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Fluid Inclusions Studies in Quartz from Fissures of Western and Central Alps

By Bernard P. Poty (Nancy)*), Hans A. Stalder (Bern)**) and Alain M. Weisbrod (Nancy)*)

With 12 figures and 8 tables in the text and 1 plate

Abstract

Fluid inclusions in quartz crystals of more than 120 alpine fissures from western and central Alps (Pelvoux, Mont Blanc, Aar, Gotthard and the Pennines) have been studied.

Microthermometry (freezing and heating) gives estimates of the chemical composition of the fluids in terms of three components: water, NaCl (for all the salts) and CO_2 . Density estimates may be inferred as well. It is shown that fluids are mainly salt aqueous solutions, excepted in the Gotthard and the Pennines, where CO_2 becomes an important component.

Chemical analyses of fluids occuring in inclusions give the relative amounts of the following solute components: K, Na, Ca, Mg and to a lesser extent, Cl and SO₄. The K/Na ratio seems to be well buffered in feldspars-bearing rocks. After it has been shown that albite grew in the disordered structural state, even at rather low temperatures, the equilibrium constants of the system adularia- high albite-KCl-NaCl aqueous solution are calculated and used as a tentative geothermometer. Pressures and depths of formation have been estimated using the calculated (from microthermometry results) densities of the fluids.

Estimated temperatures increase from the Pelvoux massif (340° C) to the Gotthard massif (505° C) . On the other hand the pressure (2.5 to 3 Kbar) and the overburden (9.5 to 11 km) are rather constant all over the Mont Blanc and Aar massives.

Zusammenfassung

Flüssigkeits- und Gaseinschlüsse von Quarzkristallen aus mehr als 120 alpinen Zerrklüften der West- und Zentralalpen (Pelvoux-, Mont Blanc-, Aar- und Gotthard-Massiv, Penninikum) sind untersucht worden.

Mikrothermometrische Messungen der flüssigen und gasförmigen Einschlussfüllungen auf dem Kühl- und Heiztisch führen zu Abschätzungen der drei Komponenten H_2O ,

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n° 1 – 54500 – Vandœuvre-les-Nancy and E.N.S.G., B.P. 452 – 54000 – Nancy (France). **) Naturhistorisches Museum Bern, Bernastrasse 15, 3005 Bern (Switzerland). NaCl (als Mass für alle Salzanteile) und CO_2 , und ebenso zur Abschätzung der Dichten der fluiden Phasen. Es wird gezeigt, dass die Einschlussfüllungen vorwiegend aus Salzlösungen bestehen; einzig im Gotthardmassiv und im Penninikum ist zudem CO_2 eine wichtige Komponente.

Chemische Analysen der fluiden Phasen in den Einschlüssen geben die Relativgrössen der Kationen K, Na, Ca und Mg. Auch die Relativgrössen der Anionen Cl und SO₄ wurden zum Teil bestimmt.

Das K/Na-Verhältnis scheint in feldspatführenden Gesteinen gut gepuffert zu sein. Es wird gezeigt, dass der Albit (der alpinen Zerrklüfte) in der ungeordneten Strukturvarietät auskristallisiert ist, auch bei relativ tiefen Temperaturen. Deshalb kann versucht werden, die Gleichgewichtskonstanten des Systems Adular-Hochalbit/KCl-NaCl der wässerigen Lösung als geologisches Thermometer zu gebrauchen. Der Bildungsdruck und die Gesteinsüberlagerung werden ebenfalls abgeschätzt, dies auf Grund der Bildungstemperatur und der Dichte der Einschlussfüllungen (letztere als Ergebnis der Mikrothermometrie).

Die abgeschätzten Temperaturen nehmen vom Pelvoux-Massiv (340° C) zum Gotthard-Massiv (505° C) zu. Auf der andern Seite sind der bestimmte Druck (2,5 bis 3,0 Kbar) und die Gesteinsüberlagerung (9,5 bis 11 km) für das Mont-Blanc- und das Aar-Massiv mehr oder weniger konstant.

INTRODUCTION

Alpine fissure minerals from Switzerland have been described by KENNGOTT (1866), KOENIGSBERGER (1919, 1940), P. NIGGLI (1940), PARKER (1940, 1953) and STALDER, DE QUERVAIN, E. NIGGLI and GRAESER (1973, revised edition of PARKER, 1953). Most alpine fissure minerals from the French alps have been described by LACROIX (1893–1913). In the Soviet union, where fissure minerals occur commonly in the Urals and the Caucasus a considerable amount of work has also been done. GRIGORIEV (1960) gives a review of these works.

The relations between fissure mineral formation and alpine metamorphism are still being discussed. Radiometric dating isotopic measurements and fluid inclusion studies should help to gain an understanding of these relationships.

Radiometric dating (Rb/Sr and K/Ar) have been performed on several fissure minerals (JÄGER et al. 1967, LEUTWEIN et al. 1970, ARNOLD 1972, PURDY and STALDER 1973). Some of the published data are listed in table I. Radiometric dating can only indicate the time when minerals ceased being in interaction with fissure solutions. In the Mont-Blanc massif ages for fissure minerals (adularia and muscovite) are quite distinct from those of country rock minerals. Ages of fissure minerals might be true ages. In the Aar massif some K/Ar values are useless because of demonstrated argon over-pressures during growth. In the Simplon area the K/Ar method gives a consistent difference of 4 my between country rock and fissure minerals. But in many places country rock and fissure mineral give the same cooling age.

The presence of fluid inclusions in alpine fissure minerals has been known for a long time but attention has been focused on them only in the last decade.

Locality	Method	Mineral (f) fissure (r) rockforming	Age in million years (different rocks or minerals)	Literature
Mt-Blanc-Massif				
Fissure Nr. 64–207	Rb- Sr	Mu (f)	14,8	LEUTWEIN et al. (1970)
	K-Ar	Mu (f)	$13,5\pm2;\ 13,4\pm2$	LEUTWEIN et al. (1970)
Fissure Nr. 68–001	K-Ar	Ad (f)	18,3 ± 2 ; 15,8 ± 2	LEUTWEIN et al. (1970)
Gotthard - $Massif$. ,
Freilaufstollen Val Casatscha-	K-Ar	Mu (f)	$16,0\pm0,5$	Purdy, Stalder (1973)
Sta Maria	Rb- Sr	Bi (f)	$\begin{array}{c} 16,3\pm0,8;16,7\pm0,9;16,7\pm0,7\\ 16,0\pm0,9 \end{array}$	ARNOLD (1972)
	Rb- Sr	Bi (r)	$16,8\pm1,7$; 17,9 $\pm2,0$; 16,6 $\pm2,0$	Arnold (1972)
Pennines				
Simplontunnel different fissures	K-Ar	Mu (f)	9,8 \pm 0,3; 10,2 \pm 0,4	Purdy, Stalder (1973)
	K-Ar	Ad (f)	$8,5\pm0,1;\ 8,2\pm0,1$	PURDY, STALDER (1973)
	K-Ar	Mu (r)	14	PURDY, JÄGER (in prep.)
	Rb-Sr	Bi (f)*)	14,3 \pm 2,7; 12,8 \pm 0,9; 12,7 \pm 2,4 12,0+2,3	Jäger et al. (1967)
	Rb- Sr	Bi (r)	$11,0\pm1,0$	Jäger et al. (1967)
Cavagnöö, Val Bavona	K-Ar	Mu (f)	$13,7\pm0,4$	PURDY, STALDER (1973)
	K-Ar	Ad (f)	$13,6\pm0,2$	PURDY, STALDER (1973)
	*\	E:	and an a farmer and a farmer	(

Table I. Selected Ages of Alpine Fissure Minerals

*) Fissure biotites but not from open fissures.

As early as 1906 KOENIGSBERGER and MÜLLER gave two chemical analyses performed on big inclusions from the Aar massif. Those two analyses were probably the first ever made on a metamorphic fluid. DEICHA (1949, 1955) began a series of studies performed with a variety of methods. Thermometry under the microscope (microthermometry), was the main method used by YPMA (1963) on Belledonne, STALDER (1963) on the Grimsel (Aar), POTY (1967–1969) on quartz from La Gardette (type locality of Dauphiné habit) and Mont-Blanc, STALDER and TOURAY (1970), MULLIS et al. (1973) on quartz from the northern calcareous alps. Besides those essentially microthermometric studies TOURAY (1968) gave mass spectrometric analyses of gases.

All these studies gave a picture of fluids associated with a phase of alpine metamorphism. This paper continues in broadening the region of sampling (Pelvoux, Mont-Blanc, Aar, Gotthard and the Pennines), with increasing the number of samples, and with improving the accuracy of microthermometry data. Chemical analyses of leachates are given and a new method determining temperature is proposed leading to T-P estimates of alpine fissures formation.

I. Data from Microthermometry

Analytical techniques

Generally speaking, microthermometry consists of measuring, under the microscope, the temperatures of physical changes undergone by the fluid phases trapped in inclusions.

Investigations were performed at Nancy and Fribourg with the aid of identical heating and freezing stages, designed to work in the -160 to $+380^{\circ}$ C range. These apparatus were built in the CRPG at Nancy.

The lower temperatures are obtained by nitrogen gas previously cooled in liquid nitrogen and flowing through the stage. After inclusions have been frozen, they are slowly warmed at a constant rate of 1 to 2° C/minute, by controlling the gas flow with a precision-valve. High temperatures are obtained with a heating resistance.

Temperatures are measured with a platinum resistance sensor set very close to the sample. Measurements are standardized against the melting points of various high purity chemicals, such as hexane (-94.3° C), heptane (-90.6° C), n-dodecane (-9.6° C) and distilled water (0.0° C). Natural CO₂-inclusions from Calanda (checked for purity by TOURAY, 1968; melting point: -56.6° C) were also used. Standardization at high temperatures was made with special chemicals from Merck Corporation.

The precision of measurements is about $\pm 0.2^{\circ}$ C in the range -12 to $+31^{\circ}$ C, and $\pm 1.5^{\circ}$ C in the range -150 to $+350^{\circ}$ C.

The procedure of investigation has been described in detail by ROEDDER (1962). For two phase inclusions, the following temperatures were measured: lower melting point (solidus or eutectic temperature), higher melting point (liquidus, often called "freezing point" in this paper), dissociation of possible hydrates, homogenization. Possible occurrence of CO_2 was checked with a crushing stage (DEICHA, 1955; ROEDDER, 1970). For inclusions with liquid CO_2 , the temperature of melting of frozen CO_2 and the temperature of homogenization of the two CO_2 -rich phases were also determined. Quite often, inclusions with an immiscible CO_2 -rich phase decrepitated before reaching the temperature of bulk-homogenization of the fluid.

Although various components are present as solute species, aqueous solution is usually described in terms of the theoretical NaCl-H₂O system. Thus, it is possible to calculate an "NaCl equivalent" content from the liquidus temperature.

The determination of salt content in the CO_2 -rich inclusions is difficult, because of the formation of CO_2 -hydrate (CO_2 -5.7 H₂O). Less CO_2 -hydrate crystallizes when a fast cooling rate is used than with a slow cooling rate (1° C/min.). In the latter case, salt concentration increases considerably because more water is used during hydrate formation. In other studies, differences up to 1° C in the freezing point of the aqueous solutions, due to variable cooling rates, were observed. The data given in this paper are for the rapid cooling rate of 20° C/min.

RESULTS

Many quartz samples and a few axinites (Pelvoux) from more than 120 fissures were investigated. Most inclusions appear in healed fractures (second-ary inclusions), but some primary inclusions were also recognized.

The most important results are listed in table II and III, and need some general comments.

Belvedere, Furka (Q 113) and Grasso di Froda I (kr 24)								
Nr. of the inclusion	Melting temp. aqueous solution		Equival. [NaCl] weight %	Homogen. temperature				
Q 113–1	-0,9		1,5	194				
\hat{Q} 113–2	-2,0	-	3,4	207				
Q 113-3	-2,1	-	3,6	208				
kr 24–1	-2,75	9,9	4,6	245				
m kr~24-2	-2,9	9,6	4,8	272				
kr 24–4	-3,0	9,1	5,0	274				

Table II. Examples of variations of salt and CO₂ content in fluid inclusions

Fluh, St. Niklaus (6713)

Nr. of the inclusion	Homogen. of the liquid and the gas. CO ₂ -phases	Vol. percent of the CO_2 -phase	Density of the CO_2 - phase at the homog. temp.	Remarks
$\begin{array}{c} 6713/1,\ 2\\ 6713/3-5\\ 6713/8\end{array}$	29 L 24 G 20 G	$90\\85\\45$	0,63 0,22 0,19	Salt content of the H_2O -rich phase \pm con- stant at 5,5% [NaCl] eq.

1. Although age relationships between secondary inclusions are difficult to detect, it seems that the older inclusions are more concentrated in salt. Examples of average values are given in table II. The largest variations are observed in the neighbourhood of evaporitic sediments (e.g. = Camperio, Bitsch, Lengenbach).

2. Some older fluids are CO_2 -rich. In such inclusions, a strong decrease of the CO_2 content from the inner to the outer part of the quartz crystals is usually observed (see table II). On the other hand, late CO_2 -rich fluids are well known all over the Alps, mainly in the so-called sceptre-quartz; they are not considered in this paper.

3. Besides CO_2 , other trace gases are present in the inclusions, as shown by mass-spectromeric analyses (TOURAY, 1968; ZIMMERMANN and POTY, 1970). Although quantitative estimations are not possible with microthermometry, such gases have been detected here. At Kammegg, for instance, an unknown hydrate was discovered and probably indicates the presence of hydrocarbon.

However, the melting temperature of CO_2 -rich phase in the inclusions investigated here departs only slightly from the melting temperature of pure $CO_2(-56.6^{\circ} \text{ C})$. Out of 40 determinations, 33 are between -56 and -57° C , and the others remain very close to these values. Therefore, the vapor phase of three-phase inclusions from alpine quartz contains almost pure CO_2 .

4. The dissociation temperatures of the CO_2 hydrate that have been measured here are scattered in the range +7 to +11.5° C. According to LARSEN (1956), these variations can be related mainly to the salt content of the aqueous solution (a high salt content decreases the dissociation temperature).

5. Some data about the solidus temperature of the aqueous solutions have already been given and discussed (POTY and STALDER, 1970). The new data confirm the former conclusions, and we have abstained from any discussion here.

6. Although the calculated compositions of the inclusions are based upon estimates and involve some simplifications, they give more reliable values than many of the mass-spectrometric analyses, for the later often give only average composition of different generations of inclusions. Moreover, the interpretation of mass-spectrometric analyses is made difficult by problems such as irreversible adsorption of some gases on surfaces.

Table III. Microthermometry results

H_2O -rich phase (1)–(3)

- (1) Liquidus temperature ("freezing point") in °C.
- (2) Concentration of NaCl (wt %) of the theoretical NaCl-H₂O aqueous solution, the freezing point of which is the same as the freezing point of the real solution, given in column (1). Liquidus curve from Scatchard and Prentiss (1933) and Rodebush (1918).
- (3) Density of the aqueous solution refered to the $NaCl-H_2O$ system.

CO_2 -rich phase (4)–(6)

- (4) Homogenization temperature of the two CO_2 -rich phases, in three phase inclusions (G = homogenization in vapor state, L = homogenization in liquid state). In two phase inclusions, the CO_2 content is estimated from runs with the crushing stage, and indicated as follows:
 - 0: no CO₂ detectable,
 - x: traces of CO₂ (about 1,5 wt %),
 - xx: some CO_2 (about 3 wt %),
 - xxx: some more CO_2 (about 4,5 wt %).
- (5) Volume percent of the CO_2 -rich phase from a microscopic estimation.
- (6) Density of the CO_2 phase at the homogenization temperature from the P-V-T data after different authors in Landolt-Börnstein (1960).
- (7) Bulk homogenization temperature of the fluid phases in °C. The number of fluid phases at room temperature is given in brackets. The values refer to the representative interval or representative highest value.
- (8) to (10) Estimated composition of the fluid phase, in terms of the components NaCl- H_2O-CO_2 .
- (8) H_2O in weight percent.
- (9) Equivalent [NaCl] in weight percent.
- (10) CO_2 in weight percent; calculated from volume percentage (estimated under the microscope) and from the density of homogenized CO_2 . The content of CO_2 in the aqueous solution coexisting with liquid CO_2 was taken to be 5.5 wt % at 25° C and 63 bars (Wiebe and Gaddy, 1940). This mean value is too high for salt-rich inclusions, too low for pure water.
- (11) Number of studied inclusions.
- (12) Notes, remarks:
 - (7) = (1) Homogenization and cryometry were made on the same inclusions.
 - (7) \sim (1) Homogenization and cryometry were made on analogous inclusions.
 - (7) \neq (1) Homogenization and cryometry were made on different inclusions, homogenization generally on younger ones.
 - G: during the total homogenization the liquid phase disappears. Without remarks: the gaseous phase always disappears.

Notes (12)		hydraw 2,-1.5 Bydraw Bydraw Bydraw		(17)-(12) (17)-(12)-(12)-(12)-(12)-(12)-(12)-(12)-(12
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Table IV. Chemical analyses of fluid inclusions

(all the chemical analyses were made at the CRPG, Nancy, under the direction of K. Govindaraju and M. Vernet)

Locality	≠ analysis	K ppm	Na ppm	Ca ppm	Mg ppm	Cl	S04 ppm	K/Na at	Ca/Na at	Mg/Ca at	<u>Ca + Mg</u> K + Na at	$ \frac{SO_4}{Cl} \sum_{at} $	anal an anal cat eq
Pelvoux													
La Gardette	53	1.60	13.20	.194	.032			.071	.008	.272	.010		
11	56 243	.325 1.58	2.25 22,125	.09 .195	.019	36.5	.33	.085 .042	.230 .005	.348 .042	,029 .005	.003	1.02
	251 323	1.09	12.87 17.02	.076	.015	22.5 29.20	.21	.050 .054	.003	.325	.004	.003	1.08
Alpe d'Huez Col de Maronne	54 131	2.24	12.40	.522	.105			.106	.024	.332	.029		
Rampe des Commères	87	7.60	29.40	27.60	.073			.152	.539	.004	.470		
17	137 138	8.70 10.31	34.30 39.00	22 .131 25 . 74	.060			.149 .155	.370 .379	.004 .004	.323 .329		
Mont Blanc													
Trient	116	.200	.75	.037	.006			.157	.028	.267 .175	.005		
" Charlet Straton	117 42	.700	2.73	.132	.014			.159	.018	.288	.020		
11	43 62	1.20 3.46	4.17 16.00	.154 2.60	.019			.169	.093	.036	.085		
n n	76 130	.455	1.72	.079	.010			.156	.026	.209	.027		
" Le Tour Noir	248 37	2.425	10.368 6.55	1.731	.008	24.2	.05	.137	.096	.008	.085	.001	1.14
11	40	1.76	10.70	.222	.037			.097	.012	.275	.014		
11	41 324	2.17	8.03 1.203	.218	.056	2.40	<.01	.159	.016	.423	.019	.000	1.10
Col des cristaux Les Courtes I	326 242	.593 2.59	2.486 8.834	.004	.005	5.16 17.5	.04	.141	.001	2.06	.002	.003	1.18
" II	115	.455	1.55	.037	.010			.173	.014	.446	.017		
" II " III	114 112	.810	3.0	.085 .105	.012			.159	.026	.251	.028		
" III " III	111 110	.517	1.59 1.785	.138	.013			.191	.050	.155	.048		
Grands Montets	118	.517	1.945	.057	.010			.156	.017	.289	.019		
Les Rachasses	119 108	.74 .37	2.69	.118	.015			.153	.048	.182	.049		
" Bochard	109 325	.45 3.854	2.78 18.52	.072 5.356	.014	50.00	.05	.095	.015	.321	.018 .148	.000	1.20
Talèfre	67	2.35	8.60	.790	.029	23.20	.07	.161	.053	.061	.048	.001	1.05
Pierre Joseph Les Périades	327 113	3.301 1.53	12.03 4.96	.210	.019			.181	.024	.149	.025		
" Les Mottets	331 126	.818	3.006	.146	.003	6.40	.01	.160	.028	.034	.025	.0006	1.14
n n	127 128	.305	1.17 1.47	.024	.004			.153	.012	.275	.013		
H .	250	3.565	18.495	.446	.024	32.5	.20	.113	.014	.089	.013	.002	1.00
Tunnel	125 129	2.53 1.22	11.97 6.70	1.346 3.103	.021			.124	.065	.026	.059 .240		
Aar													
Gauli "	245 330	.69	3.990 .715	.188	.011	6.8 1.48	.21	.101	.027	.096	.027	.011	.98 1.16
Kammegg	79	1.63	6.05	.275	.033			.158	.026	.198	.027		
Hinter Wasen Kl. Lauteraarhorn	35 23	5.60	23.4 1.10	.447 .014	.036			.168	.007	.589	.010		
Unterer Kessiturm Gerstengletscher	144 33	.580	2.30	.616	.051			.148	.154	.136	.152		
	34	2.32	7.82	.534	.024			.174	.039	.074	.036		
" Tiefengletscher	141 25	.510 1.165	1.87 3.90	.040 .184	.020			.176	.027	.179	.027		
" Göschenen I	26 319	1.66	4.70 5.530	.116	.016	11.10	.07	.208	.014	.227	.014	.002	1.11
Göschenen II	317	1.600	6.830 1.758	.243	.005	13.88	.20	.138	.020	4.12	.017	.005	1.18 1.13
Brig Bärfel	328 329	.337	5.56	.296	.002	11.08	.17	.138	.031	.011	.027	.006	1.09
Gletsch "	88 332	.235	.710	.000	.003	.85	.02	.195 .174	.000	1.67	.003	.009	.96
Rhonegletscher "	19 143	•95 •590	3.45	.098 .146	.016			.162 .197	.016	.268 .225	.018 .049		
Gotthard													
Faulhorn	320	.237	1,196	/	/	2.24 10.8	.20	.117	.045	.101	.044	.033	1.16
Muttengletscher Ri della Fibbia	246 86	1.185	5.865	.459 .078	.028			.137	.007	.254	.007		
Grasso di Froda II Grasso di Froda I	247 314	.15 .141	2.035	.000	.006	2.00	1.04	.043	.000		.003	.191 .030	(.84) (.44)
Piz Garviel	45 142	1.66 1.54	4.55 3.65	.970 .474	.031			.215	.122	.053 .070	.106		
Pennines													
Gischihorn	36	.925	3.01	.068	.012			.181	.013	.291	.014		
" Gh. del Cavagnöö III	140 318	.080	.245 .194	.020	.002	. 35	.14	.192	.047	.165	.046	.148	1.45
Lago Bianco	315 316	.691	5.860	.276	1	10.60	.02	.069	.027		.025	.0007	1.05
	110	.UEI			1,	.40							
Fusio Aquarossa Campo Blenio	312 89	.165	1.684	.576	.021	2.65	.25	.058	1.223	.060	1.094	.035	1.03

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Pelvoux Massif

High salinity, with freezing temperatures sometimes lower than the NaCl-H₂O eutectic, characterizes the fluids from the Pelvoux massif. This is due to the presence of divalent cations which are expected in fissures located in amphibolites and/or close to the Triassic sediments. Homogenization occurs at about 180° C.

Mont Blanc Massif

Freezing temperatures range from -3.5 to -7.0° C (6 to 10.5 wt% equivalent NaCl, with an average around 8.5 wt%).

In this massif, fissures are located in a quite homogeneous granite, far enough from sediments, so that the values measured here are probably controlled by irreversible leaching of the granite and/or by reactions forming new minerals.

 CO_2 concentration is generally quite low (less than 2 wt _0^{\prime}). However, it can reach 5 wt $_0^{\prime}$ in some places (Tour Noir) or even higher values (Les Courtes, Les Amethystes, POTY 1969). These high concentrations are always related to late fluids in vertical sheared zones.

Homogenization temperatures range between 150° and 210° C, with 36°_{0} of the values falling in a narrow range (170 to 185° C) (fig. 1). Higher values are always related to the presence of CO₂, indicated by liquid CO₂ or by CO₂ hydrate formation.

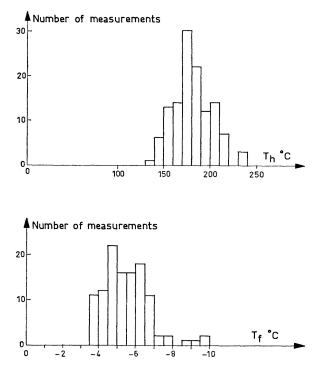


Fig. 1. Frequency distribution diagrams of microthermometric measurements of fluid inclusions from Mont-Blanc massif. Upper diagram: homogenization temperatures; lower diagram: freezing (liquidus) temperatures.

Aar Massif

 CO_2 rich inclusions, observed in quartz associated with ankerite, are rare in the Aar massif. The salt content of aqueous solutions varies from zone to zone. It is about 8 wt% NaCl eq. in the area of crystalline schists (North of the massif) and in the eastern part of the massif, though significant differences are present. In the granitic zone of the Central Aar massif (Grimsel area), the salt content is fairly high (11 wt%). It decreases again (5.5 wt%) in the southern gneisses. Some extreme values (western area) can be explained by the proximity of evaporites.

If the CO_2 -rich inclusions are not taken into account, the homogenization temperatures fall into a rather narrow range, with an average around 190 to 200° C. This point has been quoted long ago by KOENIGSBERGER and MÜLLER (1906).

Gotthard Massif

The salt content of aqueous solutions from the Gotthard massif is about 7 wt % NaCl eq. The variations are small, with two exceptions (extremely high salt content).

In comparison with the Aar massif, fluid inclusions from Gotthard are quite rich in CO_2 . In most of the quartz crystals, inclusions containing more than 10 wt% CO_2 can be seen. Nevertheless, aqueous solutions with two phases are still dominant; they are more recent than many of the CO_2 -rich fluids.

The homogenization temperatures of three-phase inclusions range between 250 and 295° C.

Pennines

The salt content is about 6 wt % NaCl eq. It is quite constant except in the youngest inclusions where it decreases considerably. The very high salt content at Camperio might be related to a gypsum deposit in this area (inclusions with halite crystals have been found there).

The Pennines are even richer in CO_2 than the Gotthard massif. Threephase inclusions with a density of CO_2 above the critical value (0.475 g/cc) are common. But 3-phase inclusions with a density of CO_2 lower than 0.475 g/cc and 2-phase inclusions are also abundant. The CO_2 content decreases with time, and the youngest inclusions contain only a small amount, if any, of this component.

The homogenization temperatures of 3-phase inclusions range from 260 to 360° C. The average value is much higher than in the Gotthard and Aar massives. The low homogenization temperatures recorded in 2-phase inclusions occuring in "needle quartz" from Gh. del Cavagnöö (208° C) and Paltano (220° C), support field evidence which indicate that these crystals belong to the latest crystallization event in the area.

II. Chemical Analyses of Leachates

Seventy nine chemical analyses of fluid inclusions were performed, using the procedure proposed by ROEDDER (1963), with a few improvements. The analyzed ions are K, Na, Ca, Mg and, for 27 samples, Cl and SO_4 .

Preparation of the Samples

Cleaning the samples was carried out as follows:

- Immersion for one hour in boiling concentrated high purity nitric acid (Merck 454).
- Immersion for one hour in boiling distilled and deionized water (DDW).
- Immersion for five days in a continuously stirred mixture of DDW and anionic + cationic resins. This step is of critical importance: a duration of less than 5 days yields bad blanks and unreliable results.

After cleaning, each sample was inserted into a stainless steel tube (Sandvik 3 R 12 - OD 20 mm - length 140 mm) closed at one end. This tube was previously cleaned by immersion during one hour in boiling nitric acid, and another hour in boiling DDW.

Crushing and Leaching

Crushing was performed with a 7 Kbar press, and the powdered sample leached with DDW on a Millipore assembly (Pyrex funnel and base, stainless steel filter support screen, 47 mm filter, 0.45 μ m) to give 60 ml of solution.

Owing to the low concentration values found in the leachates (a few hundreth of a ppm to a maximum of 15 ppm for most of the samples), the possibility of contamination was seriously considered. A blank was made for each sample in the following way: just before crushing, DDW was poured into the stainless steel tube containing the sample, and this water was then passed through the filter assembly.

All those steps, from cleaning the surfaces with water and resins to obtaining the bottle of leachate, were performed in a room exclusively devoted to this purpose.

Analytical Methods

K, Na, Ca and Mg were analyzed with a Perkin-Elmer 303 atomic absorption apparatus with digital concentration readout. Sensitivity of determinations is a few thousands of a ppm for all those elements, and accuracy is believed to be close to one hundreth of a ppm.

Concentrations in the blanks were usually less than a few thousands of a ppm and were considered perfect in this case. Sometimes they were in the 0.01–0.1 ppm range. When they were equal to or more than 0.1 ppm, contamination was considered to be too large. This occurred in six out of the 79 analyses. However, the results were kept because the concentrations in the six corresponding leachates were high enough.

Chloride was analyzed with a colorimetric method, using ferric ammonium sulfate and mercuric thiocyanate as indicated by ROEDDER (1963). This method allows determinations down to 0.01 ppm Cl and works fairly easily in routine.

Sulfate determinations (methylene blue colorimetric method) were more difficult. As

suggested by ROEDDER (1963) the procedure of JOHNSON and NISHITA (1952) was used first. But the procedure described by GUSTAFSSON (1959) appears to yield more reproducible results, and later analyses (312 and after) were done by that method. Sensitivity is 0.01 ppm and accuracy is believed to be close to 0.05 ppm.

RESULTS

Only a few of the significant components of the solutions were analyzed. H^+ , Li^+ , Fe, HCO_3^- and F^- are probably the most important ions not looked for. Since the quantity of water in the sample was not determined, the actual concentration of individual ions in the original fluid can be estimated through microthermometry results.

On the other hand ratios between analyzed cations and anions are known with good accuracy for most samples. The atomic ratios: K/Na, Ca/Na, Mg/Ca, Ca + Mg/K + Na, SO_4/Cl and the balance of charges between analyzed anions and analyzed cations are especially significant.

It should be noted also that the terms "ion", "cation" and "anion" refer to a theoretical totally ionized aqueous solution. This does not mean that those components were actually present as simple ions at the time of quartz growth, or in the analyzed leachates either.

K/Na

The values are clustered by regions on five different frequency distribution diagrams (fig. 2).

The diagram for the Mont Blanc massif shows an especially sharp peak, close to 0.161 and scatter from 0.095 to 0.191.

In order to get such a sharp peak two conditions have been met:

- a) solutions have been buffered by some mineral association and we will see that albite and adularia made the buffer;
- b) analyses must have been accurate and this is due to the quality of material from the Mont Blanc massif.

The scatter of the results probably has several causes:

- The cavities studied here were probably not formed at exactly the same time, and differences in PT conditions may have been large, leading to different K/Na ratio of the fluid. This point should be important in the Pennines.
- The nature of the anions is known to affect the equilibrium K/Na ratio for a given reaction (IIYAMA, 1970). Most experimental studies were made with chloride as anion. But for example the presence of HCO_3^- (not analyzed in this work) lowers the K/Na ratio. Total concentration and abundance of other cations should not affect the results very much.

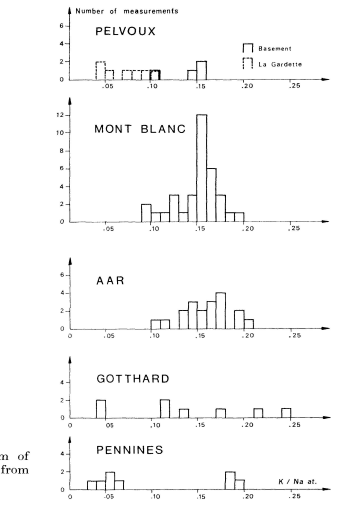


Fig. 2. Frequency distribution diagram of K/Na (atomic ratio) in fluid inclusions from Alpine fissures.

- Errors in chemical analyses do not seem to be a major cause of the scatter of the results since K and Na concentrations are always over 0.3 ppm, the value above which our technique appears to be accurate enough.

In the Pelvoux massif, the data are clustered into two groups. The cluster around 0.15 is related to cavities in the basement not very close to Triassic sediments (Rampe des Commères). The other is related to cavities or veins (La Gardette) in the basement at the contact with Triassic sediments including dolomites.

Here HCO_{3}^{-} , not measured but quite likely present, lowers the K/Na ratio as shown by IIYAMA (1970).

In the Aar massif, values are clustered in the manner of the Mont Blanc massif. The peak is not so sharp and this is probably due to the small number of analyses. Its value lies around 0.175, a bit higher than in Mont Blanc (0.161); the maximum found (0.208) is also higher.

The cluster of values for Mont Blanc and Aar suggest that the cavities were made in mainly one process, which is also suggested by field evidence. In the Gotthard and the Pennines the data are, on the contrary, quite scattered. But the structural history of these regions is more complex. Crystallization in some clefts began with scapolites and ended with zeolites (e.g. at Camperio). Such an evolution in paragenesis suggests a long duration of growth and a decrease of temperature. This has to be reflected in the chemistry of the solutions.

Ca/Na

The values of this ratio are so low (less than 0.05) that no variation can be suspected. There are a few exceptions in the Pelvoux massif, which can be explained by the occurrence of tremolite in veins (fig. 3).

Owing to the large amount of oligoclase that has been incongruently and irreversibly leached during hydrothermal processes, one could expect a higher Ca content in the fluids. The low values measured here indicate that the Cacontent of solutions equilibrated with new-precipitated minerals such as calcite, epidote ... is very low under the P and T conditions that prevailed during the formation of alpine veins.

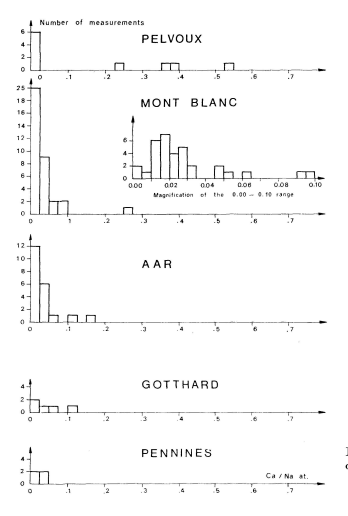
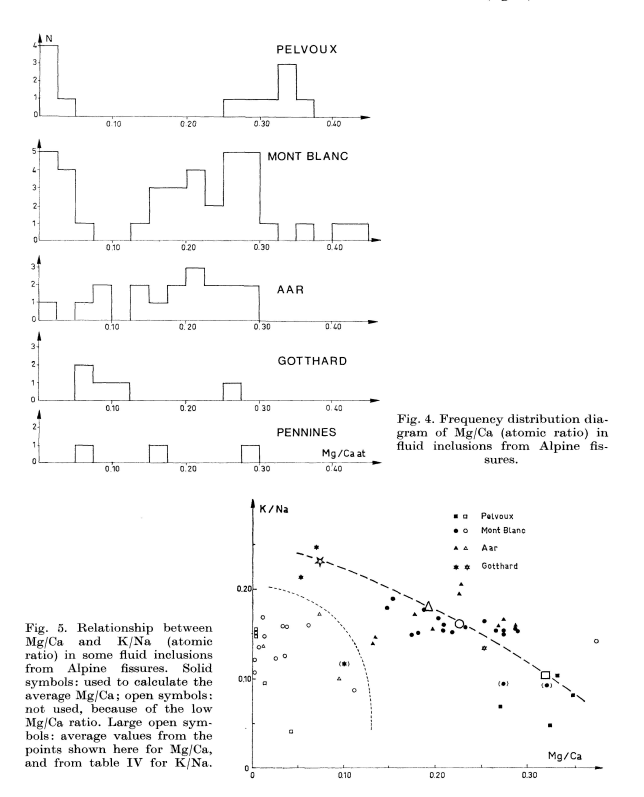


Fig. 3. Frequency distribution diagram of Ca/Na (atomic ratio) in fluid inclusions from Alpine fissures.

Mg/Ca

The results for Mg/Ca ratio are more scattered than those for K/Na. In Pelvoux, Mont Blanc and Aar, two sets of values are found (fig. 4).



- 1. 0.01 to 0.10 increasing from Pelvoux to Aar; these low values are found mainly in veins that occur in rocks containing Ca-rich minerals.
- 2. 0.13 to 0.37, decreasing from Pelvoux to Aar.

One of the most commun buffers of the Mg/Ca ratio in veins is the calcitedolomite association. Because these two minerals do not always occur at equilibrium in alpine veins, and because Fe-bearing carbonate (ankerite) is usually found in place of dolomite, buffering is imperfect. However, the decrease of Mg/Ca from Pelvoux to Aar (and probably Gotthard) – i.e. with increasing temperature – agrees with experimental data (Holland, 1967). If these values are plotted agains K/Na, a rough trend appears, as seen on fig. 5.

(Ca + Mg)/(K + Na)

This ratio does not show any systematic change from one massif to another. Concentration of divalent cations is much lower than concentration of alkalis, generally by two orders of magnitude. However, some values are fairly high (0.1 to 0.5) and this is related to a Ca-rich and (K + Na)-poor surrounding; the example of the "Rampe des Commères" is particularly significant, where the fissures are located in amphibolites.

It can be seen that abnormal values of the K/Na ratio are usually obtained when (Ca + Mg)/(K + Na) is high.

SO₄/**Cl**

In La Gardette vein, close to the Triassic sediments, SO_4/Cl of the fluid phase is amazingly low: 3×10^{-3} . This means that at high T and P only halides will be dissolved and anhydrite will remain. In the Mont Blanc massif values are in the same range, slightly higher in the Aar massif (8×10^{-3}) and much more scattered in the Gotthard and the Pennines (1×10^{-3} to 35×10^{-3}), with some as high as 190×10^{-3}).

Anions-Cations Charge Balance

The ratio \sum analyzed anions $/\sum$ analyzed cations in equivalent (i. e. $\sum n_A z A^{z-} / \sum n_M z M^{z+}$) is amazingly constant. Out of 25 values, only two are lower than 1.00 (0.98 and 0.96) and two are higher than 1.20 (1.25 and 1.44). More than 50% fall in the range 1.00–1.10; the average value is 1.08.

Since all species were not analyzed, only a tentative interpretation can be made. Of the data that are missing, those concerning Fe and carbonate should be the most important. Because the latter is probably more abundant than the former, the excess of negative charges is still larger than the values obtained above. This excess must be balanced by hydrogen (occuring as H^+ and in acid molecules, complex anions or complex molecules), the concentration of which was therefore rather constant and fairly high in the solutions of alpine veins.

III. Discussion: The Place of Fissures in Alpine Metamorphism

Results given by microthermometry and chemical analyses of leachates may help understanding the formation of alpine fissures and their place in alpine metamorphism.

First, the use of K/Na ratio of fluids buffered by adularia and albite is discussed in terms of geothermometry. Knowing the temperature determined in this way, pressure can be estimated through microthermometry results, using the isochores.

These PT data are used then to estimate depths and to discuss the origin of the fluids.

Use of K/Na Ratio in Fluid Inclusions from Quartz in Equilibrium with Adularia and Albite as a Geothermometer

Equilibrium between two alkali feldspars and an alkali-chloride solution can be written:

$$Orthoclase + NaCl_{aq} \gtrsim Albite + KCl_{aq}$$

The equilibrium conditions are determined by the following set of equations:

$$\mu_{\text{KCl}_{aq}} - \mu_{\text{NaCl}_{aq}} = \mu_{\text{Or}} - \mu_{\text{Ab}},$$

$$\mu_{\text{Or}} \text{ in K-spar} = \mu_{\text{Or}} \text{ in Na-spar},$$

$$\mu_{\text{Ab}} \text{ in K-spar} = \mu_{\text{Ab}} \text{ in Na-spar}.$$

If m_{KC1} and m_{NaC1} are the molality of total KCl and NaCl in the solution at equilibrium with two feldspars, a relation is calculated:

$$\label{eq:linear_line$$

where a_{ic} and a_{iaq} are the activities of components other than K and Na in solid phases and in the solution, respectively.

The last equation can be approximated by:

$$\operatorname{Ln} \frac{m_{KCl}}{m_{NaCl}} \sim f(T, P).$$

This function has been calculated, using the data from ORVILLE (1963) and HEMLEY (1967) (see appendix I). Fig. 6 gives the corresponding curve.

The uncertainty associated with this curve depends on the uncertainty of the experimental data and on the validity of the various approximations:

1. The only available data consist of 3 values at 2 Kbar, 670, 600, 500° C (ORVILLE, 1963) and 3 values at 1 Kbar, 500, 400 and 300° C (HEMLEY, 1967).

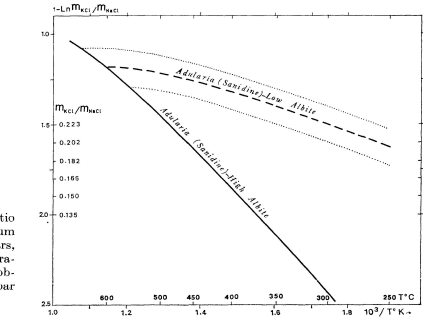


Fig. 6. K/Na atomic ratio in solutions at equilibrium with two alkali feldspars, as a function of temperature. The curves are probably valid in the 1–3 kbar pressure range.

Estimation of the effect of pressure upon the equilibrium is rather difficult, but it seems that, in the pressure range expected for the formation of alpine veins (2 to 3 Kbar), this effect is not very important. Nevertheless, the pressure effect on the equilibrium is a source of possible error.

2. The ratio $\gamma_{\text{KCl}}/\gamma_{\text{NaCl}}$ (stoichiometric activity coefficients of KCl and NaCl) in solution is approximated as a linear function of 1/T (see Appendix I). At low temperatures, the resulting error might be fairly large.

3. The equation of state for a crystalline solution of sanidine has been used in the calculations. The entropy of mixing (and therefore the Gibbs free energy of mixing) of adularia – which is the actual feldspar occuring in the alpine veins – is not known. However, according to WALDBAUM and ROBIE (1971), the differences in these properties between sanidine and adularia must be small.

4. Using theoretical values for the solvus is certainly better than taking actual analyses of coexisting feldspars in explored veins, for those minerals may have re-equilibrated after their formation.

5. The sources of error resulting from neglecting the components other than K and Na must be discussed accurately. These components are mainly divalent cations (Ca²⁺, Mg²⁺, Fe²⁺ and Fe³⁺...), anions such as SO_4^{2-} , HCO₃⁻ and various undissociated complexes.

Analyses of fluid inclusions show that concentrations of Ca^{2+} , Mg^{2+} and SO_4^{2-} are quite low in solutions, and it is probably the same for other components. Thus, it is unlikely that these components will significantly disturb the nearly ideal mixing of KCl-NaCl aqueous solution.

In the solid phases calcium changes the activities of the feldspar endmembers. Owing to the very low anorthite content of the albites dealt with, this effect can be neglected.

FOURNIER and TRUESDELL (1973) discussed the effect of calcium upon equilibrium conditions. The general equation they give (eq. 9, p. 1264) is relevant only when the solution is "semi-buffered" by (Na, K, Ca) minerals (adularia-plagioclase for example). As soon as equilibrium is assumed between the solution and two non-calcic alkali feldspars, and even if other calcic minerals are precipitated, calcium is not involved in the equilibrium constant.

The original Na-feldspar in the rocks surrounding the veins in Mont Blanc and Aar massives is usually an oligoclase. But the co-occurrence of albite in those rocks clearly indicates an irreversible and incongruent dissolution of the plagioclase. This Ca-bearing feldspar feeds the solution, yielding solute species which, in turn, can be precipitated as adularia, albite, muscovite, epidote, calcite . . . Of course, our derivations and interpretations assume local equilibrium between solution and new-formed minerals (particularly adularia and albite, which is the only Na-feldspar occurring in the veins) but the same assumption cannot be defended for the system plagioclase-solution (see, for instance, HELGESON 1967, 1968, 1969).

Structural state of albite at the time of growth

There is very little doubt about the nature of K-feldspars equilibrated with hydrothermal solutions in both experiments and natural veins (sanidine-like structural state). On the other hand, there is no evidence clearly indicating the state of Na-feldspars under the same conditions. Synthetizing low-albite requires special conditions, such as high alkalinity, etc. . . ., and many facts show that albite can reasonably be assumed to crystallize in a disordered state during hydrothermal experiments.

Albite crystals in veins are now ordered. Taking in account the similarities (concentrations of solutes in solutions, for instance) between natural growth and hydrothermal experiments, the effect of water upon the structural state of synthetic albite (MARTIN, 1969) and the systematic disordered state of hydrothermally grown K-feldspars, one can expect that natural hydrothermal albite grew first as high albite. Even if ordering is a fast process, the growing albite can be imagined with a more or less ordered core, coated by a disordered rim in equilibrium with the solution (BROWN, W., personal communication).

In order to test this hypothesis, an approximate curve has been calculated for the adularia (sanidine) – low albite – solution equilibrium (see appendix II, fig. a 6). Using both curves of fig. 6 and the example of the cavity Les Courtes I (Mont Blanc Massif) the following results are obtained: — Characteristics of the fluids:

Homogenization temperature, 185° C. Salinity, 7.7% wt (NaCl) for a pure aqueous solution (no CO₂ present). K/Na at = 0.170.

On the adularia-high albite curve this K/Na ratio gives an equilibration temperature of 420° C and, plotting this value on the isochore (LEMMLEIN and KLEVTSOV, 1961), pressure appears to be close to 2850 bars.

Using the Adularia-low albite curve, an equilibration temperature lower than 200° C is obtained (235° C if the maximum probable error in enthalpy measurements of low- and high-albite is taken into account). In those two cases, using the Adularia-low albite curve, pressure would have been either close to vapour pressure or between 500 and 600 bars. Boiling having never been observed for pure aqueous solutions, vapour pressure is excluded. Obviously the other value is much too low and cannot be defended. The low albite-adularia curve also gives a rather low temperature for green schist facies minerals. But results are still worse for pressure.

Radiometric dating of fissure minerals (K-Ar and Rb-Sr) from the Mont Blanc massif (LEUTWEIN et al. 1970) gives a value close to 15 my. Even if this age is a cooling one, it can be used to get a rough idea of the amount of cover over fissure minerals at the time of their growth. CLARK and JÄGER (1969), using geochronologic and heat flow data, evaluated the rate of erosion in the Alps at 0.7 ± 0.3 mm/year i.e. 0.7 km/my. This value means an erosion of 10.5 km overburden, over cavities, in 15 my. If fluid pressure was close to rock pressure (average rock density 2.7) pressure would have been around 2850 bars. The agreement of this pressure with that calculated above using the adularia-high albite-solution geothermometer is remarkable.

In conclusion it seems reasonable to assume that the sodium feldspar at equilibrium with the solution during its growth was disordered. Thus, using the calculated adularia – high albite – solution curve fig. 6, growth temperatures can be calculated from K/Na ratios of leachates. From microthermometry results (homogenization and freezing temperatures) a PT relationship (constant density: isochore) is determined, which in turn, can be used to calculate the pressure.

Temperatures, Pressures and Depths of Formation of Alpine Fissures

Pressure and depth data calculated in this manner are given for a number of alpine cavities (table V). The number was limited by two requirements: feldspars must have been stable in the cavity or nearby in the country rock, and isochores must be known with good accuracy. This is only possible in the case of pure aqueous solutions. Fluids containing more than a small amount

Pelvoux	K/Na	Tempera- ture ¹)	Homogeni- zation Temp. ²)	NaCl W % ²)	Pressure bars	Depth km³)
Alpe d'Huez	0.106	335	183	17	1700	6.3
Mont Blanc						
Trient	0.154	400	170	7.8	2850	10.5
Charlet Straton	0.154	400	195	8	2400	8.9
Tour Noir	0.159	410	215	8.5	$>\!2300$	$> \! 8.5$
Les Courtes I	0.170	420	185	7.7	2850	10.5
II	0.166	415	183	8.0	2800	10.4
III	0.171	420	175	7.3	3000	11.1
Grands Montets	0.159	410	182	7.8	2750	10.2
Rachasses	0.153	400	196	9.4	2400	8.9
Talèfre	0.161	410	190	9.0	2600	9.6
Pierre Joseph I	0.161	41 0	200	9.0	2500	9.3
Les Périades	0.171	420	160	9.2	3200	11.9
Les Mottets	0.162	410	196	8	2500	9.3
Average	0.161	410	178	8	2800	10.4
Aar						
Kammegg	0.158	405	182	4.8	2750	10.2
Tiefengletscher (Gletschhorn	0.191	450	208	9.5	2800	10.4
Gletsch	0.184	440	207	6.7	2800	10.4
Gerstengletscher	0.168	420	187	11.8	2800	10.4
Average central zone	0.179	430	196	8	2800	10.4
Gotthard						
Ri della Fibbia	0.137	> 380	258	5.8	> 1400	> 5.2
Piz Garviel	0.231	>505	252	9.0	> 2700	$>10^{-10}$
Pennines						
Gischihorn	0.186	445	265	4.2	2050	7.6

Table V. Estimated temperature and pressure of formation of some alpine fissures

¹) from K/Na curve fig. 6.

²) from Table I.

³) assuming $P_{\text{fluids}} = P_{\text{solids}}$ and for a rock density of 2.7.

of CO_2 (5 weight %) must be excluded. In addition to the effect on the isochores a high CO_2 content may disturb the equilibrium constant of the system two feldspars – aqueous solutions.

Temperatures are high and show, as do radiometric data (table I), that alpine fissures did not appear very late in alpine metamorphism. This is not surprising. In cavities like Camperio for instance, the succession of minerals began with scapolites. In the Mont Blanc massif temperatures were not easy to estimate using only mineral associations. Data given here show that the process of fissure formation was rather unique in temperature conditions and probably unique also in time of formation. It seems to be the same in the Aar massif, at least in the central zone. Kammegg shows a lower temperature than in the central zone and this was expected from its northern location. In the Gotthard massif Piz Garviel gives the highest recorded temperature $= 505^{\circ}$ C. But many other values listed in tables III and IV indicate a more complicated history of fissures in this area, as well as in the Pennines.

Pressures are well clustered for the Mont Blanc massif. If fluid pressure is taken to be equal to rock pressure, the depth of formation may be estimated assuming a density of 2.7 for the average rock (table V). These calculations give values close to 9 to 10 km. Such a depth is quite reasonable if we consider the thickness of the Nappe de Morcles which was probably over the Mont Blanc massif at the time of formation of the fissures. These data suggest also that fluid pressure was probably very close to rock pressure. Opening of fissures would require such an identity of fluid and solid pressures. If the former is much less than the latter, depth would be much higher which is difficult to admit.

The constancy of pressure and depth (2.5 to 3 Kbar, 9.5 to 11 km) all over the Mont Blanc and Aar massifs indicates that this region was at the same structural level at the time of formation of the fissures. The differences of temperatures, from 400° C in the Mont Blanc and northern part of the Aar, to 450° C in the central zone of Aar, are thus not related to the depth of burying and can be tentatively interpretated by the occurrence of a hot center probably located in the central part of the Pennines.

Gradients at the time of fissure formation calculated from those data are the following:

Average Mc	ont Blanc	$37^{\circ}/{ m km}$
Aar, Kamm	egg	$37^{\circ}/{ m km}$
Central	zone	$42^{\circ}/{ m km}$
Gotthard	Piz Garviel	$< 48^{\circ}/\mathrm{km}$
Pennines	Gischihorn	$55^{\circ}/\mathrm{km}$

These values are average values and it should be kept in mind that the gradients were probably not identical in granites or gneisses and in overlying sediments.

Origin of Fluids

It appears from this work that water is the major component of the fluid phase in alpine veins.

Because the rocks located below the levels investigated here should be quite dehydrated, the origin of the water must be found in overlying and surrounding sediments, or in circulation from the surface. Although the migration of meteoric or connate water has been demonstrated in many places by various authors, mainly in studies on oxygen and hydrogen isotopes, the same process has not been expressed yet for the Alps, as far as we know. If the hypothesis of participation of meteoric water is correct, that means that circulation down to more than 10 km is possible.

Two kinds of occurrences of CO_2 have been recognized.

 CO_2 occurs as an early fluid (Pennines Gotthard, and to a lesser extent, Aar). This early CO_2 was not found in Pelvoux and Mont Blanc massives. Its concentration decreases during the progressive opening of fissures. This indicates a decrease of temperature with time, a fact firmly established by both the succession of minerals and fluid inclusions studies. This early CO_2 could be related to the metamorphism of carbonate bearing rocks.

High concentrations of CO_2 are also found in late fluids, all over the Alps. This CO_2 -rich fluid is generally responsible for the skeletal habit of quartz crystals (sceptres). No decarbonation can be invoked for supplying CO_2 at that time. The origin of a CO_2 -rich fluid is more likely related to the unmixing of a previously homogeneous CO_2 -H₂O mixture as the whole rock-mass was cooling.

APPENDIX I

Sanidine (Adularia) - High Albite-Solution Equilibrium. General Relations

The equilibrium between a KCl-NaCl aqueous solution and two alkali feldspars can be described by the reaction:

$$Or^{Or} + Na^+ \rightleftharpoons Ab^{Ab} + K^+$$
 or $Or^{Or} + NaCl \rightleftharpoons Ab^{Ab} + KCl$.

Or^{or} and Ab^{Ab} represent respectively the orthoclase end-member in the K-rich feldspar and the albite end-member in the Na-rich feldspar.

The equilibrium condition is:

$$\mu_{\rm KCl} - \mu_{\rm NaCl} = \mu_{\rm Or}^{\rm Or} - \mu_{\rm Ab}^{\rm Ab} \tag{1}$$

with

$$\mu_{0r}^{\text{or}} = \mu_{0r}^{\prime} + \text{RT} \ln a_{0r}^{\text{or}}.$$
(2)

$$\mu_{0r} = \mu_{0r} + 10 \pm 2100_{0r}, \qquad (2)$$

$$\mu_{Ab}^{Ab} = \mu_{Ab} + R T \ln a_{Ab}^{Ab}.$$
 (3)

 $\dot{\mu_{0r}}$ and $\dot{\mu_{An}}$ are the chemical potentials (Gibbs free energies) of pure orthoclase and pure albite.

The chemical potentials of KCl (aqueous) and NaCl (aqueous) can be written (see, for instance, DENBIGH, 1968, p. 306):

$$\mu_{\rm KCl} = \mu_{\rm K}^{*} + \mu_{\rm Cl}^{*} + R \, T \, \ln \gamma_{\rm KCl} \, m_{\rm KCl}, \qquad (4)$$

$$\mu_{\text{NaCl}} = \mu_{\text{Na}}^{*} + \mu_{\text{Cl}}^{*} + \text{R T Ln} \gamma_{\text{NaCl}} m_{\text{NaCl}}.$$
 (5)

 μ_i^* is the chemical potential of the species i in the usual reference state. The symbols m_{KCl} and m_{NaCl} refer to the total molality of the electrolytes, and γ_{KCl} and γ_{NaCl} are the stoechiometric activity coefficients of KCl and NaCl in the solution.

Combining equations (2), (3), (4), (5) with (1), and rearranging:

$$\operatorname{Ln}\frac{\mathbf{m}_{\mathrm{KCl}}}{\mathbf{m}_{\mathrm{NaCl}}} - \operatorname{Ln}\frac{\mathbf{a}_{\mathrm{Or}}^{\mathrm{Or}}}{\mathbf{a}_{\mathrm{Ab}}^{\mathrm{Ab}}} = -\frac{\mu_{\mathrm{K}}^{*} - \mu_{\mathrm{Or}}^{*} - (\mu_{\mathrm{Na}}^{*} - \mu_{\mathrm{Ab}}^{*})}{\mathrm{R}\,\mathrm{T}} - \operatorname{Ln}\frac{\gamma_{\mathrm{KCl}}}{\gamma_{\mathrm{NaCl}}} \tag{6}$$

and with

$$\mu_{\rm K} + -\mu_{\rm 0r} - (\mu_{\rm Na}^* - \mu_{\rm Ab}) = (\Delta \,\rm G)^{\rm P, \, T} \,. \tag{7}$$

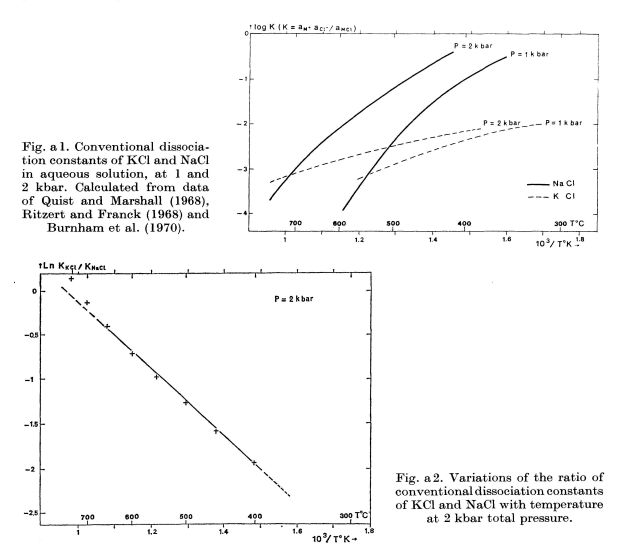
$$Ln \frac{m_{\rm KCl}}{m_{\rm KCl}} - Ln \frac{a_{\rm 0r}^{\rm or}}{m_{\rm Ab}} = -\frac{(\Delta \,\rm G)^{\rm P, \, T}}{m_{\rm Ab}} - Ln \frac{\gamma_{\rm KCl}}{m_{\rm Ab}} \,. \tag{8}$$

$$\operatorname{Ln}\frac{\operatorname{In}_{\mathrm{KCl}}}{\operatorname{m}_{\mathrm{NaCl}}} - \operatorname{Ln}\frac{\operatorname{a}_{\mathrm{Or}}}{\operatorname{a}_{\mathrm{Ab}}^{\mathrm{Ab}}} = -\frac{(2 \text{ G})^{\prime}}{\mathrm{R T}} - \operatorname{Ln}\frac{\gamma_{\mathrm{KCl}}}{\gamma_{\mathrm{NaCl}}}.$$
(8)

Activity coefficients in the solution

The activity coefficients γ_{KCl} and γ_{NaCl} take care not only of the non-ideality of the solution and the choice of the molality scale, but also of the occurrence, in the solution, of undissociated neutral complexes KCl₀ and NaCl₀.

Data from ROBINSON (1961), STERN and ANDERSON (1964), GAMMON, BORCSIK and HOLLAND (1967), WELLMAN (1970), indicate that aqueous solutions of KCl and NaCl (with a constant total molality) mix ideally, or very nearly so, in a wide range of temperature and pressure. Therefore, it can be assumed that, in equation (8), the part of



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 $\gamma_{\rm KCl}/\gamma_{\rm NaCl}$ that does not take in account the partial dissociation of the electrolytes is constant.

Fig. a 1 and a 2 have been calculated from the data of QUIST and MARSHALL (1968) and RITZERT and FRANCK (1968) on the conventional dissociation constants K_{KCl} and K_{NaCl} of KCl and NaCl and the specific volume of water (BURNHAM et al., 1970). These figures show that, at constant pressure (P = 2 kbar), the value of $Ln(K_{Kcl}/K_{NaCl})$ can be approximated by a linear fonction of 1/T, at least in between 300° C and 650° C.

Now, if the variation (ΔC_p) is neglected in the calculation of (ΔG) (eq. 7), the quantity $(\Delta G)/RT$ is a linear fonction of 1/T. Hence, the right side of eq. (8) is also a linear fonction of 1/T and we can write:

$$\operatorname{Ln}\frac{\mathrm{m}_{\mathrm{KCl}}}{\mathrm{m}_{\mathrm{NaCl}}} - \operatorname{Ln}\frac{\mathrm{a}_{\mathrm{Or}}^{\mathrm{Or}}}{\mathrm{a}_{\mathrm{An}}^{\mathrm{An}}} \approx \alpha + \frac{\beta}{\mathrm{T}},\tag{9}$$

where A and B are constants.

Activities of components in the feldspar phases

The chemical potential of the orthoclase end-member in an alkali feldspar can be written:

$$\mu_{\rm or} = \mu_{\rm or,id} + \mu_{\rm or,ex}$$

or
$$\dot{\mu_{\rm or}} + R T \ln \gamma_{\rm or} N_{\rm or} = \dot{\mu_{\rm or}} + R T \ln N_{\rm or} + \mu_{\rm or,ex}.$$
 (10)

 μ_{or} is the Gibbs free energy of pure orthoclase, at the same P and T. N_{or} is the molar fraction K/K + Na in the feldspar, and γ_{or} is the activity coefficient of orthoclase in the same feldspar. From (10) we get

$$\gamma_{\rm Or} = e^{\mu} o_{\rm r, ex/BT}. \tag{11}$$

The excess chemical potential $\mu_{\text{or,ex}}$ is (Gibbs-Duhem equation):

$$\mu_{\rm Or,\,ex} = G_{\rm ex} - N_{\rm Ab} \frac{\partial G_{\rm ex}}{\partial N_{\rm Ab}}.$$
(12)

Assuming a regular assymptric mixture model for the feldspars (Thompson, 1967), the excess free energy G_{ex} is:

$$G_{ex} = A N_{Ab} N_{0r}^2 + B N_{0r} N_{Ab}^2.$$
(13)

A and B are the Margules parameters. They are independent of composition, and are constant at given P and T.

Calculating $\frac{\partial G_{ex}}{\partial N_{Ab}}$ and reporting this value and G_{ex} in (12) gives:

$$\mu_{\rm Or, ex} = (2 \,\mathrm{A} - \mathrm{B}) \,\mathrm{N}_{\rm Ab}^2 + 2 \,(\mathrm{B} - \mathrm{A}) \,\mathrm{N}_{\rm Ab}^3 \tag{14}$$

$$\mu_{Ab ex} = (2 B - A) N_{0r}^2 + 2 (A - B) N_{0r}^3.$$
(15)

Using the equation of state given by WALDBAUM and THOMPSON (1969), we get directly

$$A = 6326.7 + 0.0925 P - 4.6321 T,$$
(16)

$$B = 7671.8 + 0.1121 P - 3.8565 T.$$
(17)

From the same authors, we get also the composition $N_{Or}^{(Ab)}$ and $N_{Ab}^{(Or)}$ of two coexisting feldspars versus T and P, i.e. the solvus values (WALDBAUM and THOMPSON, 1969, table 1).

Activities of the feldspars and $\operatorname{Ln} a_{0r}^{0r}/a_{Ab}^{Ab}$ at 1 and 2 kbar are given below (table a 1 and fig. a 3 and a 4).

and also

T°C	K-rich feldspar		Na-rich	feldspar	$-\mathrm{Ln}\frac{\mathbf{a}_{\mathrm{Or}}^{\mathrm{Or}}}{\mathbf{a}_{\mathrm{Ab}}^{\mathrm{Ab}}}$						
	$\mathbf{N_{or}^{or}}$	$\mathbf{a}_{\mathrm{Or}}^{\mathrm{Or}}$	$\mathbf{N_{Ab}^{Ab}}$	$\mathbf{a_{Ab}^{Ab}}$							
	P = 1 kb										
662	0.333	0.649	0.667	0.892	0.318						
606	0.585	0.713	0.855	0.917	0.253						
512	0.762	0.812	0.935	0.952	0.159						
428	0.866	0.886	0.969	0.974	0.094						
335	0.940	0.946	0.988	0.989	0.045						
241	0.980	0.981	0.997	0.997	0.016						
195	0.990	0.990	0.999	0.999	0.008						
		Р	= 2 kb								
675	0.333	0.649	0.667	0.893	0.318						
648	0.506	0.680	0.807	0.905	0.285						
599	0.628	0.734	0.877	0.925	0.232						
504	0.790	0.831	0.944	0.957	0.142						
391	0.908	0.919	0.980	0.982	0.066						
296	0.963	0.966	0.993	0.993	0.028						
201	0.990	0.990	0.999	0.999	0.008						

Table a 1

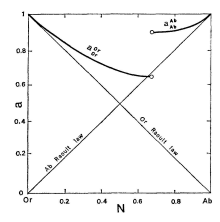


Fig. a3. Activities of orthoclase and albite end members in coexisting alkali feldspars at temperatures up to the critical temperature and pressure of 1 to 2 kbar.

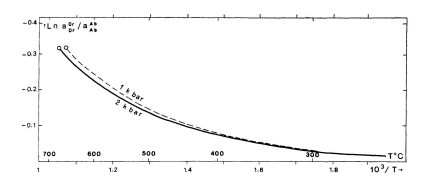


Fig. a4. Negative logarithm of the ratio of activities of orthoclase and albite in coexisting alkali feldspars as a fonction of temperature, at 1 and 2 kbar.

Smoothing the experimental data

The values of the coefficients α and β in equation (9) can be calculated from the experimental data of ORVILLE (1963) and HEMLEY (1967) at 2 and 1 kbar (table a 2).

T°C	10³/T °K	${ m m_{KCl}/m_{NaCl}}$	${f Ln} {m_{ m KCl}/m_{ m NaCl}}$	$- Ln a_{0r}^{0r} / a_{Ab}^{Ab}$	$Ln m_{KCl}/m_{NaCl}$ – $Ln a_{Or}^{Or}/a_{Ab}^{Ab}$
$\begin{array}{c} 670 \\ 600 \\ 500 \\ 500 \\ 400 \\ 300 \end{array}$	$1.060 \\ 1.145 \\ 1.294 \\ 1.294 \\ 1.486 \\ 1.745$	$\begin{array}{c} 0.351^{1})\\ 0.302^{1})\\ 0.227^{1})\\ 0.232^{2})\\ 0.161^{2})\\ 0.086^{2}) \end{array}$	-1.05 1.20 1.48 1.46 1.82 2.45	$\begin{array}{c} 0.31 \\ 0.23 \\ 0.135 \\ 0.145 \\ 0.08 \\ 0.03 \end{array}$	$egin{array}{c} -0.74 \ -0.97 \ -1.345 \ -1.315 \ -1.74 \ -2.42 \end{array}$

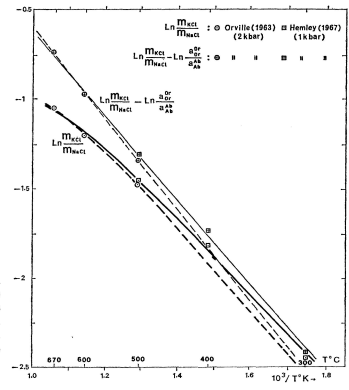
Table a2

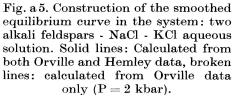
¹) ORVILLE, 1963, 2 kb, mica-free. ²) HEMLEY, 1967, 1 kb, with muscovite.

The values of $\text{Ln}\,m_{\text{KCl}}/m_{\text{NaCl}} - \text{Ln}\,a_{\text{Or}}^{\text{Or}}/a_{\text{Ab}}^{\text{Ab}}$ plotted against 1/T fit a straight line (fig. 5). Actually, two lines can be calculated

- 1. one using ORVILLE's data only (P = 2 kb),
- 2. the other using the data of both Orville and Hemley, assuming that the difference in pressure (1 and 2 kb) yields a variation which can be neglected.

From thoses straight lines, the smoothed equilibrium cuves giving $\text{Ln}\,m_{\text{KCl}}/m_{\text{NaCl}}$ versus 1/T can be easily computed (fig. a5).





APPENDIX II

Adularia - Low Albite - Solution Equilibrium

Equation (8) (appendix I) refers to equilibria between two feldspars and a solution, whatever the structural states of the feldspars are.

Using the subscripts H and L to indicate adularia – high albite and adularia – low albite equilibria, respectively, it can be written, hence:

$$\operatorname{Ln}\left(\frac{m_{\mathrm{KCl}}}{m_{\mathrm{NaCl}}}\right)_{\mathrm{H}} - \operatorname{Ln}\left(\frac{a_{\mathrm{Or}}^{\mathrm{Or}}}{a_{\mathrm{Ab}}^{\mathrm{Ab}}}\right)_{\mathrm{H}} = -\frac{\left(\varDelta \,\mathrm{G}_{\mathrm{H}}\right)^{\mathrm{P,\,T}}}{\mathrm{R\,T}} - \operatorname{Ln}\left(\frac{\gamma_{\mathrm{KCl}}}{\gamma_{\mathrm{NaCl}}}\right)_{\mathrm{H}},\tag{1}$$

$$\operatorname{Ln}\left(\frac{m_{\mathrm{KCl}}}{m_{\mathrm{NaCl}}}\right)_{\mathrm{L}} - \operatorname{Ln}\left(\frac{\mathbf{a}_{\mathrm{Or}}^{\mathrm{Or}}}{\mathbf{a}_{\mathrm{Ab}}^{\mathrm{Ab}}}\right)_{\mathrm{L}} = -\frac{\left(\Delta \operatorname{G}_{\mathrm{L}}\right)^{\mathrm{P, T}}}{\mathrm{R T}} - \operatorname{Ln}\left(\frac{\gamma_{\mathrm{KCl}}}{\gamma_{\mathrm{NaCl}}}\right)_{\mathrm{L}}.$$
(2)

Substracting (1) from (2) and remembering that $\gamma_{\text{KCl}}/\gamma_{\text{NaCl}}$ is constant (at given P and T) with the assumption of ideal mixing of KCl and NaCl in aqueous solutions:

$$\left(\operatorname{Ln}\frac{\mathrm{m}_{\mathrm{KCl}}}{\mathrm{m}_{\mathrm{NaCl}}} - \operatorname{Ln}\frac{\mathrm{a}_{\mathrm{Or}}^{\mathrm{Or}}}{\mathrm{a}_{\mathrm{Ab}}^{\mathrm{Ab}}}\right)_{\mathrm{L}} - \left(\operatorname{Ln}\frac{\mathrm{m}_{\mathrm{KCl}}}{\mathrm{m}_{\mathrm{NaC}}} - \operatorname{Ln}\frac{\mathrm{a}_{\mathrm{Or}}^{\mathrm{Or}}}{\mathrm{a}_{\mathrm{Ab}}^{\mathrm{Ab}}}\right)_{\mathrm{H}} = -\frac{(\varDelta \,\mathrm{G}_{\mathrm{L}})^{\mathrm{P,T}} - (\varDelta \,\mathrm{G}_{\mathrm{H}})^{\mathrm{P,T}}}{\mathrm{R}\,\mathrm{T}}.$$
(3)

We know (equation 7, appendix I) that:

$$\Delta \mathbf{G} = \boldsymbol{\mu}_{\mathbf{K}^{+}}^{*} - \boldsymbol{\mu}_{\mathbf{N}a^{+}}^{*} - (\boldsymbol{\mu}_{\mathbf{Or}}^{-} - \boldsymbol{\mu}_{\mathbf{Ab}}^{-}).$$

Therefore: $(\Delta \mathbf{G}_{\mathbf{L}})^{\mathbf{P}, \mathbf{T}} - (\Delta \mathbf{G}_{\mathbf{H}})^{\mathbf{P}, \mathbf{T}} = \boldsymbol{\mu}_{\mathbf{Ab} \mathbf{L}}^{-} - \boldsymbol{\mu}_{\mathbf{Ab} \mathbf{H}}^{-} = - (\Delta \mathbf{G}_{\mathbf{Ab} \mathbf{L} \to \mathbf{H}})^{\mathbf{P}, \mathbf{T}}.$

And with (3):

$$\left(\operatorname{Ln}\frac{\mathbf{m}_{\mathrm{KCl}}}{\mathbf{m}_{\mathrm{NaCl}}} - \operatorname{Ln}\frac{\mathbf{a}_{\mathrm{Er}}^{\mathrm{Or}}}{\mathbf{a}_{\mathrm{Ab}}^{\mathrm{Ab}}}\right)_{\mathrm{L}} - \left(\operatorname{Ln}\frac{\mathbf{m}_{\mathrm{KCl}}}{\mathbf{m}_{\mathrm{NaCl}}} - \operatorname{Ln}\frac{\mathbf{a}_{\mathrm{Or}}^{\mathrm{Or}}}{\mathbf{a}_{\mathrm{Ab}}^{\mathrm{Ab}}}\right)_{\mathrm{H}} = \frac{\left(\varDelta \operatorname{G}_{\mathrm{Ab} \operatorname{L} \to \mathrm{H}}\right)^{\mathrm{P, T}}}{\operatorname{R} \operatorname{T}}.$$
(4)

Because of the lack of knowledge of the excess entropy of mixing of low albite – microcline solid solutions, the quantity

$$\operatorname{Ln}\left(\frac{a_{Or}^{Or}}{a_{Ab}^{Ab}}\right)_{H} / \left(\frac{a_{Or}^{Or}}{a_{Ab}^{Ab}}\right)_{L}$$

cannot be computed. However, according to WALDBAUM and ROBIE (1971), it seems that the structural state of K-Na feldspars has little effect on their mixing properties. Therefore, it can be approximated:

$$\left(\frac{\mathbf{a}_{Or}^{Or}}{\mathbf{a}_{Ab}^{Ab}} \right)_{\mathbf{H}} / \left(\frac{\mathbf{a}_{Or}^{Or}}{\mathbf{a}_{Ab}^{Ab}} \right)_{\mathbf{L}} \approx 1$$

$$\left(\operatorname{Ln} \frac{\mathbf{m}_{\mathrm{KCl}}}{\mathbf{m}_{\mathrm{NaCl}}} \right)_{\mathbf{L}} - \left(\operatorname{Ln} \frac{\mathbf{m}_{\mathrm{KCl}}}{\mathbf{m}_{\mathrm{NaCl}}} \right)_{\mathbf{H}} = \frac{\left(\Delta \operatorname{G}_{\mathrm{Ab} \operatorname{L} \to \mathbf{H}} \right)^{\mathrm{P. T}}}{\mathrm{R T}}.$$

$$(5)$$

and

If we assume a first-order transition for the reaction Ab L \gtrsim Ab H, the Gibbs free energy of reaction $(\varDelta G_{Ab L \rightarrow H})^{P,T}$ can be computed from the following data:

- * $(\Delta H)^{1,298} = 2.6 \pm 0.3$ KCal (Waldbaum and Robie, 1971).
- * The transition $AbL \gtrsim AbH$ is likely to occur at about 600°C (McConnel and McKie, 1960; Holm and Kleppa, 1968).
- * $(H_T H_{298})$ and $(S_T S_{298})$ are from Robie and Waldbaum (1968).

The entropy of transition, calculated from those data $((\Delta S_{AbL} \rightarrow H)^{1,298} \approx 2.55 \text{ e.u.})$ is significantly lower than the entropy given in ROBIE and WALDBAUM (1968), i.e. 4.47 e.u. However, we get at 700° C a value of about 3.9 e.u., which agrees with the values estimated by HOLM and KLEPPA (1968).

With $(\Delta V_{Ab L \to H}) \approx 0.0086 \text{ cal/bar}$, the following values of $(\Delta G_{Ab L \to H})$ are computed (Table a 3).

Τ°K	$(\Delta \mathrm{H})^{1,\mathrm{T}} \mathrm{cal}$	(⊿ S) ^{1, T} eu	(⊿ G) ^{1, T} cal	(2 G) ^{2 kb, T} /rt
500 600 700 800 900	2837 2954 3072 3189 3306	3.15 3.36 3.53 3.67 3.80	$1262 \\ 938 \\ 601 \\ 253 \\ -114$	$1.28 \\ 0.80 \\ 0.44 \\ 0.17 \\ -0.05$

 $Table \,\, a\, 3$

The uncertainty of $(\Delta G)/RT$ is:

$$\delta\left(\frac{\Delta\,\mathrm{G}}{\mathrm{R}\,\mathrm{T}}\right)\,=\,\frac{1}{\mathrm{R}}\left(\frac{\delta\,(\Delta\,\mathrm{H})}{\mathrm{T}}+\delta\,(\Delta\,\mathrm{S})\right).$$

With $\delta(\Delta H) = 300$ cal, $\delta(\Delta S) = 0.15$ eu, $\delta(\Delta G/RT)$ varies from 0.24 at 900° K to 0.37 at 500° K. Those are large numbers, but the uncertainty of $\Delta G/RT$, and therefore on Ln (m_{KCl}/m_{NaCl}) can be reduced, owing to the following considerations:

* The value of $(\Delta H)^{1,298} = 2.6$ Kcal has been reproduced several times within less than 100 cal (HOLM and KLEPPA, 1968; HOVIS et al., 1970; WALDBAUM and ROBIE, 1971), and the uncertainty of ± 300 cal is certainly overestimated.

* The transition low albite – high albite is unlikely to occur at temperatures lower than 490° (C and probably 550° C) or higher than 670° C.

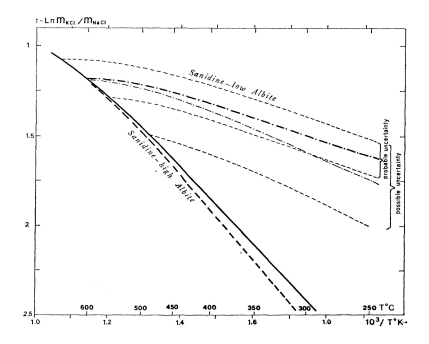


Fig. a6. The Equilibrium curves of the system Sanidine - Low Albite - KCl -NaCl aqueous solution.

Assuming the occurrence of high albite in ORVILLE'S (1963) and HEMLEY'S (1967) experiments (see discussion in text), we can draw the curve of equilibrium: KCl-NaCl aqueous solution – adularia – low albite, using the equation:

$$\mathrm{Ln}\left(\frac{\mathrm{m}_{\mathrm{KCl}}}{\mathrm{m}_{\mathrm{NaCl}}}\right)_{\mathrm{L}} = \mathrm{Ln}\left(\frac{\mathrm{m}_{\mathrm{KCl}}}{\mathrm{m}_{\mathrm{NaCl}}}\right)_{\mathrm{H}} + \frac{\left(\operatorname{\Delta G}_{\mathrm{Ab}\;\mathrm{L}\to\mathrm{H}}\right)^{\mathrm{P},\,\mathrm{T}}}{\mathrm{R}\;\mathrm{T}}.$$

This curve is presented on Fig. a 6 and Fig. 6.

APPENDIX III

List of the Localities

Most of the localities from which the analyzed specimens were obtained are described in the topographical mineralogy of Switzerland («Die Mineralfunde der Schweiz») by STALDER, DE QUERVAIN, NIGGLI and GRAESER, 1973. For bibliographical data refer to the above mentioned book.

In addition to the sample localities (including the coordinates of the map of Switzerland, Landeskarte der Schweiz, $1:25\ 000$ and $1:50\ 000$) the country rock type, the mineral paragenesis in the cleft, and the classification of the paragenesis according to PAR-KER (FG) (see STALDER et al., 1973) are included in the following.

Abreviations of the Mineral Names

$\mathbf{A}\mathbf{b}$	==	Albite	Flu	=	$\mathbf{Fluorite}$
Act	=	Actinolite, asbestiformeous	\mathbf{Gal}	=	Galena
		in the Aar Massif	\mathbf{Hem}	=	Hematite
Ad	=	Adularia	11	\equiv	Ilmenite
\mathbf{An}	=	Anatase	Mon	=	Monazite
Ank	=	Ankerite	\mathbf{Mus}	=	Muscovite
\mathbf{Ap}	=	Apatite	$\mathbf{P}\mathbf{y}$	=	Pyrite
\mathbf{Br}	=	Brookite	Qz	=	Quartz
\mathbf{Ce}	=	Calcite	\mathbf{Rut}	=	\mathbf{Rutile}
Chl	=	Chlorite	\mathbf{Sid}	=	Siderite
Dol	=	Dolomite	\mathbf{Tit}	=	Titanite
$\mathbf{E}\mathbf{p}$	=	Epidote	Tou	=	$\mathbf{Tourmaline}$

Mont Blanc Massif

Trient, approximately 568.0/97.2, alt. 2000 m. Gneisses, Qz, Ad, Chl, Cc.

Le Tour, 566.9/93.7, alt. 3400 m. Granite. Qz, Ad, Chl, Cc, Ab in granite.

Charlet Straton, 566.5/89.6, alt. 3150 m. Granite. Ep, Qz, Ad, Flu, Chl, Cc, Ab in granite.

Le Tour Noir, 568.2/88.3, alt. 3300 m. Granite. Qz, Ab, Mus, Ank, Potassium Feldspar in granite.

Col des Cristaux, 567.1/85.4. Granite. Qz, Ad, Chl, Cc, Ab in granite.

Les Courtes I, 566.8/86.7, alt. 3000 m. Granite. Qz, Ad, Chl, Ab in granite.

Les Courtes II, 566.7/86.6, alt. 3300 m. Granite. Qz, Ad, Chl, Ab in granite.

Les Courtes III, 566.8/86.5, alt. 3550 m. Granite. Qz, Ad, Chl, Ab in granite.

Les Grands Montets, 562.9/88.7, alt. 3150 m. Granite. Qz, Ad, Chl, Cc, Ab in granite.

Les Rachasses, 562.7/90.2, alt. 2500 m. Gneisses. Qz, Chl, Cc.

Bochard, 560.8/88.5. Granite. Qz, Ad, Chl, Flu.

Pointe Isabelle, 567.0/85.3, alt. 3150 m. Granite. Qz, Ad, Chl, Flu, Ab in granite.

Pierre Joseph, 566.2/84.0, alt. 2850 m. Granite. Ep, Qz, Ad, Flu, Chl.

Petites Jorasses, 565.5/81.4, alt. 2900 m. Granite. Ep, Qz, Ad, Chl, Cc.

Les Periades, 562.9/80.5, alt. 3000 m. Granite. Ep, Qz, Ad, Chl, Cc, Ab in granite.

Les Mottets, 559.4/88.2, alt. 1900 m, gallery of the power station. Gneisses. Qz, Chl, Cc.

Aar Massif

- Jeizel, Goppenstein, VS, approximately 625.1/133.6. Amphibolite, prehercynic basement. Qz, Ad, Cc, Ep, Act. FG 2 a.
- Meiggbach, Goppenstein, VS, approximately 624.0/134.9. Amphibolite, prehercynic basement. Qz, Ad, Chl, Ep, Tit. FG 2 a.
- Jägiknubel, Lötschental, VS, "at the bottom of the Jägiknubel, right side of the Langgletscher". Amphibolite, prehercynic basement. Qz, Ep, Act. FG 2 a.
- Gauli, Oberhasli, BE, 659.5/163.1. Chlorite-sericite schists, prehercynic basement. Qz, Chl, Ad. FG 2 a.
- Kammegg, Guttannen, BE, 667.05/168.00. Amphibolite, prehercynic basement. Qz, Ad, Chl, Ep, Act, Tit, Scheelite. FG 2 a.
- Mittagfluh (Guttannen)-Trift, BE, Gallery of the hydro-electric power station Oberhasli. Chlorite-sericite schists, prehercynic basement. Qz, Cc, Chl, Ad, Py. FG 2 a.
- Trift, Oberhasli, BE, approximately 671.0/171.1. Chlorite-sericite schists, prehercynic basement. Qz, Ad, Ab, Chl, Cc, An, Il. FG 2 a.
- Horefelli, Voralp, UR, 682.7/169.6. Chlorite-sericite schists, prehercynic basement, near the granite contact. Qz, Ad, Chl, Flu, Tit. FG 2 a.
- Vorder Ried I, Amsteg, UR, 663.25/179.65. Sericite schists, prehercynic basement. Qz (two generations, scepterquartz), Ank, Mus. FG 1 c.
- Vorder Ried II, Amsteg, UR, 663.25/179.65. Amphibolite, prehercynic basement. Qz, Ab, Ad, Act, Ep. FG 1 d.
- Griesserental, Maderanertal, UR. Sericite schists, prehercynic basement. Qz, Ad, Chl. FG 1 a.

Bändertal, Maderanertal, UR. Sericite schists, prehercynic basement. Qz, Ad, Chl. FG 1 a.

- Staldental, Maderanertal, UR. Sericite schists, prehercynic basement. Qz, Ad, Chl, Il, FG 1 a.
- Reistital, Maderanertal, UR, approximately 704.5/181.65. Sericite schists, prehercynic basement. Qz, Ad, Chl, Il. FG 1 a.

Alpgnofer Platten, Maderanertal, UR. Windgällen volcanics, ignimbrite. Qz, Chl. FG 1 a.

- Ober Sand-Limmernboden, gallery of the hydro-electric power station, Linthal, GL, approximately 716.0/186.5. Tödi granite. Qz, Cc, Py. FG 1 a.
- Alp Cavrein, Val Gronda, GR, 707.5/180.7. Chlorite-sericite schists. Qz, Ad, Chl, An, Br. FG 1 a.
- Hinter Wasen, Fieschergletscher, VS, 654.8/149.7. Central Aar granite. Qz, Amethyst, Chl, Il, Rut. FG 4 b.
- Kleines Lauteraarhorn, Grimsel, BE, approximately 654.0/158.4 Crystalline schists. Qz, Ank, Tetrahedrite, Chalcopyrite. FG 2 b.
- Vorderer Zinggenstock, Grimsel, BE, 663.1/156.3. Central Aar granite. Qz, Chl, Flu, Phenakite. FG 4 a.
- Juchli, Grimsel, BE, approximately 668.0/158.5. Underground discovery of 1942. Grimsel granodiorite. Qz, Chl, Cc, Flu, Py. FG 4 a.

- Sommerloch I, underground, outlet gallery of the power station Oberaar, Grimsel, BE, 668.65/158.90. Grimsel granodiorite. Qz, Chl, Cc, Flu, Milarite, Ap. FG 4 a.
- Sommerloch II, old historical fissure, Grimsel, BE, 668.9/159.25. Grimsel granodiorite. Qz, Chl. FG 4 a.
- Unterer Kessiturm, underground, penstock of the power station Oberaar, Grimsel, BE, 668.22/157.45. Grimsel granodiorite. Qz, Mus, Cc, Sphalerite, Ank, Sid, Mon etc. FG 4 c.
- Gerstengletscher, discovery of 1948, Grimsel, BE, 670.60/160.60. Grimsel granodiorite. Qz, Chl, Py. FG 4 a.
- Gletschhorn, discovery of 1868, Tiefengletscher, UR, 676.3/163.65. Central Aar granite. Qz, Chl, Flu, Gal. FG 4 a.
- Tiefengletscher, UR, 677.3/162.6. Central Aar granite. Qz, Chl. FG 4 a.
- Göschenen I, Gotthard road tunnel N2, safety gallery, meter 105 from the northern portal, UR. Central Aar granite. Qz, Chl, Cc. FG 4 a.
- Göschenen II, Gotthard road tunnel N2, safety gallery, meter 1925 from the northern portal, UR. Central Aar granite. Qz, Chl, Cc, Flu, Ap. FG 4 a.
- Göschenen III, Gotthard road tunnel N2, safety gallery, meter 2137 from the northern portal, UR. Central Aar granite. Qz, Cc, Ap, Chl, Tit. FG 4 a.
- Fedenstock, westside, finder: X. Gnos, UR. Central Aar granite. Qz, Ab, Hem. FG 3 c.
- Furggbach, Baltschiedertal, VS, 635.2/132.65. Sericite schists, prehercynic basement. Qz, Ad, Chl, Hem, Ap. FG 5 d.
- Brig, fissure along the railway track of BLS, VS, 640.3/129.23. Augengneiss, prehercynic basement. Qz, Chl, Ank, Hem. FG 5 c.
- Bel Grat, Belalp, VS, 639.3/139.3. Biotite-gneiss, migmatic complex of the prehercynic basement. Qz, Chl. FG 5.
- Münstigergletscher, Goms, VS, 660.3/151.78. Gneiss, prehercynic basement. Qz, Ad, Chl, Hem, Ap, Chabazite. FG 5 c.
- Bärfel, Oberwald, Goms, VS, 670.80/155.20. Amphibolite of the migmatite zone of St. Niklaus, prehercynic basement. Qz, Ad, Chl, Ap, Tit. FG 5 d.
- Trübtensee, Grimsel, BE, 667.15/156.9. Gneis-Schiefer-Zwischenzone, prehercynic basement. Qz, Ad, Chl, Hem. FG 4 d.
- Gletsch, underground, exploration gallery, VS, 670.78/157.0. Amphibolite of the migmatite zone of Gletsch, prehercynic basement. Qz, Chl, Ad, Tit, Cc, Ep. FG 5 d.
- Rhonegletscher, finder: C. Simmen, 1960, Gletsch, VS. Sericite gneiss of the Gneis-Schiefer-Zwischenzone, prehercynic basement. Qz, Ad, Ap. FG 5 d.
- Unter Sass, Rhonegletscher, VS, 672.1/158.6. Gneis-Schiefer-Zwischenzone, prehercynic basement. Qz, Ad, Chl, Ap, Py. FG 5 d.
- Belvedere, Furka, VS, 673.1/158.63. Main migmatite zone of Gletsch, prehercynic basement. Qz, Sulfosalts, An. FG 5 d.
- Grosstal, Urseren, UR, approximately 683.9/163.8. Gneiss, prehercynic basement. Qz, Ad, Chl, Ap, etc. FG 5 d.
- Göschenen IV, Gotthard road tunnel N2, safety gallery, meter 3565 from the northern portal, UR. Gneiss, prehercynic basement. Qz, Cc, Py, Chl, Laumontite. FG 5 d.
- Mettlen, Andermatt, UR, 689.3/165.7. Permocarboniferous. Qz, Chl. FG 6.
- Russeinbrücke, Disentis, GR, 711.45/175.60. Russeindiorite. Qz, Chl, Hem, Ep. FG 6 a.
- Plattenzüg, Calanda, GR, 754.6/189.95. Spilite of the Taminser crystalline rocks. Qz, Tou, Ab, Chl. FG 15 g.
- Tschengels, Calanda, GR, 751.27/190.2. Rötidolomite, Trias. Qz, Dol, Flu. FG 15 h.
- Bitsch, near Brig. Penstock of the Electra Massa. Prehercynic basement, gneiss. Qz, Chl, Ce, Act. FG 5 d.

Gotthard Massif

- Mooshubel, Mühlebach, VS, 654.65/139.7. Phyllite, prehercynic basement. Qz, Mus, Ank, Sid, Rut. FG 5 b.
- Lamme, Fiesch, VS, approximately 654.75/139.88. Sericite schists, prehercynic basement. Qz, Ab, Cc, Mus, etc. FG 5 b.
- Faulhorn, Oberwald, VS, 672.05/155.50. Sericite schists, perhercynic basement. Qz, Rut, Ce, Barite, Py. FG 5 b.
- Muttengletscher, Gletsch, VS, approximately 675.0/155.8. Gurschen gneiss, prehercynic basement. Qz, Ab, Mus, Chl. FG 5 b.
- Piz Prevat, Gotthard, UR, 691.0/159.2. Paragneiss, prehercynic basement. Qz, Ab, Mus, Chl, Rut, An. FG 7 b.
- Ri della Fibbia, Gotthard, TI, 686.37/156.1. Fibbia granite-gneiss. Qz, Ad, Chl, Mus, Rut, Stilbite. FG 7 d.
- La Fibbia, nord-east side of the mountain, Gotthard, TI. Fibbia granite-gneiss. Qz, Ad, Hem, Mus, Chl. FG 7 d.
- Ri d'Albinasca, Airolo, TI, 687.1/153.4. Amphibolite, Tremolaserie s. l. Qz, Chl, Mus, Ad, Act. etc. FG 7 i.
- Alpe di Fieud, Gotthard, TI, 685.85/153.5. Amphibolite, Tremolaserie s. l. Qz, Chl, Mus, Rut, Act. FG 7 i.
- Grasso di Froda I, Val Canaria, gallery Unteralp-Val Canaria of the power station Ritom, meter 480 from the southern portal, TI. Biotite-gneiss, prehercynic basement. Qz, Ab, Mus, Chl, Ank, Sid, Rut, Py. FG 7 j.

Grasso di Froda II, Val Canaria, gallery Unteralp-Val Canaria of the power station

- Ritom, meter 490 from the southern portal, TI. Biotite-gneiss, prehercynic basement. Qz, Ab, Mus, Chl, Ank, Sid, Rut, Mon, Py. FG 7 j.
- Grasso di Froda III, Val Canaria, gallery Unteralp-Val Canaria of the power station Ritom meter 1450 from the southern portal, TI. Biotite-gneiss, prehercynic basement. Qz, Ad, Ab, Mus, Chl, Ce, Py. FG 7 j.
- Poncioni Negri, Val Canaria, TI, 695.4/157.8. Amphibolite, prehercynic basement. Qz, Amethyst, Ab, Mus, Sid. FG 7 i.
- Alpe Stgegia I, Lukmanier, GR, 705.0/160.0. Crystallina granodiorite. Qz, Ab, Ad, Chl, Tit. FG 7 e.
- Alpe Stgegia II, Lukmanier, GR, 704.72/160.66. Gneissic lense in the hercynic granite. Qz, Ab, Chl, Cc, Tou, Ap. FG 7 e.
- Drual, Platta, Val Medel, GR, 708.15/168.4. Sericite schists with arsenopyrite, prehercynic basement. Qz, Ank, Gal. FG 6 f.
- Piz Miez, Val Medel, GR, 706.7/159.2. Medelser granite. Qz, Ad, Chl, Mus, Rut, Tou. FG 7 e.

Val Casatscha, Val Medel, GR, 708.25/160.00. Medelsergranite. Qz, Act, Mus, Ap. FG 7 e. Piz Garviel, Val Medel, GR, 706.9/161.0. Crystallina granodiorite. Qz, Chl. FG 7 e.

Pennines

- Gh. del Cavagnöö I, Val Bavona, TI, 680.68/145.25. Psammite gneiss of the Lebendun nappe. Qz, Ad, Mus, Chl, Py, An. FG 10 e.
- Gh. del Cavagnöö II, Val Bavona, TI, 680.68/145.25. Psammite gneiss of the Lebendun nappe. Qz, Ad, Ab, Chl, Mus, An, Rut, Py. FG 10 e.
- Gh. del Cavagnöö III, Val Bavona, TI, 680.76/145.30. Psammite gneiss of the Lebendun nappe. Qz, Ad, Ab, Cc, Mus, An, Py, Ank. FG 10 e.

- Gh. del Cavagnöö IV, Val Bavona, TI, 680.62/145.15. Psammite gneiss of the Lebendun nappe. Qz. FG 10 j.
- Cavagnöö, supply gallery of the power station Maggia, close to the entrance of the gallery. Gneiss. Qz, Ad, Mus, Chl, Rut, Ank, Py. FG 10 f.

Poncione di Braga, Alpe Masnaro, Val di Peccia, TI. Gneiss. Qz, Chl, Mus, Hem. FG 10 e.

- Lago Bianco, Val Bavona, TI, approximately 682.3/145.3. Gneiss. Qz, Mus, Cc, Ank. FG 10 f.
- Pizzo dell'Arzo, Val Bavona, TI, 680.65/144.55. Gneiss of the Lebendun nappe. Qz, Ad, Mus, Rut. FG 10 e.
- Fusio, Val Lavizzara, TI, 694.0/145.0. Gneiss. Qz, Ad, Chl, An. FG 10 e.
- Pizzo Madone I, Val Torta, TI, 685.6/149.3. Bündner schists. Qz, Mus, Rut, Pyrrhotite. FG 10 d.

Pizzo Madone II, Val Torta, TI, 685.25/149.13. Bündner schists. Qz, Chl, Tou. FG 10 d. Alpe di Ravina, Airolo, TI. Bündner schists. Qz, Mus, Py. FG 10 d.

Pizzo Gararesc, Val Cassinello, Val Bedretto, TI. Gneiss. Qz, Ab, Ad, Mus, Cc. FG 10 e.

Laghetti, Naret, Val Sambuco, TI, 688.4/147.12. Granite gneiss. Qz, Chl. FG 10 e.

- Naret, Val Sambuco, TI. Amphibolite. Qz, Ab, Chl, Tit, Il. FG 10 h.
- Val di Chironico, Val Leventina, TI, 703.0/140.5. Gneiss. Qz, Ad, Tit. FG 10 h.
- Pizzo Barone, Val di Chironico, Val Leventina, TI, 700.5/140.5. Biotite gneiss. Qz. FG 10 h.
- Iragna, Riviera, TI, 717.9/131.0. Granite gneiss of the Leventina. Qz, Ad, Chl, Ep, Tit. FG 10 g.
- Aquarossa, Val Blenio, TI, 715.5/145.4. Gneiss. Qz, Ad, Ab, Mus, Cc, Tou, Rut, Sid etc. FG 10 e.
- Prugiasco, Val Blenio, TI, 714.72/146.17. Granite gneiss. Qz, Chl, Mus, Ab, Rut, Ap, Ank, Tou. FG 10 f.
- Arvigo, Val Calanca, GR, 729.0/128.9. Granite gneiss. Qz, Ad, Chl, Mus, Cc, Ep, Tit, Zeolites etc. FG 10 g.
- Camperio, Olivone, TI (fissure nr 70 of A. Wagner), 711.70/153.65. Bündner schists, calciferous garnet schists of the lower Stgir serie. Qz, Ank, Cc, Chl, Mus, Tou, Rut, Scapolite etc. FG 10 d.
- Campo Blenio, Val Blenio, TI, 714.8/156.2. Bündner schists. Qz, Ab, Chl, Mus, different iron ores. FG 10 d.

Paltano, Val Bedretto, TI, 677.94/147.06. Bündner schists. Qz, FG 10 i.

Piz Stgir, Greina, GR, 721.54/165.8. Bündner schists. Qz, Cc. FG 8 h.

Val Largia, Lumnezia, GR, 725.98/169.58. Bündner schists. Qz, Cc. FG 8 h.

- Stockchnubel, Gornergletscher, VS, 630.2/92.2. Mica schists from the Stockchnubel-Schuppenzone. Qz, Ad, Cc, Chl, Ep, Laumontite. FG 12 c.
- Fluh, St. Niklaus, Mattertal, VS, approximately 628.2/112.8. Trias quartzite. Qz, Ab, Mus. FG 13 a.
- Simplontunnel I, meter 5342-49 from the northern portal. Berisalschists. Qz, Chl, Mus, Ank. FG 11 f.
- Simplontunnel II, meter 5528-32 from the northern portal. Berisalschists. Qz, Amethyst, Chl. FG 11 f.
- Simplontunnel III, meter 5730-38 from the northern portal. Berisalschists. Qz, Chl, Cc, Hem. FG 11 f.
- Simplontunnel IV, meter 6462-68 from the northern portal. Berisalschists. Qz, Chl, Mus, Ad, Ank. FG 11 f.
- Gibelmatten, Längtal, Binntal, VS. Gneiss of the Monte Leone nappe. Qz. FG 11 d.

Lengenbach, Feld, Binntal, VS, 660.16/135.14. Sugar-grained dolomite. Qz, Hyalophane, Mus, Sulphides, Sulfosalts etc. FG 11 b.

Gischihorn, Kriegalptal, Binntal, VS, 659.65/130.05. Gneiss of the Monte Leone nappe. Qz, Tou, Hem, Rut, Mus, Chernovite. FG 11 d.

Albrunhorn, Binntal, VS, 664.5/135.4. Gneiss of the Monte Leone nappe. Qz, Ep. FG 11 d. Herkumme, Blinnental, VS. Bündner schists. Qz, Cc, Rut, Ank, Sphalerite. FG 11 a.

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