# Petrogenesis of alkalic and calcalkalic volcanic rocks of Mormon Mountain Volcanic Field, Arizona

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Abstract. The Cenozoic Mormon Mountain Volcanic Field (MMVF) of northern Arizona is situated in the transition zone between the Basin and Range and the Colorado Plateau. It is composed of alkalic to sub-alkalic basalts and calcalkalic andesites, dacites, and rhyodacites. Despite their spatial and temporal association, the basalts and the calcalkalic suite do not seem to be co-genetic. The petrogenesis of primitive MMVF basalts can be explained as the result of different degrees of partial melting of a relatively homogenous, incompatible element-enriched peridotitic source. The variety of evolved basalt types was the result of subsequent fractional crystallization of olivine, spinel, and clinopyroxene from the range of primitive basalts. Crustal contamination seems to have occurred, but affected only the highly incompatible element abundances. The formation of MMVF calcalkalic rocks is most readily explained by small to moderate amounts of partial melting of an amphibolitic lower crust. This source is LREE-enriched but depleted in Rb and relatively unradiogenic Sr ( ${}^{87}$ Sr/ ${}^{86}$ Sr ~0.7040). Calcalkalic rhyodacites may also be derived from andesitic parents by fractional crystallization. The overall petrogenesis of the MMVF complex is the result of intra-plate volcanism where mantle-derived magmas intrude and pass through thick continental crust.

# Introduction

The MMVF developed near the southwestern margin of the Colorado Plateau during the late Cenozoic (15–3 Ma). The MMVF is similar to other volcanic fields that mark the western, southern, and eastern boundaries of the Plateau [e.g. SW Utah (Lowder 1973; Best and Brimhall 1974); San Francisco Volcanic Field (Robinson 1913; Moore et al. 1976)] with respect to the variety of rock types present (alkalic basalts and calcalkalic andesites, dacites, and rhyodacites). These volcanic areas lie within the transition zone between the Basin and Range and the Colorado Plateau. This zone is characterized by numerous normal faults, high heat flow (Blackwell 1978), and a gradual thickening of the continental crust (Warren 1969), all of which might be expected to have some influence or be interrelated in some way with the petrogenesis of late Cenozoic lavas of this area (Lowder 1973). This paper considers the petrogenesis of MMVF and is part of a larger study of the Cenozoic volcanism of the transition zone undertaken in part to better understand the relationship between tectonism and volcanism in the region (Gust et al. 1984), and the genesis of anorogenic (in the sense of non-plate collision zone) calcalkalic magmatism.

There are several important issues in the petrogenetic history of the MMVF suite that deserve detailed discussion: 1) the origin of the variety of MMVF basalts; 2) the extent of superimposed contributions of fractional crystallization and crustal assimilation on the initial diversity of basalt compositions created by variable partial melting conditions; 3) the relationship between the basalts and the andesites; 4) the origin of the variety of intermediate and silicic rock types in MMVF; and 5) the importance of interaction of mantle-derived magmas with continental crust.

# **Regional geology**

MMVF extends over about 3,500 km<sup>2</sup> south of Flagstaff, Arizona (Fig. 1). Multiple flows of basaltic lavas, erupted from monogenetic cinder cones and NW-trending fissures, form a thin plateau-capping cover on sedimentary strata. These fissures parallel local fault trends and the margin of the Colorado Plateau (Mogollon Rim), suggesting a regional structural control on the volcanic eruptions. Mormon Mountain, the largest silicic center in the field, is composed of hornblende andesite flows overlain by dacite flows. A rhyodacite dome occurs near the southern flank of Mormon Mountain and a smaller rhyodacite dome crops out 10 km further south in a basaltic cinder cone. Small andesite domes occur throughout the field. The volume of basalt exceeds that of the silicic lavas, but precise estimates are difficult to make.

Basalts from Anderson Mesa and Oak Creek Canyon, delimiting the northern and western boundaries of the field, yield K – Ar ages of  $5.9 \pm 0.9$  Ma and  $6.0 \pm 0.3$  Ma, respectively (Damon et al. 1974). Basalts from the southern edge of the field, near Casner Butte and Pine are older (9–15 Ma, Peirce et al. 1979). A dacite from Mormon Mountain is dated at  $3.1 \pm 0.1$  Ma (R.B. Moore pers. comm. 1977), and an andesite from Table Mountain in the southern part of MMVF is dated at  $3.6 \pm 0.1$  Ma (T.M. Harrison pers. comm. 1980), both dates by K – Ar methods. Detailed stratigraphic relationships between different volcanic units are not clear in the field.

MMVF overlies Permian Coconino sandstone that caps a thick sequence of Paleozoic sedimentary rocks. Precambrian granites, schists, and gneisses comprise the upper crust of the Colorado Plateau. The lower crust consists of amphibolite, granulite, and eclogite as inferred from geophysical investigations (Prodehl 1970)

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Fig. 1. Location and sample locality maps of MMVF (*stippled area*). The inset shows the location of MMVF with respect to the Colorado Plateau and the Basin and Range. Outlined areas are other Tertiary volcanic fields which rim the Colorado Plateau

and petrological studies of high-pressure xenoliths in explosive volcanics (McGetchin and Silver 1972; Arculus and Smith 1979; Ehrenberg and Griffin 1979; Stoeser 1973; Keating et al. 1985).

# Petrology

## **Basalts**

MMVF basalts range from olivine nephelinite to olivine tholeiite with basanite and alkali olivine basalt being the most abundant. The basalts are aphanitic to fine-grained, with intersertal to intergranular textures; most are vesicular and some contain calcite or zeolite-filled amygdules. Olivine and clinopyroxene phenocrysts with euhedral spinel inclusions are common. Plagioclase phenocrysts are rare, except in hawaiites and olivine tholeiites. Modal phenocryst abundances range from 10–50%, and some clinopyroxene phyric lavas could be called ankaramites.

The composition of olivine phenocrysts within the basaltic suite ranges from  $Fo_{89}$  to  $Fo_{62}$ . Phenocrysts of clinopyroxene are sectorand oscillatory zoned and range from diopside to salite in the basanites and some alkali olivine basalts, and diopside to augite in other alkali olivine basalts, hawaiites and olivine tholeiites (Fig. 2). Octahedral spinel inclusions in clinopyroxene and olivine range from Mg-Al-rich chromites to chromian titanomagnetites. Spinels near the forsteritic core of host olivine phenocrysts have higher Mg numbers and Cr/(Cr+Al) ratios than spinels nearer the rim. Rare ilmenite phenocrysts or xenocrysts in basanites are magnesian – (13 wt.% MgO) and manganese-rich (1.3 wt.% MnO). Plagioclase phenocrysts (An<sub>82-70</sub>) are small (<2 mm), euhedral laths often occurring as glomerocrysts with clinopyroxene.

Groundmass olivine ( $Fo_{80-43}$ ) is partly altered to iddingsite. Groundmass clinopyroxene contains up to 11.0 wt.% Al<sub>2</sub>O<sub>3</sub> and 2.5 wt.% TiO<sub>2</sub>. Plagioclase laths in the groundmass are highly variable in composition, ranging from An<sub>89</sub> to An<sub>37</sub>. Minor groundmass phase assemblages include: 1) nepheline (Ca<sub>4</sub>Na<sub>83</sub>K<sub>13</sub>), sanidine and alkalic residuum (basanites); 2) anorthoclase and K-rich residuum (alkali olivine basalts); and 3) coexisting leucite (Ca<sub>2</sub>Na<sub>2</sub>K<sub>96</sub>) and nepheline (Ca<sub>10</sub>Na<sub>77</sub>K<sub>13</sub>) (olivine nephelinite). Apatite is ubiquitous and associated with anhedral patches of felsic-alkalic residuum. Unexsolved titanomagnetite with a limited compositional range and rare, unexsolved, Mg- and Mn-poor ilmenite occur in the groundmass of some basalts.

## Megacrysts and inclusions

Some basanites contain megacrysts of olivine and clinopyroxene whose compositions overlap those of the most mafic phenocrysts in the lavas (Fig. 2). Three types of ultramafic inclusions are also present: 1) spinel wehrlite; 2) olivine websterite; and 3) spinel websterite. They have equigranular to poikilitic textures with minimal evidence of recrystallization and deformation. Wehrlites and olivine websterites are composed of olivine (Fo<sub>85</sub> to Fo<sub>81</sub>), Cr diopside



Fig. 2. Compositions of clinopyroxenes from MMVF dacites, and andesites, basalts, and ultramafic inclusions plotted in the pyroxene quadrilateral. Filled symbols are phenocrysts and open symbols are groundmass phases

and Mg-Al chromites with olivine websterites also containing orthopyroxene (En<sub>85</sub>, 2.5 wt.% Al<sub>2</sub>O<sub>3</sub>). Spinel websterites contain diopsidic to augitic clinopyroxene, orthopyroxene (En<sub>80</sub>, 2.3 wt.% Al<sub>2</sub>O<sub>3</sub>), and chromian titanomagnetite.

## Intermediate and silicic rocks

MMVF hornblende andesites, two-pyroxene dacites, and hornblende rhyodacites are mildly porphyritic with hyalopilitic textures. Plagioclase phenocrysts are abundant in all rocks except the Round and Table Mountain andesites, and are absent in the Mormon Mountain andesites and dacites. Euhedral phenocrysts in the Hutch Mountain andesite exhibit strong reverse zoning  $(An_{39} \text{ cores} - An_{64} \text{ rims})$  and are accompanied by rare albite xenocrysts and quartz and magnetite fragments. In contrast, plagioclase phenocrysts of the Round and Table Mountain andesites are normally zoned  $(An_{63} - An_{56})$ . Rhyodacites contain two groups of phenocrysts consisting of large (up to 1 cm), oscillatory-zoned, rectangular crystals  $(An_{45} - An_{30})$  and small (<3 mm), normally-zoned, equant crystals  $(An_{33} - An_{24})$ .

Andesites and rhyodacites contain abundant large amphibole phenocrysts. Compositions range from ferroan pargasitic hornblende (andesite) to ferroan pargasite (rhyodacites). Subhedral to euhedral augites (Fig. 2) with sector and subtle concentric zoning occur in the Mormon Mountain andesites and dacites. Augites in the andesites contain about twice as much  $TiO_2$  and  $Al_2O_3$ (0.71 and 4.5 wt.%, respectively) as do augites in the dacites. Orthopyroxene phenocrysts of bronzite-to-hypersthene composition occur only in the pyroxene dacites and usually in glomerocrysts with clinopyroxene.

Groundmass are dominated by plagioclase and glass with accessory opaques, clinopyroxene, quartz, alkali feldspar, and apatite. Small laths of plagioclase range from  $An_{65-37}$  (andesites),  $An_{51-42}$  (dacites) to  $An_{40-25}$  (rhyodacaites). Light-to-dark-brown glass is only slightly devitrified and rhyolitic in composition with high  $K_2O/Na_2O$  ratios (2–6). Opaque phases are either titanomagnetite or titanomagnetite (less rich in ulvospinel) and ilmenite.

## Geochemistry

Sixty-nine samples from MMVF were selected for major element analysis and 34 were analyzed for trace elements. Sample locations

Sample	Ol. Nephelinite	Basanite		Alkali Ol. Basalt		Hawaiite		Ol. Tholeiite
	MMT-24	MMT-29	MMT-11	MMT-2	MMT-34	MMT-16	MMT-5	MMT-10
SiO <sub>2</sub>	43.50	45.40	48.92	47.77	47.73	51.86	52.64	50.81
TiO	1.80	2.34	1.63	1.74	1.40	1.30	1.31	1.46
Al <sub>2</sub> O <sub>2</sub>	12.65	14.88	17.53	15.37	17.00	15.43	18.62	16.47
FeO	9.89	10.76	9.26	9.58	11.47	8.39	8.72	9.61
MnO	nd	nd	0.15	0.15	nd	0.14	0.13	nd
MgO	14.57	10.59	5.75	9.99	7.33	8.63	4.39	7.13
CaO	12.13	10.71	10.90	10.48	10.48	9.18	8.32	10.48
Na <sub>2</sub> O	3.35	3.93	3.93	3.50	3.29	3.24	3.99	2.99
K <sub>2</sub> O	1.23	0.65	1.45	0.77	0.98	1.36	1.36	0.71
$P_2O_5$	0.89	0.73	0.48	0.63	0.32	0.47	0.52	0.33
Mg # <sup>b</sup>	75.6	67.4	56.6	68.6	57.3	68.4	51.4	60.9
CaO/Al <sub>2</sub> O <sub>3</sub>	0.96	0.72	0.62	0.68	0.62	0.60	0.45	0.63
CIPW normativ	e mineralogy							
Diop	31.26	22.45	20.79	19.68	17.52	15.57	7.70	16.83
Hyp					• •	9.64	11.71	12.62
Oliv	23.29	19.51	10.90	19.12	17.32	10.42	4.34	5.97
Or	4.54	3.85	8.55	4.54	5.80	8.02	8.03	4.18
Ab		14.59	20.96	21.46	20.98	27.45	33.73	25.32
An	15.86	21.02	25.90	23.93	28.74	23.53	28.91	29.42
Ne	15.34	10.12	6.68	4.44	3.69			
Leu	2.13		• •				••	••
Mt	2.39	2.60	2.24	2.32	2.77	2.03	2.11	2.32
Ilm	3.41	4.44	3.09	3.30	2.66	2.46	2.49	2.77
Ap	1.94	1.60	1.05	1.37	0.70	1.02	1.14	0.73
100 An								
$\overline{An + Ab}$	100.0	62.9	57.6	55.1	60.6	46.2	46.1	53.7

Table 1. Major element geochemistry and CIPW norms of representative MMVF basalts<sup>a</sup>

<sup>a</sup> Normalized to 100.0% volatile free. Norms calculated with FeO = 0.85  $\Sigma$  FeO

<sup>b</sup> Mg # = 100 Mg / Mg + 0.85  $\Sigma$  FeO



Fig. 3. Classification of representative MMVF basalts by normative hypersthene-nepheline vs. normative AN (100 An/An + Ab) (adapted from Best and Brimhall 1974). *Filled squares*-basanites, *open squares*-alkali ol. basalts, *filled circles*-hawaiites, *open circles*ol. tholeiites



Fig. 4. Classification of MMVF intermediate and silicic rocks by  $K_2O$  vs. SiO<sub>2</sub> (Peccerillo and Taylor 1976). *Circles*-Mormon Mountain complex samples, *squares*-other MMVF samples. *Filled symbols* are analyses given in Table 3

Table 2. Trace element geochemistry of representative MMVF basalts

Sample	Ol. Nephelinite	Basanite		Alkali Ol.	Alkali Ol. Basalt		Hawaiite	
	MMT-24	MMT-29	MMT-11	MMT-2	MMT-34	MMT-16	MMT-5	MMT-10
Rb	24	22	15	11	7	21	16	6
Ba	1,135	905	1,475	1.120	470	1.150	1.395	640
Sr	1,065	1,040	1,015	1,040	540	960	1.120	565
Pb	5	8	12	9	5	14	18	6
La	92	51	45	41	19	42	52	26
Y	24	21	20	19	20	18	19	20
Zr	240	218	144	171	120	146	165	111
Nb	83	57	49	48	22	53	53	23
Ga	14	17	17	17	19	16	21	18
Cr	714	410	77	404	167	495	6	238
Ni	366	216	46	190	108	249	17	87
V	225	229	220	212	206	180	177	205
Sc	29	24	23	22	27	23	19	28
Cu	70	52	57	58	93	65	39	98
Zn	83	93	73	83	88	72	95	90

are given in Fig. 1. Major element analyses are each averages of 30 individual electron microprobe analyses of glasses from duplicate fusions on Mo strips in an Ar-pressurized atmosphere (Brown 1977; Nicholls 1974). Fused beads of BCR-1 and AGV-1 were used as internal standards. Na<sub>2</sub>O, K<sub>2</sub>O, and P<sub>2</sub>O<sub>5</sub> were determined by atomic absorption for all 34 samples selected for trace element analysis. Selected samples were also analyzed in triplicate by atomic absorption to verify the fused bead analyses. Results obtained by these other procedures are in good agreement with the fused bead data. Abundances of Ba, Rb, Sr, Pb, La, Y, Zr, Nb, Ga, Cr, Ni, Sc, V, Cu, and Zn were determined by x-ray fluorescence spectrometry using the methods of Norrish and Chappell (1977). Only selected analyses are listed in the accompanying tables; additional analyses are available from the authors. Rare earth elements (REE), U, Th, Cs, Hf, Sn, and Mo were analyzed by spark source mass spectrography with the methods of Taylor and Gorton (1977). Sr-isotope ratios were measured at the Australian National University using Re triple filaments and normalized to <sup>86</sup>Sr/<sup>88</sup>Sr=0.1194 (McCulloch and Perfit 1981). Replicate measurements of NBS 987 yielded an average  ${}^{87}$ Sr/ ${}^{86}$ Sr of 0.71033+6.

## **Basalts**

MMVF basalts are varied and include olivine nephelinite, basanite, alkali olivine basalt, hawaiite, and olivine tholeiite (classification of Best and Brimhall 1974) (Table 1; Fig. 3). Mg numbers  $[100 \text{ Mg}/(\text{Mg}+\text{Fe}^{2+}) \text{ with } \text{Fe}^{2+}=0.85 \text{ total}$ Fe (Nicholls and Whitford 1976)] range between 67 and 54. Abundances of large-ion lithophile (LIL) elements (also  $P_2O_5$  and  $K_2O$ ) are similar among basanites, alkali olivine basalts, and hawaiites, but vary within each group by a factor of 3 (Table 2). Incompatible elements are lowest in the olivine tholeiites and highest in the olivine nephelinite. Contents of the compatible trace elements (Ni, Cr, Sc, and V) are similar for the various basalt groups. Ni and Cr contents exhibit a slight positive correlation with Mg number whereas Sc and V do not vary systematically with Mg number. High Ni (366 ppm) and Cr (714 ppm) content of the olivine nephelinite (MMT-24) (Mg  $\pm 75$ ) may be

Sample	Andesite				Dacite			Rhyodacite	
	MMT-6	MMT-21	MMT-22	MMT-25	MMT-7	MMT-9	MMT-13	MMT-3	MMT-18
SiO <sub>2</sub>	60.05	59.92	60.70	57.90	62.06	66.04	68.03	69.97	68.80
TiO <sub>2</sub>	1.00	0.68	0.74	0.86	0.70	0.33	0.53	0.26	0.41
Al <sub>2</sub> Õ <sub>3</sub>	17.33	19.07	18.00	20.05	17.14	16.02	16.72	14.66	16.14
FeO	5.60	5.04	4.91	5,44	5.41	4.30	3.10	3.31	2.73
MnO	nd	nd	nd	nd	0.10	0.05`	nd	nd	nd
MgO	3.69	2.09	2.95	1.80	2.63	2.02	1.12	1.33	1.60
CaO	5.88	6.23	5.93	6.33	5.42	3.98	3.69	3.16	3.39
Na <sub>2</sub> O	4.38	4.84	4.61	5.07	4.33	4.98	4.47	4.81	4.34
K <sub>2</sub> O	1.79	1.75	1.79	1.98	1.94	2.09	2.17	2.35	2.44
$P_2O_5$	0.27	0.38	0.35	0.56	0.29	0.19	0.17	0.14	0.15
LOI	0.32	0.76	0.53	0.07	0.76	0.09	0.58	0.85	1.80
Mg # <sup>b</sup>	58.1	46.6	55.8	37.2	50.6	49.7	43.2	45.8	55.2
K <sub>2</sub> O/Na <sub>2</sub> O	0.41	0.36	0.39	0.39	0.45	0.42	0.49	0.49	0.56
CIPW norma	tive mineral	ogy							
Diop	4.28	2.92	3.49	1.68	2.94	2.86		2.85	
Hyp	13.42	9.79	11.41	9.95	11.81	9.27	6.31	6.17	7.07
Qtz	8.58	7.61	9.07	3.58	12.39	16.19	22.98	23.35	23.28
Ōr	10.56	10.34	10.59	11.71	11.46	12.35	12.84	13.90	14.24
Ab	37.04	40.92	39.04	42.88	36.61	42.14	37.80	40.73	36.30
An	22.36	25.16	23.13	26.11	21.61	15.19	17.31	11.45	15.74
Mt	1.35	1.22	1.19	1.31	1.31	1.04	0.75	0.80	0.65
Ilm	1.91	1.30	1.41	1.64	1.33	0.63	1.01	0.49	0.82
Ap	0.59	0.83	0.77	1.23	0.63	0.42	0.37	0.31	0.33
Cor							0.68		0.53

Table 3. Major element geochemistry and CIPW normative mineralogies of representative MMVF intermediate and silicic rocks<sup>a</sup>

<sup>a</sup> Normalized to 100.0%, volatile free. CIPW norms calculated using FeO = 0.85 FeO

 $^{\rm b}~$  Mg #  $\,=\,100$  Mg / Mg +  $\,0.8\,\varSigma$  FeO

Sample	Andesite				Dacite			Rhyodacite	
	MMT-6	MMT-21	MMT-22	MMT-25	MMT-7	MMT-9	MMT-13	MMT-3	MMT-18
Rb	16	14	16	18	19	21	23	21	16
Ba	1,310	1,300	1,520	2,200	1,440	1,445	1,570	1,525	1,565
Sr	1,145	1,530	1,520	1,390	1,220	1,110	1,065	985	715
Pb	17	17	18	23	18	21	22	20	20
La	42	47	51	70	39	44	29	23	21
Y	17	13	11	17	12	11	7	8	4
Zr	111	145	146	175	124	122	121	107	126
Nb	33	28	33	69	29	38	19	17	12
Ga	18	20	20	20	18	18	18	17	15
Cr	34	3	21	21	26	27	23	17	17
Ni	34	8	23	4	30	26	21	14	14
V	121	69	74	83	102	64	49	41	44
Sc	15	10	10	8	13	9	8	6	6
Cu	36	14	12	12	22	26	24	14	12
Zn	66	74	73	75	66	59	65	46	48

Table 4. Trace element geochemistry of representative MMVF intermediate and silicic rocks

the result of olivine ( $\pm$  spinel) accumulation. Less than 50 ppm Ni is present in the basalts (MMT-5, MMT-20) with the lowest Mg numbers, which are also distinctive because of their high Al<sub>2</sub>O<sub>3</sub> contents (>18 wt.%).

## Intermediate and silicic rocks

MMVF andesites, dacites, and rhyodacites (Table 3) are calcalkalic as defined by normative plagioclase versus

 $Al_2O_3$  (Irvine and Baragar 1971),  $K_2O$  versus  $SiO_2$  (Peccerillo and Taylor 1976) (Fig. 4), and the alkali-lime index (Peacock 1931). The Hutch Mountain andesite (MMT-25) is more alkalic, contains more  $Al_2O_3$  (20 wt.%), and has a lower Mg number (37) than any other MMVF andesite. The MMVF suite as a whole is characterized by high Ba (1,300–2,200 ppm) and Sr (715–1,530 ppm), but low Rb (14–23 ppm) and Y (4–17 ppm) (Table 4). Concentrations of Ba, Rb, Pb, and Zr increase and La, Y, Nb, and Sr



Fig. 5. Incompatible trace elements vs.  $SiO_2$  for MMVF intermediate and silicic rocks. Symbols are as in Fig. 4

 
 Table 5. Additional trace elements of andesite, dacite, and rhyodacite from Mormon Mountain volcanic complex<sup>a</sup>

	Andesite	Dacite	Rhyodacite
	MMT-6	MMT-9	MMT-3
La	42.0	44.0	23.0
Ce	73.1	82.2	44.4
Pr	7.7	7.0	5.1
Nd	27.3	24.1	18.7
Sm	4.3	2.6	2.7
Eu	1.20	0.76	0.88
Gd	3.0	1.4	1.8
ТЬ	0.49	0.23	0.21
Но	0.38	0.19	0.22
Er	0.93	0.37	0.48
Yb	0.75	0.20	0.32
Ba <sup>b</sup>	1,310	1,445	1,525
Cs	0.43	0.79	0.58
U	1.47	1.56	0.95
Th	5.0	5.6	2.7
Zr <sup>b</sup>	111	122	107
Hf	2.2	1.9	2.3
Sn	0.85	0.70	0.70
Мо	1.3	1.7	1.3
SiO <sub>2</sub> (wt.%)	60.05	66.04	69.97

<sup>a</sup> All abundances in ppm

<sup>b</sup> XRF values

decrease with increasing SiO<sub>2</sub> contents (Fig. 5). The abundances of Ba, Pb, La, Y and Nb are unusually high in the Hutch Mountain andesite. Chondrite – normalized REE abundances of Mormon Mountain andesite (MMT-6) and rhyodacite (MMT-3) are LREE-enriched and strongly fractionated (La/Yb-= 55 and 73) (Table 5; Fig. 6). The rhyodacite contains less total REE than the andesite. REE's of the Mormon Mountain dacite (MMT-9) are more fractionated (La/Yb=220) and cross both the andesite and the rhyodacite REE patterns (Fig. 6). Small positive Eu anomalies are present in the dacite and rhyodacite, but not in the andesite. Abundances of the compatible trace elements (Ni, Cr, Sc, V, Cu, and Zn) decline from andesites to rhyodacites.



Fig. 6. Chondrite-normalized REE diagram for andesite, dacite, and rhyodacite from the Mormon Mountain complex. Reference andesite from Bagana is typical of island arc andesites and is shown for comparison

Table 6. <sup>87</sup>Sr/<sup>86</sup>Sr isotopic ratios of samples from MMVF

Sample	Rock type	<sup>87</sup> Sr/ <sup>86</sup> Sr	Sr (ppm)	Rb (ppm)
MMT-6	hb. andesite	$0.70332\pm 2$	1145	16
MMT-7	pyx. dacite	$0.70400\pm8$	1220	19
MMT-3	rhyodacite	$0.70363 \pm 5$	985	21
MMT-22	hb. andesite	0.70378 + 4	1520	16
MMT-5	hawaiite	$0.70367 \pm 7$	1120	16
MMT-24	ol. nephelinite	0.70361 + 8	1065	24
SW-6	pyx. granulite	$0.70372 \pm 1$	1013	0.5
NBS 987	standard	0.71033	••	

# Strontium isotopes

 $^{87}$ Sr/ $^{86}$ Sr isotopic ratios of two basalts and a nephelinite are within the range of 0.7036 to 0.7037 (Table 6). The andesite, dacite, and rhyodacite suite ranges from 0.70332 to 0.70400. A pyroxene granulite xenolith from nearby Williams, Arizona is characterized by a low Rb/Sr ratio (5 × 10<sup>-4</sup>) and a Sr isotopic ratio of 0.70372.

# Discussion

## **Basalts** petrogenesis

Our working hypothesis to account for the range of primitive basalt major and trace element geochemistry, despite their near identity in terms of <sup>87</sup>Sr/<sup>86</sup>Sr ratio, involves variable degrees of partial melting of a common upper mantle source as the primary process. Secondary processes, such as fractional crystallization and assimilation of crustal rock types, have operated to varying extent on this range of primitive basalt types. These processes further diversify the geochemistry of MMVF basalts.

Primitive basaltic magmas that are little modified by extensive fractional crystallization or crustal contamination should have high Mg numbers (68 to 75) and high contents of compatible trace elements (e.g. Frey et al. 1978). MMVF alkali olivine basalts, basanites, and nephelinite with these characteristics contain abundant olivine phenocrysts, suggesting that they have fractionated olivine. Correction for this fractionation to a uniform Mg number of 71 (appropriate for a melt in equilibrium with a mantle of Mg number 89) requires adjustments in trace element abundances. For incompatible elements this is simply a dilution effect, however, adjustment of Ni, Cr, Sc, and V abundances is complicated by the marked affinities of olivine and spinel for these elements (Frey et al. 1978). No attempt was made to model the abundances of these siderophile elements in the mantle source.

Different pressures, temperatures, and degrees of melting or source heterogeneities all affect melt composition. Assuming a uniform source, pyrolite (Ringwood 1966), the effects of differing degrees of partial melting can be evaluated (eg. Frey et al. 1978) for MMVF primitive basalts. The degree of partial melting may be defined by the  $P_2O_5$ content of the primitive basalt and the pyrolite source (0.06 wt.%). TiO<sub>2</sub> and CaO contents of the basalt provide qualitative estimates of the degree of melting as both decrease as the percentage of melting increases. The CaO/Na<sub>2</sub>O ratio increases as the degree of melting increases due to the dilution of jadeitic components in earlier formed melts by diopsidic components.

Although quantification is highly model dependent, the olivine nephelinite, with its low contents of SiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub> and high contents of MgO, CaO, TiO<sub>2</sub>, and P<sub>2</sub>O<sub>5</sub>, should have resulted from small degrees of partial melting. The CaO/Na<sub>2</sub>O ratio is high relative to the estimated degree of melting, but may be explained by bonding of Ca with CO<sub>3</sub> in the melt (Frey et al. 1978). Experimental studies on olivine nephelinite indicate that the associated volatile phase must be CO2-rich relative to H2O during melting and consequently, the magmas must be enriched in CO<sub>3</sub> (Brey 1978). Primitive basanites have higher SiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub> contents and lower MgO, CaO, TiO<sub>2</sub>, and P<sub>2</sub>O<sub>5</sub> contents than the olivine nephelinite, suggesting greater degrees of partial melting. Primitive alkali olivine basalts result from even greater degrees of partial melting than basanities. Given the model adopted, percentages of partial melting calculated vary from 7% for the olivine nephelinite to about 20% for the alkali olivine basalts. Although the absolute numbers may be substantially in error (c.f. McKenzie 1984), the overall sense of low to high percentages of melting in the order suggested seems probable.

Abundances of incompatible trace elements of primitive MMVF basalts correlate with the estimated degree of partial melting, decreasing as the amount of melting increases. Results of non-modal batch partial melting calculations suggest that the source of MMVF basalts was enriched in various incompatible elements relative to a model primitive mantle (Wood 1979). The order of this enrichment, from greatest to least, is Nb, Ba, La, Sr, K, Rb, Pb, Zr, and Y (Fig. 7). The causes of these enrichments are highly speculative but is note-worthy that strong enrichments of these elements are also present in the MARID suite of ultramafic nodules in kimberlites (Dawson and Smith 1977). Metasomatism of the mantle by  $H_2O + CO_2$  fluids enriched in incompatible elements (Best 1975; Boettcher et al. 1979), and veining of the mantle by highly alkalic melts (Wood 1979) have been suggested to account for these types of mantle enrichments.



Fig. 7. Calculated enrichment (relative to primitive mantle (Wood 1979)] in the source of the ol. nephelinite and a basanite from MMVF



Fig. 8.  $Al_2O_3$  vs. MgO for all MMVF basalts. *Arrows* indicate fractionation trends of liquids crystallizing clinopyroxene, olivine, and plagioclase. The MMVF trend is best explained by a combination of olivine and clinopyroxene fractionation and absence of plagioclase fractionation

## Fractional crystallization and crustal contamination

Fractional crystallization of the different MMVF primitive basalts accounts for the variety of the MMVF basaltic suite. Decreases in CaO and increases in Al<sub>2</sub>O<sub>3</sub> with decreasing MgO indicate strong olivine and clinopyroxene control during fractionation (Fig. 8). Plagioclase fractionation is relatively unimportant, and distinctly unlike that of many ocean floor and island arc basaltic suites. Quantitative models of fractional crystallization based on major element contents and petrologic characteristics are derived from least squares mixing calculations (Wright and Doherty 1970). These models suggest that moderate amount (8% to 32%) of olivine, clinopyroxene, and spinel have been fractionated from primitive MMVF basalts. The Cr, V, and Sc contents are controlled by spinel and clinopyroxene fractionation. Extremely high Kd's of Ni and Cr for olivine and spinel, respectively, account for the observed three of fourfold drop in the concentrations of these elements (Sato 1977) prior to clinopyroxene fractionation. Decreases in Sc content in



Fig. 9. Chondrite-normalized Ba/La ratios vs.  $P_2O_5$  for all MMVF basalts. The dark band represents the expected range of Ba/La ratios of partial melts from a peridotitic mantle source (*large circle*). Estimates of possible lower crustal materials which might contaminate the basalts are shown by arrows. Symbols are in Fig. 3

the more evolved basalts are compatible with clinopyroxene fractionation (Irving 1978).

Rayleigh fractionation calculations (Minster et al. 1977) using constraints of least squares modeling do not successfully reproduce the observed characteristics for some incompatible trace elements of MMVF basalts. These results, which change only slightly with the choice of partition coefficients [from Arth (1976); Irving (1978)] require the derivative magmas to be less enriched in the highly incompatible elements (Rb, Ba, Sr, K, Nb, La) than they actually are. This disparity is consistent with the complex, but geologically reasonable open-system magma chamber process described by O'Hara and Mathews (1981). However, the limited amount of basaltic volcanism and the strongly silicaundersaturated character of it in the MMVF seems unlikely to be products of a near steady-state magma chamber with periodic magma replenishments and evacuations.

The differences between calculated and actual abundances of some incompatible trace elements can also be explained by a combination of fractional crystallization and assimilation of incompatible element enriched crust (De-Paolo 1981). Irregularities in the trends of incompatible trace elements of some primitive basalts with respect to degree of melting support this general concept. MMVF basalts were erupted through about 40 km of continental crust and this passage may have resulted in selective contamination (Watson 1982). If basalts are contaminated by melts derived from small amounts of crustal anatexis, P should be retained in the crustal residue in apatite due to its low solubility in felsic melts (Watson and Capobianco 1981). The presence of residual apatite in the source would also diminish the concentrations of REE in the melt. On the other hand, Ba, K and Pb are concentrated in felsic melts being derived from the partial melting of alkali feldspar (Thompson et al. 1983). Watson (1982) has shown that assimilation of a feldspar melt in a basalt produces enrichments of various alkali elements over Si, Al, and the REE. These observations suggest that assimilation of crustal materials by basaltic magmas should result in selective enrichment of Ba, K, and Pb over P and the REE. Weaver and Tarney (1983) and Thompson et al. (1983) present several spidergrams of possible crustal contaminants that are consistent with these generalizations. In all cases, these contaminants are characterized by positive Ba/La ratios (chondritenormalized) and low abundances of P. The use of these parameters, Ba/La or Pb/La versus  $P_2O_5$  may help assess the nature of crustal contamination in MMVF basalts. These ratios should decrease slightly as the degree of partial melting increases if the magmas are from a homogenous source. MMVF basalts exhibit trends opposite to those expected from only partial melting (Fig. 9) and although this can be due in part to clinopyroxene fractional crystallization, it is also consistent with crustal contamination.

## Basalt and andesite – co-genetic?

Successful least-squares mixing models for several basaltandesite parent-daughter pairs are possible using either the observed basaltic or andesitic mineralogies as the fractionating assemblage. However, in using these models simple Rayleigh fractionation calculations are inconsistent for most of the incompatible trace elements. Almost all calculations predict excessive abundances of Rb, Pb, K, Ba, Zr, Nb, La, and Y in the andesites. Calculated La/Y ratios, which qualitatively model (LREE/HREE ratios, are variable and do not compare favorably with observed La/Y ratios. Calculated Sr contents are more enriched than observed values. Sr isotopic data argue most convincingly against a simple co-genetic relationship between MMVF andesites and basalts as <sup>87</sup>Sr/<sup>86</sup>Sr isotopic ratios of the basalts fall within a narrow range, 0.7036 to 0.7037, with the ratios of the andesites being either higher or lower.

An exception to this general conclusion is the Hutch Mountain andesite (MMT-25). This andesite, which is geochemically distinctive from other MMVF andesites, probably formed by fractional crystallization of a basaltic parent. A model using a fairly evolved hawaiite (Mg number = 52) as the parent produces a close match to the observed major and trace element contents of the andesite. The occurrence of this andesite as a small plug within a hawaiite cinder cone supports this interpretation.

Combined wall-rock assimilation and fractional crystallization significantly enriches the highly incompatible elements in a magma over that expected from simple fractional crystallization (DePaolo 1981). Unlike MMVF basalt however, MMVF andesites are depleted in incompatible elements relative to the predictions of fractional crystallization models; combination of fractional crystallization with assimilation of incompatible element-enriched crustal material should accentuate this disparity. Thus, fractional crystallization combined with assimilation is probably an inappropriate mechanism to explain the origin of MMVF andesites.

#### Origin of MMVF calcalkalic rocks – crustal anatexis

An alternative petrogenetic hypothesis assumes andesite to be a primary magma. Some experimental petrologists have suggested that andesite could be generated by partial melting of amphibolite (Boettcher 1973), hydrous quartz eclogite (Ringwood 1974) or hydrous peridotite (Mysen and Boettcher 1975), but subsequent studies have concluded that calcalkalic low-Mg andesite can not be in equilibrium with a hydrous peridotite (Green 1976) or hydrous eclogite



Fig. 10. AB-OR-QZ ternary projection of MMVF andesites, dacites, and rhyodacites. Similarities between the trend of this suite and the trends of partial melting of amphibolite (tholeiitic and alkalic composition – Helz 1976) suggests that the MMVF suite may result from different degrees of amphibolite fusion

 Table 7. Input parameters for non-modal, sequential batch partial melting calculations for Mormon Mountain andesite

Proportions of minerals in source	Amphibole	Plagioclase	Apatite	Magnetite
Initial	0.500	0.470	0.015	0.015
First melt	0.300	0.650	0.035	0.015
Second melt	0.300	0.650	0.050	0.000
Third melt	0.800	0.200	0.000	0.000

Apatite exhausted at 42.8% melting. End of first melting interval. Trace elements for andesite source calculated at 45.0% melting

at high pressures (Stern and Wyllie 1978; Gust 1982). Experiments on amphibolites (Holloway and Burnham 1972; Helz 1973/76) indicate that moderate degrees of melting (30%-50%) produce liquids similar to MMVF andesites, dacites, and rhyodacites (Fig. 10). These similarities suggest that amphibolite anatexis is of least a tenable hypothesis for the origin of the MMVF calcalkalic suite.

Trace element characteristics of a source for the Mormon Mountain andesite were calculated using an amphibolite anatexis model. Partial melting equations depend upon many poorly constrained variables including variations in source paragenesis, mode of melting, and trace element partitioning with respect to melt composition (Hanson 1980). Calculation of an amphibolite source uses input parameters loosely constrained by the experimental data of Helz (1976) (Table 7). The results of these calculations are shown in Fig. 11 where the source of the Mormon Mountain andesite is bracketed by compositions calculated using low-Si andesite and dacite partition coefficients (Hanson 1978). A generalized composition of the source of the Mormon Mountain andesite is moderately LREE-enriched and HREE-depleted. The contents of Rb, Sr, Ba, Y, Zr, Nb, and the REE's are not typical of model estimates for either upper



Fig. 11. Calculated amphibolite source(s) of andesite and rhyodacite of Mormon Mountain. Three sets of partition coefficients used in the calculation are indicated in the figure. The amount of overlap between the most likely coefficient sets represents the best approximation of the source

or lower crust (eg. Taylor and McLennan 1981). However, trace element analyses of crustal xenoliths indicate that the crust is much more complex than any of these generalized models and considerable variability in trace element composition is expected. The relatively low <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the MMVF andesites indicate that their source must have low Sr isotopic ratios resulting from a long-term depletion in Rb. Low abundances of Rb in these lavas are also consistent with a Rb-poor source.

Different source mineralogies (pyroxene or garnet amphibolite) than the one used in the partial melting calculations show that addition of clinopyroxene at the expense of amphibole and plagioclase requires a more incompatible element-depleted source. Addition of garnet, replacing plagioclase, lessens the HREE-depletion of the source.

Mormon Mountain dacites and the MMVF rhyodacites could have been produced by smaller degrees of partial melting than those envisioned to produce andesitic magmas from amphibolite. Partial melting calculations for the trace elements suggest that andesite and rhyodacite could have been derived from a source with similar trace element characteristics (Fig. 11). Anomalies in Zr and the heavy REE abundances between the two sources may represent trace amounts of residual zircon in the rhyodacite source (Watson 1979). The petrogenesis of the dacite also may involve garnet or zircon as a residual phase in its source as suggested by its depleted HREE signature.

## Fractional crystallization in the calcalkalic suite

Derivation of the dacites and rhyodacite of Mormon Mountain by fractional crystallization of the andesite is possible only if the proposed parent-daughter relationship is compatible with geological, temporal, petrologic, and geochemical characteristics of the suite. Different phenocryst assemblages of the andesite and dacites (wet versus dry), crossing REE patterns, and approximate constancy in contents of highly incompatible elements are inconsistent with a fractional crystallization origin for the dacites. On the other hand, these types of tests show that the rhyodacite could have been produced from andesite by fractional crys-



Fig. 12. Results from Rayleigh fractionation calculations for andesite (MMT-6) to rhyodacite (MMT-3) presented as the ratio of calculated abundances over observed abundances. The large compositional change between the parent-daughter pair is reflected in the use of different sets of partition coefficients. The closest approximation to the observed abundances in the rhyodacite (ratio = 1) is achieved by using dacite coefficients

tallization. Least squares calculations suggest that about 50% crystallization and removal of amphibole, plagioclase, apatite, and magnetite is required. Coarse-grained inclusions, dominated by amphibole and plagioclase, are found in the dacites, and may be products of this fractionation. Rayleigh fractionation calculations for incompatible trace elements neither confirm nor deny this hypothesis due to the extreme dependency of element partitioning between amphibole and melt on magma composition. The REE's (except Ce and Yb) are best modeled using dacitic coefficients, but Zr, Nb, Y, and Sr abundances are inconsistent with the data (abundances and Kd's) at hand (Fig. 12). Enrichments in Rb, Ba, U, Th, and Ce are predicted using either set of partition coefficients and contrast strongly with the observed depletions. In general, therefore, fractional crystallization of parental basalt magmas seems less likely as a genetic process in the generation of evolved melts of MMVF than melting of crustal sources.

## Conclusions

The petrogenesis of MMVF is complicated and consists of two distinct aspects: the origin of the basalts, and the origin of the calcalkalic rocks. We have shown that these different suites are not likely co-genetic. Except in one case, MMVF andesites cannot be derived from MMVF alkalic basalts by fractional crystallization with or without crustal assimilation. Evidence for magma-mixing is not apparent. Therefore, the andesites are interpreted to be products of crustal anatexis. The only direct connection between basalts and andesites would seem to be the possible thermal contribution from hot basaltic magmas to the anatexis of the sources of the andesites.

The conclusions of this study are:

1. MMVF basalts are derived by varying degrees of partial melting of an incompatible element-enriched, metasomatized mantle. Fractional crystallization of olivine and clinopyroxene increases the compositional spectrum. Contamination by continental crust is subtle, and apparently restricted to highly incompatible elements.

2. Calcalkalic andesites, dacites, and rhyodacites may be generated by partial melting of a wet basaltic source (amphibolite) in the lower crust.

3. Rhyodacites may be produced by fractional crystallization of andesitic parents.

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