# EXPERIMENTAL CONTROL OF FLUORINE REACTIONS IN HYDROTHERMAL SYSTEMS

J. L. Munoz, Department of Geological Sciences, University of Colorado, Boulder, Colorado, 80302 and H. P. Eugster, Department of Earth and Planetary Sciences, Johns Hopkins University, Baltimore, Maryland, 21218.

#### Abstract

The solid polyphase buffer technique has been adapted to permit quantitative study of synthetic fluorine-bearing hydrothermal systems. This method involves three nested capsules; the innermost capsule contains the charge which is open to the fluorine buffer assemblage in the sealed second capsule. The outermost capsule contains an oxygen buffer; the hydrogen fugacity  $(f_{\rm H_2})$  in equilibrium with the fluorine buffer assemblage must be externally fixed. The buffer reactions which have been studied experimentally are:

wollastonite+2HF; fluorite+quartz+H<sub>2</sub>O (WFQ) anorthite+2HF; fluorite+sillimanite+quartz+H<sub>2</sub>O (AFSQ) calcite+2HF; fluorite+graphite+H<sub>2</sub>O+O<sub>2</sub> (CFG)

Fugacities and compositions of the gas phase in equilibrium with the given buffer phases have been calculated from thermodynamic data and are presented as a function of temperature, total pressure, and  $f_{\rm H_2}$ . In all cases, the concentration of HF in the gas (in wt%) never exceeds 5 percent, and is more commonly one to eight orders of magnitude smaller than that. For the WFQ and AFSQ buffers, the remainder of the gas is composed of H<sub>2</sub>O and H<sub>2</sub>. When graphite is present (CFG buffer), the equilibrium bulk composition of the gas phase is primarily a mixture of CO<sub>2</sub>, H<sub>2</sub>O, CO, CH<sub>4</sub>, and H<sub>2</sub> (the exact composition depending mostly on temperature and  $f_{\rm H_2}$ ), with very minor amounts of HF.

Tests of these buffers using synthetic (F, OH) phlogopite as a charge indicate that equilibrium between buffer and charge requires less than two weeks at 775°C and 6–10 weeks at 550°C. The compositions of phlogopites in equilibrium with these 3 fluorine buffers range from F-phlogopite49 OH-phlogopite51 (mole %, CFG fluorine buffer with  $f_{\rm H_2}$  specified by the nickel-nickel oxide buffer, 550°C) to F-phlogopite30 OH phlogopite2 (CFG fluorine buffer with  $f_{\rm H_2}$  specified by the hematite-magnetite buffer, 700°C). The 700° data are most complete and demonstrate that phlogopite is extremely effective in removing fluorine from the gas phase. Extrapolating from this calibration, it appears that the most fluorine-rich natural phlogopite (75 mole % F-phlogopite) coexisted with a gas phase containing less than 0.05 mole % HF. Judging from natural biotites, a common range for igneous and metamorphic fluids would be 0.001–0.005 mole % HF. As more calibrations become available, the behavior of HF in natural environments can be established quantitatively.

### Introduction

Fluorine is a common constituent of many micas and amphiboles and such accessory minerals as apatite, tourmaline, and topaz. In these minerals, fluorine freely substitutes for hydroxyl. Published data indicates that fluorine greatly enhances the thermal stability of micas and amphiboles (see, Van Valkenberg and Pike, [1952]). Hence, both the composition and the stability relations of these minerals are, in part,

dependent upon the magnitudes of the H<sub>2</sub>O and HF fugacities of their environment. In order to calibrate these effects in synthetic systems, it is necessary to develop methods which permit control of these fugacities at elevated pressures and temperatures in a predictable and quantitative manner. The most simple and direct method is to vary the bulk composition of the gas phase. This approach is unsatisfactory for fluorinebearing systems because of the high solubility of silicates in HF at low temperatures, analytical difficulties, and inadequate knowledge of the PVT (and fugacity) relations in the system O-H-F under the conditions of experimental interest. Alternatively, the composition of the gas phase and its fugacities may be buffered by appropriate crystalline phases. This solid polyphase buffer approach has been widely applied to the study of redox reactions in hydrothermal systems (e.g., Eugster and Wones, 1962), and has recently been expanded to include multicomponent gas systems (Eugster and Skippen, 1967). The composition of the equilibrium gases as a function of total P and T is calculated from the equilibrium constants for all the important molecular species present (JANAF tables, 1960) and from the free energy data for the solid buffer phases (Robie, 1962). The fluorine buffers have been developed in cooperation with Rieder (1968), who has used them successfully to determine the thermal stability of zinnwaldites.

## FLUORINE BUFFER EQUILIBRIA

Consider a gas phase of the system O-H-F. For a given P and T, this gas phase is divariant; thus, in order to define the fugacities of all gas species present, it is necessary to specify two additional intensive parameters. One possible method is to vary independently the fugacities of O<sub>2</sub> and H<sub>2</sub> using double oxygen buffers (Munoz, 1966); this method has the severe drawback that fluorine must be initially added to the gas phase (e.g., as HF). To prevent this difficulty, we have chosen to use solid fluorine-bearing buffer assemblages to control the ratio of two fugacites, while the remaining degree of freedom is removed by imposing a given hydrogen fugacity on the fluorine buffer assemblage through a Pt or Ag<sub>7</sub> Pd<sub>3</sub> membrane. Figure 1 shows the experimental arrangement. The charge is contained in a gold foil and is surrounded by the fluorine buffer assemblage. This charge-buffer system is sealed in a Pt or Ag<sub>7</sub>Pd<sub>3</sub> membrane. The hydrogen fugacity is fixed externally, either by using a Shaw bomb, a standard oxygen buffer, or the methane buffer (see Eugster and Skippen, 1967).

We have calculated fugacities for the following buffer reactions in the system O-H-F as a function of P, T, and  $f_{\rm H_2}$ :

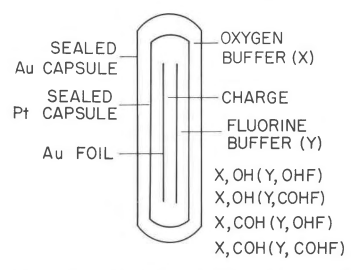


Fig. 1. Cross-sectional schematic diagram of triple capsule arrangement used for fluorine buffer experiments. The gold capsule is 1.5'' long. Notation: the fluorine buffer (Y) is enclosed by parentheses, together with the elemental composition of the gas phase. The external oxygen buffer (X) and its gas components are written to the left of the parentheses.

$$\begin{split} \text{CaSiO}_3 + 2\text{HF} &\rightleftharpoons \text{CaF}_2 + \text{SiO}_2 + \text{H}_2\text{O}^a \text{ (WFQ)} \\ \text{wollastonite} + 2\text{HF} &\rightleftharpoons \text{fluorite} + \text{quartz} + \text{H}_2\text{O} \\ \text{CaAl}_2\text{Si}_2\text{O}_8 + 2\text{HF} &\rightleftharpoons \text{CaF}_2 + \text{Al}_2\text{SiO}_5 + \text{SiO}_2 + \text{H}_2\text{O} \\ \text{anorthite} + 2\text{HF} &\rightleftharpoons \text{fluorite} + \text{sillimanite} + \text{quartz} + \text{H}_2\text{O} \text{ (AFSQ)} \end{split}$$

These calculations require as input data: values for the equilibrium constants of  $H_2O$  and HF, free energy data for the solids taking part in the buffer reaction, fugacity coefficients ( $\gamma$ ) for  $H_2O$ ,  $H_2$ , and HF, and the  $f_{H_2}$  specified by the external buffer. All of these data are functions of temperature and most of them depend upon pressure as well.

Specifically, we have

<sup>a</sup> These equations can also be written in an equivalent form:

$$CaSiO_3 + F_2 \rightleftharpoons CaF_2 + SiO_2 + 1/2 O_2$$
  
 $CaAl_2Si_2O_8 + F_2 \rightleftharpoons CaF_2 + Al_2SiO_5 + SiO_2 + 1/2 O_2$ 

in which case the equilibrium constants for the reaction are  $K_{\rm Bu}=f_{0_2}$  <sup>1/2</sup> $f_{\rm F_2}$ . We have elected to express all buffer reactions in terms of the more abundant geologic species H<sub>2</sub>O and HF. Needless to say, however, the fugacities of these species are no more or less meaningful than those of O<sub>2</sub> or F<sub>2</sub>.

$$K_{\rm H_2O} = \frac{f_{\rm H_2O}}{f_{\rm O_2}^{1/2} f_{\rm H_2}} \tag{1}$$

$$K_{\rm HF} = \frac{f_{\rm HF}}{f_{\rm H_2}^{1/2} f_{\rm F_2}^{1/2}} \tag{2}$$

- a)  $CaSiO_3 + 2HF \rightleftharpoons CaF_2 + SiO_2 + H_2O$
- b)  $CaAl_2Si_2O_8 + 2HF \rightleftharpoons CaF_2 + AlSi_2O_6 + H_2O$

$$K_{\rm B_u} = \frac{f_{\rm H_2O}}{f_{\rm HF}^2} \tag{3}$$

$$P_{\text{TOTAL}} = \frac{f_{\text{HF}}}{\gamma_{\text{HF}}} + \frac{f_{\text{H}_2\text{O}}}{\gamma_{\text{H}_2\text{O}}} + \frac{f_{\text{H}_2}}{\gamma_{\text{H}_2}} + \frac{f_{\text{F}_2}}{\gamma_{\text{F}_2}} + \frac{f_{\text{O}_2}}{\gamma_{\text{O}_2}}$$
(4)

Substituting (1), (2), and (3) into (4) gives

$$f_{\rm HF}^2 \left( \frac{K_{\rm B_u}}{\gamma_{\rm H_2O}} \right) + f_{\rm HF} \left( \frac{1}{\gamma_{\rm HF}} \right) + \left( \frac{f_{\rm H_2}}{\gamma_{\rm H_2}} - P_{\rm TOTAL} \right) = 0$$
 (5)

 $f_{\rm HF}$  is the only unknown in this quadratic equation; hence, it can be solved for any desired values of P and T.  $f_{\rm H_2}$  is controlled by a specific oxygen buffer and its magnitude has been calculated previously (see Eugster and Skippen, 1967). Knowledge of  $f_{\rm HF}$  and  $f_{\rm H_2}$ , by proper substitution in (1), (2) and (3), yields the remaining fugacities. Fugacity coefficients for  $H_2{\rm O}$  were taken from Holser (1954), for  $H_2$  from Shaw and Wones (1963) and for the remaining species from the reduced variable chart of Hougen and Watson (1946).

Note that terms involving  $f_{\rm F_2}$  and  $f_{\rm O_2}$  in the  $P_{\rm TOTAL}$  equation are neglected in (5). This can be done because the magnitudes of these fugacities are so low ( $<10^{-10}$  bars at all temperatures of interest) that they effectively do not add to the summation of partial pressures. For the same reason, equilibria involving other species in the system O-H-F (e.g., FOH, F<sub>2</sub>O) can be neglected.

The approach used for O-H-F gases can be extended to include C-O-H-F gases, provided that graphite is present. The assemblage graphite +C-O-H-F gas is also divariant for a given P and T. We have chosen the buffer reaction

$$CaCO_3 + 2HF \rightleftharpoons CaF_2 + C + H_2O + O_2$$
  
 $calcite + 2HF \rightleftharpoons fluorite + graphite + H_2O + O_2$  (CFG)

<sup>&</sup>lt;sup>1</sup> Using the more recent fugacity coefficients calculated by Anderson (1964) does not change the result significantly. A larger uncertainty, perhaps, is introduced by the necessary assumption of ideal mixing.

Table 1. Equations For Fluorine Buffers of the System C-O-H-F

$$K_{\rm W} = \frac{f_{\rm H_20}}{f_{\rm H_2} \times f_{\rm O_2}^{1/2}} \quad (1) \qquad K_{\rm HF} = \frac{f_{\rm HF}}{f_{\rm H_2}^{1/2} f_{\rm F_2}^{1/2}} \quad (2) \qquad K_{\rm CO_2} = \frac{f_{\rm CO_2}}{f_{\rm O_2}}$$

$$K_{\rm CO} = \frac{f_{\rm CO}}{f_{\rm O_2}^{1/2}} \quad (4) \qquad K_{\rm CH_4} = \frac{f_{\rm CH_4}}{f_{\rm H_2}^2} \quad (5)$$

$$CaCO_3 + 2HF \rightleftharpoons CaF_2 + C + H_2O + O_2 \quad (6)$$

$$K_{\rm BU} = \frac{f_{\rm H_20}f_{\rm O_2}}{f_{\rm HF}^2}$$

$$P_{\rm TOTAL} = \frac{f_{\rm H_20}}{\gamma_{\rm H_20}} + \frac{f_{\rm HF}}{\gamma_{\rm HF}} + \frac{f_{\rm CO_2}}{\gamma_{\rm CO_2}} + \frac{f_{\rm CO_4}}{\gamma_{\rm CO}} + \frac{f_{\rm CH_4}}{\gamma_{\rm CH_4}} + \frac{f_{\rm O_2}}{\gamma_{\rm O_2}} + \frac{f_{\rm F_2}}{\gamma_{\rm F_2}} + \frac{f_{\rm H_2}}{\gamma_{\rm H_2}} \quad (7)$$

Substituting (1)-(6) into (7) gives

$$H_{20} \left( \frac{K_{C0}}{K_{w} f_{H_{2}} \gamma_{C0}} + \frac{1}{\gamma_{H_{2}0}} \right) + f_{H_{2}0}^{3/2} \left( \frac{1}{K_{w} K_{Bu}^{1/2} f_{H_{2}} \gamma_{HF}} \right) + f_{H_{2}0}^{2} \left( \frac{K_{C0}_{2}}{K_{w}^{2} f_{H_{2}} \gamma_{C0}_{2}} \right) + \left( \frac{f_{H_{2}}}{\gamma_{H_{2}}} + \frac{K_{CH_{4}} f_{H_{2}}^{2}}{\gamma_{CH_{4}}} - P_{T} \right) = 0 \quad (8)$$

The numerical value of the second term of the above equation is almost always small enough so that it can be ignored; cancelling this term leaves a simple quadratic equation to be solved for  $f_{\rm H_20}$ . Note that the fugacity of  $\rm H_2$  must be specified; also, as was true in the O-H-F system,  $P_{\rm O_2}$  and  $P_{\rm F_2}$  are not included in the  $P_{\rm TOTAL}$  equation.

Depending on temperature and the imposed value of  $f_{\rm H_2}$ , the important species may be  ${\rm H_2O}$ , HF,  ${\rm CO_2}$ , CO, CH<sub>4</sub>, or H<sub>2</sub>. The experimental arrangement is identical to the O-H-F case, and the relevant equations are presented in Table 1.

Partial results of the fugacity calculations for the systems O-H-F and C-O-H-F appear in Table 2, and the dependence of the fugacity of HF for selected fluorine buffers, temperatures, and hydrogen fugacities is shown in Figure 2.

For O-H-F buffers, the effect of changing the external oxygen fugacity on the magnitude of the internal HF fugacity is very small, and isothermal variations in  $f_{\rm H_2O}/f_{\rm HF}$  can be obtained only by imposing very high H<sub>2</sub> fugacities on the inner capsule, so that  $P_{\rm H_2O}$  becomes much less than  $P_{\rm TOTAL}$ . Thus, the equilibrium compositions of the two buffers HM, OH (WFQ, OHF) and NNO, OH (WFQ, OHF) are essentially identical, but they have decidedly lower  $f_{\rm H_2O}/f_{\rm HF}$  ratios than the buffer QFI, OH (WFQ, OHF) (Table 2). On the other hand, C-O-H-F buffers are strongly sensitive to differences in external oxygen fugacity. This difference lies in the effect the other major components have on the magnitude of  $f_{\rm H_2O}$ .

Table 2. Calculated Gas Fugacities in Charge for Eight Typical Buffering Systems when  $P_{\rm gas}\!=\!2$  Kbar

Γemperature, °C	$-\log$	$g f_{0_2}$	$-\log f_{\mathbf{F}_2}$		$f_{\mathrm{H_2}}$	$f_{\rm H_20}$	log	fff
		1 N	HO OV	(WFO	OHF)a			
227	32.0	700	NO, OH 55.227	(11.6)	603	234.0	-3.	771
327			16 222	1	614	175 8	-2.	
427	26.2	221	46.223 39.482 34.263 30.093 26.682	1	.014	475.8 809.1 1142 1416 1650	-1.	111
527	21.8	331	39.482	3	.425	809.1	-1.	407
627	18.4	112	34,263	5	.831	1142	$\begin{array}{c} -0 \\ -0 \end{array}$	097
727	15.0	577	30.093	8	. 463	1416	-0.	115
827	13.4	140	26.682	0 1 3 5 8 11	. 33	1650	0.	356
		2. Q	FI, OH (	WFQ,	OHF)			
327	41.	2. Q. 293 321 092 026	59.829	3 2 2 2	277	$31.76 \\ 76.46$	-4.	205
427	34	321	50.270	2	891	76.46	$\frac{-2}{-1}$	834
527	20	092	43 113	2	627	145.4	-1.	817
627	25.	026	37 570	2	409	232.7	-1	042
	21.	020	43.113 37.570 33.140	2	214	332.1	0.	
727	21.	–	33.140	2 2 2	050	436.3		067
827	19.	110	29.517		050	430.3	0,	.007
		3. NI	NO, OH	(AFSQ,	OHF)	222 0	1	675
327	32.	088	51.034	(AF 5Q, 0 1 3 5 8 11	603	233.9	-1	0/3
427	26.	227	42.440	1	.614	475.7 808.7 1139 1407	-0	540
527	21.	832 414 682	36.021	3	.425	808.7	0.	287
627	18	414	31 055	5	. 831	1139	0.	908
727	15	682	27 000	8	463	1407	1	387
	13.	450	23.834	11	23	1627	1	780
827	13		23.034	11	.00	808.7 1139 1407 1627		.,,,,
		4. Q1	FI, OH (	AFSQ,	OHF)	21 76	2	100
327	41.	293	55.636	3	277	31.76		100
427	34.	321	46.487	2	2890	76.44 145.2 231.7	-0	943
527	29.	093	39.652	2	2627	145.2	-0	
627	25.	030	34.362	2	409	231.7	0	. 562
727	21	782	30 140	2	214	328.1	1	.071
827	19.	030 782 135	26.675	2	OHF) 277 890 8627 4409 214	424.2	1	.488
Temperature, °C							$f_{\rm C0}$	f <sub>0H<sub>4</sub></sub>
		5. H	M, OH (	CFG. C	COHF)			
327	30.971	51 885	0.0018	2.555	3.361	3363	0.0084	0.000
427	26.040		0.0045	1 650	2.792	3507	1.012	0.000
527	22.351	38.765	0.0091	1 186	2.372	3530	6 419	0.000
	10 476	30.703	0.0091	0.060	2.051	3595	27.01 82.78	< 10
627	19.476					3488	07 70	< 10
727	17 . 193 15 . 331	30.907	0.0210		1.824	3400	04.10	< 10
827	15.331	28.065	0.0274	0.452	1.643	3351	204.5	< 10
		6. NI	NO, OH 0.603	(CFG,	COHF)			
327	32.154	53.660	0.603	216.9	2.988	220.4	0.0215	45.07
427	26,552	45.150	1.614	327.3	1.901	1078	0.5610	27.93
527	22.589	39.122	3.425	338 1	1.264	2041	4.881	19.25
	19.605	34.575	5.831	338.1 289.2	0.852			
627	19.005	34 3/3	0.462	225.9	0.002	2912	75 64	7 03
727 827	17.271	31.025	8.463 11.334	176.0	0.580 $0.375$	2912	23.28 75.64 192.5	5.23
Temperature, °C	$-\log j_{0_2}$	$-\log f_{\rm F}$	2 /H2	∫H <sub>2</sub> O	-log/HF	JC02	fco	JUH4
				/OTC	COTTE			
		7. WM	IC, COH	(CFG,	COHF	2 -40	4 -40-4	ECO
327	41.101	7. WM 67.080	IC, COH 6.77	0.0	819 9.17	3 < 10	<10 <sup>-4</sup>	
	41.101 33.391	7. WM 67.080 55.409	6.77	0.08	819 9.17 00 6.46	3 - 0.000	0.0002	520%
327 427	33, 391	67.080 55.409 46.613	6.77 22.04 53.60	0.08 1.70	819 9.17 00 6.46 1 4.41	3 - 0.000	0.0002	520%
327 427 527	33,391 27,583	67.080 55.409 46.613	6.77 22.04 53.60	0.08 1.70	819 9.17 00 6.46 1 4.41	3 0.000 2 0.020 8 0.932	0.0002 07 0.0016 27 0.4351	520° 471°
327 427 527 627	33,391 27,583 23,062	67.080 55.409 46.613	6.77 22.04 53.60	0.08 1.70	819 9.17 00 6.46 1 4.41	3 0.000 2 0.020 8 0.932	0.0002 07 0.0016 27 0.4351	5207 4716 4049
327 427 527 627	33,391 27,583	67.080 55.409 46.613	6.77	0.08 1.70	819 9.17 00 6.46 1 4.41	3 0.000 2 0.020 8 0.932	0.0002 07 0.0016 27 0.4351 6 6.200	5207 4716 4049

Table 2. (Continued)

Temperature, °C	$-\log f_{0_2}$	$-\log f_{\mathrm{F}_2}$	$f_{\mathbf{H}_{2}}$	$f_{\mathrm{H}_2\mathrm{0}}$ —le	og∫нғ	$f_{\mathbf{CO_2}}$	$f_{\mathbf{CO}}$	$f_{\mathrm{CH_4}}$
		8. C	C, CH (C	FG, COH	F)			
327	43,453	70.609	6.78	Ó. 0055	10.937	$< 10^{-4}$	$< 10^{-4}$	5690
427	37.232	61.170	22.08	0.0205	9.342	$< 10^{-4}$	$< 10^{-4}$	522
527	32.519	54.016	54.17	0.0580	8 111	$< 10^{-4}$	$< 10^{-4}$	481
627	29.626	49.606	109.4	0.0530	7.732	$< 10^{-4}$	0.0002	443.
727	26.679	45.137	191.7	0.101	6,960	$< 10^{-4}$	0.0014	4070
827		41.773	302.3	0.134	6.476	$< 10^{-4}$	0.0055	372

 $<sup>^{\</sup>rm a}$  For buffer notation see Figures 1 and 2. More complete results and computer program available upon request (HPE).

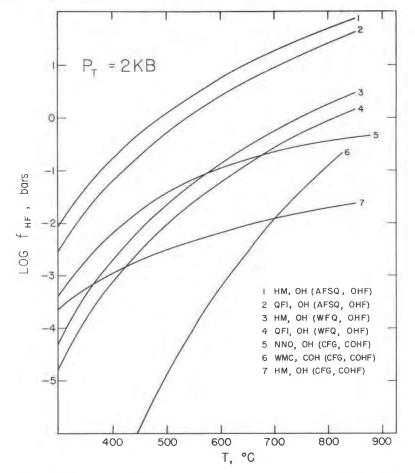


Fig. 2. Variation of HF fugacity with temperature calculated for a number of fluorine buffers at a total pressure of 2 kbar. Buffer assemblages (neglecting gas) are: HM, hematite +magnetite; NNO, nickel+nickel oxide, QFI, quartz+fayalite+iron; WMC, wustite +magnetite+graphite; AFSQ, anorthite+fluorite+sillimanite+quartz; WFQ, wollastonite+fluorite+quartz; CFG, calcite+fluorite+graphite.

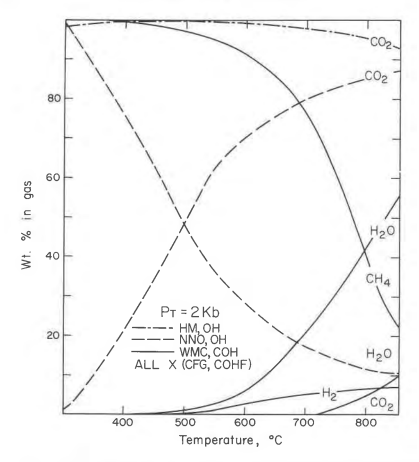


Fig. 3. Bulk composition of gas phase (major components only) in equilibrium with the CFG buffer at 2kbar plotted as a function of temperature and hydrogen fugacity. Dashdot curve: HM, OH (CFG, COHF); dashed curve: NNO, OH (CFG, COHF); solid curve: WMC, COH (CFG, COHF). HF is too minor a component to show up on this type of plot.

At 2 kbar gas pressure the partial pressure of HF never exceeds 50 bars, and is usually reckoned in tenths or hundredths of a bar. Hence, the bulk chemistry of the C-O-H-F system is very similar to that of the C-O-H system (French, 1966). Thus, for high oxygen fugacities (e.g., HM buffer), the gas phase is predominantly CO<sub>2</sub>; for intermediate oxygen fugacities (e.g., NNO buffer), H<sub>2</sub>O is dominant at low temperatures, but is rapidly replaced by CO<sub>2</sub> at higher temperatures; for very low oxygen fugacities (e.g., QFI buffer), H<sub>2</sub>+CO+CH<sub>4</sub> become the dominant species (Fig. 3).

It is important to consider the effect of these buffers on an experimental system. Consider a schematic fluorination reaction

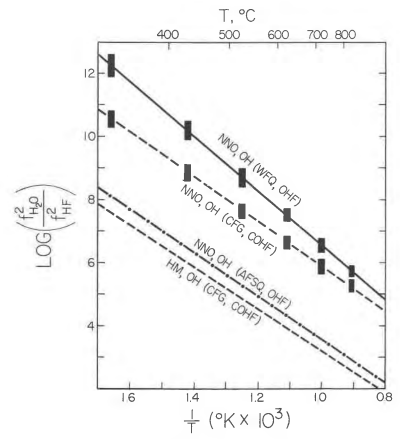


Fig. 4. Log  $(f_{\rm H2}o^2/f_{\rm Hr}^2)$  versus reciprocal temperature for four fluorine buffers. The height of the solid bars placed on the upper two curves represents the extent of uncertainty in the equilibrium constant which can be solely attributed to uncertainties in  $\Delta G^{\circ}$  values for the solid buffer phases. For symbols, see Fig. 2.

$$A(OH)_2 + 2HF \rightleftharpoons AF_2 + 2H_2O$$

Each buffer curve is plotted on a log K vs 1/T grid, with K's obtained from the calculated fugacities of H<sub>2</sub>O and HF for each buffer (Fig. 4).<sup>1</sup> The plot reveals a number of interesting facts pertinent to fluorine-OH reactions controlled by fluorine buffers: 1. Fluorination is favored by increasing temperature and, conversely, hydroxylation is favored with decreasing temperature; 2. The slopes of the buffer curves are nearly parallel; 3. The effect of external oxygen fugacity on fluorination is opposite

<sup>&</sup>lt;sup>1</sup> Assuming no solid solution.

T, °C	$\log f_{ m H_{2O}}$	$\log f_{\rm H_2}$	$\log f_{\mathrm{HF}}$	$\log a_{\mathrm{F}}$	$\log a_{\mathrm{H}^+}$
450°	2.735	0.289	-2.189	-3,49	-3.50
550°	2.945	0.592	-1.256	-3.55	-3.53
650°	3.080	0.806	-0.552	-3.74	-3.68

Table 3. Molecular and Ionized Species Calculated for the Buffer NNO (OH), WFQ (OHF) at 2  $\rm kbar^a$ 

for O-H-F and C-O-H-F buffers; for C-O-H-F buffers decreasing  $f_{\rm O2}$  favors hydroxylation, whereas for O-H-F buffers decreasing  $f_{\rm O2}$  favors fluorination, although in the latter case the effect is not detectable over ranges where  $P_{\rm H_2}$  is low compared to  $P_{\rm TOTAL}$ . (Table 2).

## EFFECT OF IONIZATION

In addition to the molecular dissociation of H<sub>2</sub>O and HF, some degree of ionization will occur, as governed by the ionization constants

$$K_{
m HF} = rac{a_{
m H}^{+}a_{
m F}^{-}}{a_{
m HF}} \ \ {
m and} \ \ K_{
m H_{2}O} = rac{a_{
m H}^{+}a_{
m OH}^{-}}{a_{
m H_{2}O}}$$

Ionization will, of course, have no effect on the fugacity ratios defined by the fluorine buffer equations. Moreover, these ionization equilibria will not effect the magnitudes of the individual calculated fugacities (i.e., will not effect the  $P_{\text{TOTAL}}$  equation) providing that the population of ionized species is always much less than the population of molecular species. From the values for the ionization constants for H<sub>2</sub>O and HF and 2kbar and temperatures of 450°, 550°, and 650°C (Barnes, et al., 1966, p. 404 and 407), values for  $a_{\rm H}^+$  and  $a_{\rm F}^-$  have been calculated for the NNO, OH (WFQ, OHF) buffer, assuming that H+, F- and OH- are the only ionized species present. The results (Table 3) predict that fluorine is present dominantly in the molecular state (as HF), and that the sum of the ionized components is sufficiently small relative to  $P_{\text{TOTAL}}$  so that they may be safely ignored. It is probably worth noting, however, that although the molecular versus the ionic approaches to fluorine (or fluoride) equilibria are very different, the results of the separate approaches must be identical. This point is emphasized by combining the equations for the ionic dissociation of H2O and HF with the restriction of electrostatic neutrality, i.e., that the sum of the number of moles of positive ions must equal the sum of the number of moles of negative ions, to form the single equation

<sup>&</sup>lt;sup>a</sup> Data for HF: Barnes, Hegelson & Ellis (1966), Table 18-4 C (p. 407). Data for  $H_2O$ : ibid, Table 18-1 C (p. 404); density of supercritical  $H_2O$  from Kennedy & Holser (1966), Table 16-1 (p. 378).

$$a_{\mathrm{H^+}} = \gamma_{\mathrm{H^+}} \bigg[ \bigg( \frac{K_{\mathrm{H\,F}} a_{\mathrm{H\,F}}}{\gamma_{\mathrm{F^-}}} \bigg) + \bigg( \frac{K_{\mathrm{H}_2\mathrm{O}} a_{\mathrm{H}_2\mathrm{O}}}{\gamma_{\mathrm{OH^-}}} \bigg) \bigg]$$

where  $\gamma_{\rm H}^+$ ,  $\gamma_{\rm F}^-$ , and  $\gamma_{\rm OH}^-$  represent the activity coefficients for the respective ions. The equation shows that once the activities of the molecular species are independently defined, the pH and hence the activities of all the ions in the gas phase are fixed.

## DEMONSTRATION OF EQUILIBRIUM

Equilibrium between fluorine buffer and charge was investigated using both crystalline hydroxyphlogopite and fluorophlogopite as starting materials. Phlogopite was chosen because it is one of the few phases showing (OH, F) solid solution for which both end members can be readily obtained in a pure state. By using both end members, the composition of the (F, OH) phlogopite solid solution in equilibrium with a given fluorine buffer at a fixed temperature can be approached from both sides. The composition of the intermediate phlogopites was determined by X rays on the basis of the d(005) assuming that a linear correlation can be made be-

Table 4. Run Table for Phlogopite Equilibrated with CFG, WFQ, and AFSQ Fluorine Buffers at 2 kbar

Starting Material	Temperature °Ca	Time, days	Final Composition
A. NNO, OH (CFG, COHE	`)		
gel	700	24	$F_{61}OH_{39}$
F-phlog	700	24	$F_{60}OH_{40}$
OH-phlog	700	24	$F_{64}OH_{36}$
OH+F-phlog	625	74	$F_{55}OH_{45}$
OH+F-phlog	550	108	$\mathrm{F}_{49}\mathrm{OH}_{51}$
B. HM, OH (CFG, COHF)			
OH-phlog	700	10	${ m F_{39}OH_{61}}$
OH+F-phlog	700	21	$F_{61}OH_{39} + F_{95}OH_{5}$
$F_{61}OH_{39} + F_{95}OH_5$	700	12	$\mathrm{F}_{96}\mathrm{OH_4}$
C. NNO, OH (WFQ, OHF)	)		
OH+F-phlog	775	10	$\mathrm{F}_{75}\mathrm{OH}_{25}$
OH+F-phlog	700	54	$\mathrm{F}_{71}\mathrm{OH}_{29}$
OH+F-phlog	550	108	$F_{60}OH_{40}$
D. NNO, OH (AFSQ, OHF	`)		
OH-phlog	700	68	$F_{95}OH_5$

<sup>&</sup>lt;sup>a</sup> Temperatures are  $\pm$ 5°C; Compositions are precise to  $\pm$ 2 mole %, but accuracy may be considerably less due to the linear extrapolation of the X-ray determination curve.

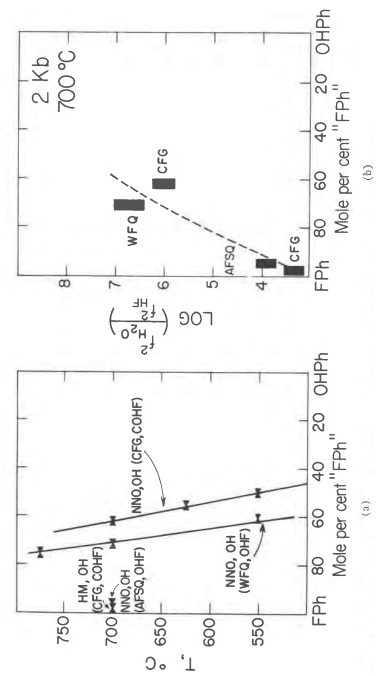


Fig. 5. a. Compositions of phlogopite in equilibrium with four fluorine buffers at 2kbar plotted as a function of temperature, Arrows show direction of movement towards the equilibrium value. FPh: fluorophlogopite; OHPh: hydroxyphlogopite.

fugacity ratio. The relative compositions of micas equilibrated with the NNO, OH (CFG, COHF) and NNO, OH (WFQ, OHF) buffers are in the b. Compositions of phlogopite equilibrated with four fluorine buffers at 2kbar, 700°C (see 5a) plotted as a function of the log of the squared opposite direction from that predicted by the fugacity ratio. tween the basal spacings of the end members and composition in mole percent. Compositions can be determined to a precision of better than  $\pm 2$  percent. Experiments were performed in standard externally heated furnaces using cold seal rod bombs at a total pressure of 2 kbar.

In most cases, a mechanical mixture of OH-phlogopite + F-phlogopite was used as a starting material once it had been established that reaction could be obtained using each phase separately; in these cases, the two-phase mixture was run under the desired conditions until a single, sharp (005) reflection was obtained from the quenched sample. The results are shown in Table 4 and Fig. 5. The AFSQ buffer equilibria were calculated using sillimanite as the aluminosilicate phase even though and alusite may be the stable polymorph at 2 kbar, 700°C (Gilbert et al. 1968). The small difference in the heats of formation of and alusite and sillimanite (0.48 kcal/mole at 968° K, Holm and Kleppa, 1966) results in a very small shift in the position of the buffer curve.

A number of facts emerge from the data. First, because the reaction can be approached from both sides, equilibrium between buffer and charge is demonstrated. Second, the run times needed to homogenize the two-phase phlogopite mixtures are considerably longer than equivalent run times needed to demonstrate the direction of a redox reaction at the same temperature. This could be either because the exchange of fluorine with hydroxyl is a slow process or because of the time required for the gas phase to equilibrate with the buffer. Third, a calibration problem is apparent. The phlogopites equilibrated with the WFQ (OHF) buffer are more fluorine-rich than those equilibrated with the CFG (COHF) buffer at the same temperature; the calculated  $f_{\rm H_2O^2}/f_{\rm H_3F^2}$  values (Fig. 4) predict the opposite result. This difficulty could arise from inadequately known  $\Delta G^{\circ}$  for the various buffer phases, or from nonideal mixing in the gas phase. Taking these possibilities in turn, the maximum errors in  $\Delta G^{\circ}$  for the various buffer phases as reported by Robie (1962) have been compiled and used to calculate maximum and minimum limits for log K as shown by the size of the error flags in Figure 4. Although this represents a fairsized source of error, it alone cannot explain the discrepancy. With regard to the second possibility, the method of reduced variables was used to determine fugacity coefficients for CO, CH4, and HF; as these are fairly crude approximations at best, the values must be viewed with reservation as possible error sources. More serious, however, these fugacity coefficients relate to pure homogeneous systems in which there are no interactions between different molecules; extrapolation of these compounding approximations to complex fluorine-bearing gases containing a large number of molecular species may prove to be unwarranted. To evaluate the problem, more data is needed using different buffers. Ultimately, the system O-H-F should be investigated systematically. Nonetheless, the phlogopite experiments are very promising. Once calibration difficulties can be resolved, phlogopite itself will provide a very useful sliding scale which can be used to record the  $f_{\rm H_2O}/f_{\rm HF}$  ratios in future experiments which involve complex fluorine-bearing hydrothermal gases.

## GEOLOGIC APPLICATIONS

The data obtained for the (OH, F)-phlogopites are much too incomplete to warrant any definite conclusions. However, the 700°C isotherm of (Fig. 5b) illustrates a very significant point. Consider the exchange reaction

$$KMg_3AlSi_3O_{10}(OH)_2 + 2HF \rightleftharpoons KMg_3AlSi_3O_{10}F_2 + 2H_2O$$
  
OH-phlogopite F-phlogopite

Assuming that the dashed curve in Figure 5b represents the equilibrium position of this reaction at 700°C and 2 kbar pressure, we can calculate the composition of the gas phase in equilibrium with an (OH, F)-phlogopite, using the relation

$$f_i = f_i^{\circ} \times X_i$$

where  $f_i$  is the fugacity of i in the gas mixture,  $f_i^{\circ}$  is the fugacity of pure i at the same P and T, and  $X_i$  is the mole fraction of i in the mixture. At 700°C, 2 kbar,  $f_{\rm H_2O}{}^{\circ}=1360$  bars,  $f_{\rm H_F}{}^{\circ}=4020$  bars. The correlation between gas and phlogopite composition was calculated from Figure 5b and is plotted in Figure 6. The distribution of fluorine between mica and gas is obviously very asymmetric, with the mica removing the fluourine nearly quantitatively from the gas. In other words, during crystallization phlogopite acts as a very efficient trap even for small traces of HF.

The effect of changes in *P*, *T* and biotite composition on the curve in Figure 6 is not known, but it is probable that a similar relationship exists for most igneous and metamorphic phlogopites and biotites.<sup>2</sup> If this is true, the gas phase in equilibrium with the most fluorine-rich phlogopite (75 mole % F-phlogopite, see Deer, *et al.*, 1962) would have contained no more than 0.05 mole percent HF. According to the biotite analyses

<sup>&</sup>lt;sup>1</sup> This conclusion seems to be at variance with the results of Noda and Ushio (1964), who were unable to grow very fluorine-rich phlogopite from an aqueous phase even with large fluorine excess in the starting material. In their experiments, however, solid KMgF<sub>3</sub> was also present and they did not determine the composition of the gas phase.

<sup>&</sup>lt;sup>2</sup> According to preliminary data of Rieder (1968), this extrapolation may not be valid for annite. It may also be questionable for the case of lepidolites, as indicated by the good correlation between lithium and fluorine contents (Munoz, 1966; Rieder, 1968).

tabulation of Foster (1960), a more common range would be 0.001–0.005 mole percent HF. At a gas pressure of 2000 bars, this would correspond to a HF fugacity of 0.04–0.2 bars. According to data of Wyllie and Tuttle (1961), this amount of HF does not affect the melting temperatures of albite and granite and probably other fluorine-free silicate assemblages; but it will strongly influence the thermal stability of (OH, F) solid solutions. Hence, mica-amphibole equilibria will also be affected by it.

Perhaps even more interesting than the low values of  $f_{\rm HF}$  and  $X_{\rm HF}$  in igneous and metamorphic gases is the ubiquitous *presence* of HF, as re-

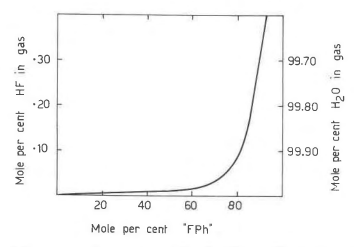


Fig. .6. Distribution of fluorine between phlogopite and gas at 700°C, 2 kbar, calculated from Figure 5b. The vertical scale is 200 times the horizontal scale FPh: fluorophlogopite. Right-hand ordinate valid only for an H–O–F gas.

flected by the mica compositions. As more calibrations become available, it will be possible to obtain a quantitative picture of the behavior of HF and hence F<sub>2</sub> in igneous and metamorphic processes.

## PRACTICAL CONSIDERATIONS: CHOICE OF FLUORINE BUFFER

An oxygen buffer is separated from the charge by a metallic membrane permeable only to hydrogen; thus, contamination of the charge by metallic ions is avoided. However, lacking a fluoride-specific membrane, the fluorine buffer gas is in direct contact with the charge and is only mechanically separated from the buffer phases by a crimped foil packet or tube. Thus, by means of exchange with the vapor phase, some transfer of material from the buffer to the charge can occur. This problem can be avoided by choosing a buffer whose components will be inert to the

charge under the desired experimental conditions. For example, for the phlogopite case described in this report, the only possible buffer contaminant is Ca, by means of the substitution of Ca for K in the interlayer site of the mica. However, in order to form Ca-phlogopite (clintonite or xanthophyllite, Deer et al., 3, p. 99) from phlogopite, alumina must be added to and silica expelled from the tetrahedral layer in order to maintain electrical neutrality; inasmuch as there is no source of additional alumina in the buffer and no excess quartz was observed in the quenched charges, it appears that contamination did not occur. Nonetheless, contamination of this type is a problem to which future users of the buffer method should be alerted.

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