kozushima, Onoharajima, and Inambajima) (Fig. 1). In accordance with the scheme of Jakes and White (1972), the volcanoes on the front are dominated by tholeiitic series whereas those behind the front are dominated by calc-alkaline series (Onuma et al., 1983). This will be used as the first-order criterion for correlation.

Compared with the Mariana and Volcano arcs, surprisingly little work has been done on the distribution of trace elements and REEs in the Izu-Ogasawara Arc. The only trace elements that have been systematically mapped are Ba and Sr (Onuma et al., 1981; Ishiki et al., 1982; Hirano et al., 1982). The Ba/Sr ratio will be used as the second-order criterion for correlation. However, this ratio is not fractionation independent. Judging from the pronounced Ti depletion front. Recently, this simplistic picture was modified by Stern et al. (1988), who found that shoshonitic volcanism may characterize the initial stages of arc construction after an episode of backarc rifting.

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Figure 8. Changes in the tholeiitic magma series affinities of glass shards isolated from tephras in Holes 792A and 792E during the last 1.0 m.y.

(Figs. 9A and 9B) and because the ash layers come from explosive volcanism, they probably represent highly differentiated compositions. Thus, this ratio is not ideal for correlation purposes. Nevertheless, because of the scarcity of other data, this is our best option.

Figure 12 compares the distribution of Ba/Sr ratios along the northern part of the Izu-Ogasawara Arc with the Ba/Sr ratios of the four tholeiitic, two calc-alkaline, and shoshonitic groups. The correlation of the various groups will be discussed separately.

Correlation of the Tholeiitic Layers

All of the tholeiitic layers except for Layer 43 are less than 1 m.y. old. Our first criterion dictates that these layers must derive from volcanoes on the volcanic front. Only volcanoes on Oshima and Hachijojima have produced tholeiites with sufficiently high Ba/Sr ratios to be source candidates (Fig. 12). Of these, Hachijojima is more probable because of its proximity to Site 792. However, the high Ba/Sr in the tholeiitic ashes relative to the lava analyses may reflect missing samples of extremely differentiated compositions on the islands.

Notsu et al. (1983) observed a remarkably narrow range of strontium isotope ratios in Quaternary volcanic rocks from the Izu-Ogasawara Arc (0.7033-0.7038). When including data from other sources (Kurasawa, 1979; Stern, 1982; Nohda and Wasserburg, 1981; Ikeda and Yuasa, 1989; Hochstaedter et al., 1990), this range only expands to 0.7032-0.7040. The 87Sr/86Sr values of the tholeiitic layers plot within or very close to this range (Fig. 13).
Figure 9. LIL-HFS diagrams averaged for each of the tholeiitic groups (A) and the calc-alkaline and shoshonitic groups (B) (modified from Pearce et al., 1981). The N-MORB data for normalization were taken from table 11.6 of Taylor and McLennan (1985), except for Yb, which is about 36% too high compared with the other heavy REEs.

Correlation of the Calc-alkaline Layers

The calc-alkaline layers span a period of 6.25 m.y. Based on the first criterion, these must derive from volcanoes behind the volcanic front. Only volcanoes on Kozushima and Niijima have produced calc-alkaline series with sufficiently high Ba/Sr ratios to be considered as sources (Fig. 12) (the uncertainty with respect to the coverage of the island samples must be kept in mind).

On the other hand, the $^{87}$Sr/$^{86}$Sr values of the Cl layers are considerably higher than ever reported for volcanism along the Izu-Ogasawara Arc (Fig. 13). These layers must have been derived from outside the arc. To the south of the Izu-Ogasawara Arc lie the Volcano and Mariana arcs (Fig. 1). Except for the old data published by Pushkar (1968), there are no reports of volcanism from the Mariana Arc with $^{87}$Sr/$^{86}$Sr values higher than 0.70362 (Woodhead and Fraser, 1985). The highest $^{87}$Sr/$^{86}$Sr value reported from the Volcano Arc is 0.70387 (Stern et al., 1984). On the other hand, the Ryukyu Arc volcanoes Kikai, Sakurajima, and Kirishima (Fig. 1) have $^{87}$Sr/$^{86}$Sr values of 0.70485–0.70539 (Notsu and Tatsunori Soya, 1987), 0.7051–0.7058 (Kurasawa et al., 1984), and 0.7045–0.7069 (Kurasawa et al., 1986), respectively. Most of the volcanoes from central Japan have $^{87}$Sr/$^{86}$Sr ratios (Notsu et al., 1989) that are too radiogenic to be regarded as sources for the Cl layers. These facts favor derivation of the Cl group layers from volcanoes on the Ryukyu Arc.

Correlation of the Shoshonitic Layers

The shoshonitic layers are 0.52 and 0.86 m.y. old. Jakes and White (1972) suggested that when shoshonitic rocks occur the age of the basement rocks is usually pre-Mesozoic. The evolution of the Izu-Ogasawara Arc is thought to have begun in the early-middle Eocene. Stern et al. (1988) found that shoshonitic volcanism may also be produced during the initial stage of arc construction following an episode of backarc rifting, and recently Lin et al. (1989) have in fact reported the occurrence of shoshonitic volcanism in the northern Mariana Arc. However, the high $^{87}$Sr/$^{86}$Sr values of the shoshonitic layers (Fig. 13) show that they cannot have been derived from the Izu-Ogasawara, Volcano, or Mariana arcs. In a comparison of volcanism along the three Japanese arcs, Aramaki and Ui (1982) found that K$_2$O-rich (2.5%–4.5%) ash-fall deposits associated with large calderas occur almost exclusively in the southwest Japan Arc (southwest Honshu and Ryukyu arcs). For this reason, the Ryukyu Arc volcanoes are the most likely sources of shoshonitic layers.

CONCLUSIONS

Glass shards from 26 megascopic tephras have been analyzed for 26 elements and $^{87}$Sr/$^{86}$Sr and $^{144}$Nd/$^{143}$Nd isotope ratios. The following three points were addressed: (1) the degree of alteration of the samples, (2) chemical characterization in a tectonic context, and (3) correlation of the tephras with volcanism along the Izu-Ogasawara and Ryukyu arcs.

The most important observations of our study are:

1. The distribution of $^{87}$Sr/$^{86}$Sr and $^{144}$Nd/$^{143}$Nd isotope ratios indicate that the glass separates contain less than 10% of palagonitized glass or secondary alteration minerals.
2. All tephras derive from destructive plate margins.
3. Correlation diagrams and cluster analyses show that the tephras may be grouped into four tholeiitic groups, two calc-alkaline groups, and one group with glass of shoshonitic affinity.
4. The grouping obtained by cluster analyses of the “more-HYG” element ratios is also reflected in the distribution of the $^{87}$Sr/$^{86}$Sr ratios.
5. The tholeiitic groups form a time series, but judging from the REE patterns and the temporal variations of the Ce/Yb ratios, the evolution is not unidirectional from strong to weak tholeiitic affinity.
6. The tholeiitic layers were derived from volcanoes on the Izu-Ogasawara Arc, most probably from the volcanic island Hachijojima. Only one group of the calc-alkaline layers can have derived from the Izu-Ogasawara Arc. The other calc-alkaline layers and the two layers with shoshonitic affinity probably derive from volcanism along the Ryukyu Arc.

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Table 3. Classification of layers based on trace element analyses.

<table>
<thead>
<tr>
<th>Group</th>
<th>Abbreviation</th>
<th>Cluster</th>
<th>Layers in each group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shoshonitic</td>
<td>S1</td>
<td>1</td>
<td>22,30</td>
</tr>
<tr>
<td>Calc-alkaline</td>
<td>C1</td>
<td>2</td>
<td>6,31,39</td>
</tr>
<tr>
<td>Tholeitic</td>
<td>T1</td>
<td>5</td>
<td>16,24,26,27,29,32,33</td>
</tr>
<tr>
<td></td>
<td>T2</td>
<td>7</td>
<td>3,4</td>
</tr>
<tr>
<td></td>
<td>T3</td>
<td>3</td>
<td>5,7,9,14,19,43</td>
</tr>
<tr>
<td></td>
<td>T4</td>
<td>3</td>
<td>20,21</td>
</tr>
</tbody>
</table>

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Figure 10. Chondrite-normalized REE patterns for the tholeitic groups (A) and
the calc-alkaline and shoshonitic groups (B). Because of instrumental
difficulties, some of the REE analyses are suspect. The following analyses are not
included in Figure 10A: La (Samples 3 and 9), Ce (Sample 26), and Eu (Sample 29).
The N-MORB data are taken from table 11.6 of Taylor and McLennan
(1985), except for Yb, which is about 36% too high compared with the other
heavy REEs. Chondrite concentrations are from Anders and Grevesse (1989).
Figure 11. Ce/Yb for the individual tholeiitic layers vs. time since deposition. The shaded area represents the evolution from tholeiitic compositions (T1) to less tholeiitic compositions (T3), and possibly a return to more tholeiitic compositions in the younger layers (T2).

Figure 13. A comparison of the $^{87}\text{Sr}/^{86}\text{Sr}$ values of the glass shards isolated from tephras in Hole 792A and Hole 792E with the range of $^{87}\text{Sr}/^{86}\text{Sr}$ values reported for volcanism along the Izu-Ogasawara Arc (see text).


