Characterization of granite matrix porosity and pore-space geometry by in situ and laboratory methods

M. Schild,1 S. Siegesmund,1,2 A. Vollbrecht1 and M. Mazurek3

1 Institut für Geologie und Dynamik der Lithosphäre, 37077 Göttingen, Goldschmidtstrasse 03, Germany. E-mail: ssieges@gwdg.de
2 Geol.-Pala¨ont. Institut, Bernoullistrasse 32, 4056 Basel, Switzerland
3 GGWW, Institutes of Geology and of Mineralogy and Petrology, University of Bern, Baltzer-Strasse 1, 3012 Bern, Switzerland

Accepted 2001 January 22. Received 2001 January 19; in original form 2000 September 7

SUMMARY
Most available studies of interconnected matrix porosity of crystalline rocks are based on laboratory investigations; that is, work on samples that have undergone stress relaxation and were affected by drilling and sample preparation. The extrapolation of the results to in situ conditions is therefore associated with considerable uncertainty, and this was the motivation to conduct the ‘in situ Connected Porosity’ experiment at the Grimsel Test Site (Central Swiss Alps). An acrylic resin doped with fluorescent agents was used to impregnate the microporous granitic matrix in situ around an injection borehole, and samples were obtained by overcoring. The 3-D structure of the pore-space, represented by microcracks, was studied by U-stage fluorescence microscopy. Petrophysical methods, including the determination of porosity, permeability and P-wave velocity, were also applied. Investigations were conducted both on samples that were impregnated in situ and on non-impregnated samples, so that natural features could be distinguished from artefacts.

The investigated deformed granites display complex microcrack populations representing a polyphase deformation at varying conditions. The crack population is dominated by open cleavage cracks in mica and grain boundary cracks. The porosity of non-impregnated samples lies slightly above 1 per cent, which is 2–2.5 times higher than the in situ porosity obtained for impregnated samples. Measurements of seismic velocities (Vp) on spherical rock samples as a function of confining pressure, spatial direction and water saturation for both non-impregnated and impregnated samples provide further constraints on the distinction between natural and induced crack types. The main conclusions are that (1) an interconnected network of microcracks exists in the whole granitic matrix, irrespective of the distance to ductile and brittle shear zones, and (2) conventional laboratory methods overestimate the matrix porosity.

Calculations of contaminant transport through fractured media often rely on matrix diffusion as a retardation mechanism.

Key words: contaminant transport, fractured media, Grimsel Test Site, permeability, porosity, seismic anisotropy.

INTRODUCTION
Matrix porosities of crystalline rocks are mostly below 1 per cent and are attributed to open intra- and intergranular microcracks. These are common features of rocks that have undergone deformation, and commonly they display complex composite systems which have formed progressively by a variety of geological processes and under varying conditions over long time spans. In granitic rocks, natural microcracking can be related to thermal contraction during cooling, tectonic stresses during stages of deformation, and stress relaxation during uplift and unroofing (Kowallis & Wang 1983). The lifetime of natural microcracks depends on the geological environment; that is, on the percolation of fluids which may cause healing or sealing of interconnected cracks.

Open (in situ) cracks may be of special interest for the following reasons:

(1) further propagation and coalescence of open cracks at different stress states finally leads to macroscopic fractures, and hence control the mechanical behaviour in the brittle regime (e.g. Hallbauer et al. 1973);
(2) interconnected, open cracks yield pathways for fluid flow and solute diffusion (e.g. France-Lanord et al. 1988);
unconnected, porosity. Rasilainen Knapp (1977), and dominates quantitatively over residual, i.e. is substantially larger than the one measured by Norton & Kauppi et al. to laboratory, and it is still unclear how they can be extrapolated in that all methods provided consistent results. The results were encouraging number of laboratory-based analytical techniques to measure the porosity of crystalline rocks. The results were encouraging among the most important parameters for energy and mass transport through rocks. Thus the frequency and geometry of cracks and pores on all scales, and their type and degree of saturation are relevant characteristics of these reservoir rocks. Several countries (including France, Canada, Sweden, Finland and Switzerland) consider crystalline rocks as hosts for radioactive and industrial waste. Most disposal concepts take advantage of the barrier function of the host rock for flow and transport. The potential of a fractured rock system to retard contaminant migration is greatly increased if the rock matrix adjacent to fractures contains an interconnected network of micropores that is accessible to diffusion (e.g. Neretnieks 1980). Diffusion into the matrix not only dilutes contaminant concentrations in fractures where advection occurs but also provides access to mineral surfaces in the matrix on which contaminants can be sorbed. The combined action of matrix diffusion and sorption on matrix minerals is efficient in retarding contaminant exfiltration and in reducing peak concentrations.

In their classical experimental study, Norton & Knapp (1977) calculated diffusion-accessible porosities on the basis of measured KCl breakthrough curves. These porosities were small compared to the total porosity, leading to the conclusion that a large part of the microscopic pore space of the investigated samples is not connected (‘residual porosity’). However, a wealth of more recent studies (e.g. Hellmuth et al. 1995; SIitari-Kauppi et al. 1998) indicate that the diffusion-accessible porosity is substantially larger than the one measured by Norton & Knapp (1977), and dominates quantitatively over residual, i.e. unconnected, porosity. Rasiainen et al. (1996) compared a large number of laboratory-based analytical techniques to measure the porosity of crystalline rocks. The results were encouraging in that all methods provided consistent results. However, all of these measurements were performed in the laboratory, and it is still unclear how they can be extrapolated to in situ conditions. The quantification of analytical artefacts is one of the major difficulties in characterizing low-porosity rocks. A number of processes may affect porosity measurements in the laboratory, including the following (see Fig. 1):

1. the generation of microcracks during drilling (due to Kirsch and bottom-hole stresses and due to the mechanical impact of drilling);
2. the stress release from lithostatic to atmospheric pressure; and
3. the generation of microcracks during sample preparation (cutting, drying).

In porous rocks, such as shales or sandstones, these effects will be small compared with the in situ porosity of the rocks. In contrast, they may be substantial in low-porosity rocks, such as fresh crystalline rocks. In addition, the connectivity of the microcrack network may also be affected, such that laboratory measurements of permeability and diffusivity may not be representative of in situ conditions.

THE ‘CONNECTED POROSITY’ EXPERIMENT

The main motivation to conduct the Connected Porosity experiment (Frieg et al. 1998) was to reduce and quantify analytical artefacts inherent in all laboratory methods and thereby obtain matrix porosities that more closely represent in situ values. Further objectives included the characterization of the pore-space geometry and pore connectivity. This field experiment was performed by Nagra (Swiss National Cooperative for the Disposal of Radioactive Waste) at the Grimsel Test Site (GTS), located in the Central Swiss Alps at a depth of about 400 m below surface.

The basic principle of the Connected Porosity experiment included the in situ injection of a low-viscosity, fluorescein-doped resin into the rock matrix under near-natural stress conditions from a set of borehole intervals, subsequent polymerization of the resin by a heat pulse, and overcoring of the whole experimental array. Over the time of the injection period, the penetration depth of the resin into the surrounding matrix was of the order of 5 cm (Mori et al. 1996, 1999). Resin-filled pore spaces can be observed and quantified in slices and thin sections of the overcore under UV light and compared with non-impregnated samples from the same locality. Samples from the Connected Porosity experiment are the basis for this paper, and the field information is evaluated and augmented by ancillary laboratory studies. The samples examined in this paper (LS, SZ, HS) were chosen along a profile across a shear zone in the Grimsel granodiorite (see Fig. 2). This shear zone consists of a fine-grained, mica-rich mylonite about 10 cm thick, which was reactivated as a brittle, gouge-bearing fault during regional uplift (Steck 1968). Samples were chosen such that porosity could be monitored as a function of the distance to the shear zone.

By its design, the Connected Porosity experiment fully eliminates artefacts created during sample preparation, and greatly reduces the effects of stress release (points 2 and 3 above). The possible generation of microcracks during drilling (point 1 above) cannot be fully avoided because resin impregnation of the rock matrix did not much exceed 1 diameter of the injection borehole.

GEOLOGICAL SETTING

The Grimsel Test Site (GTS) is situated in the Aar Massif, a basement high in the Helvetic realm of the Alps. The Aar Massif consists of a metasedimentary envelope that was intruded by Hercynian granitoids (320–280 Ma) such as the Central Aare granite and the Grimsel granodiorite; the latter is the host rock of the site investigated. All rocks of the Aar Massif have been affected by Alpine greenschist metamorphism and deformation at about 25 Ma (Dempster 1986). The plutonic rocks were transformed to gneisses. Structures in the Hercynian plutonic rocks are mostly attributed to Alpine deformation (Steck 1968; Marquer & Gapais 1985).
Peak Alpine metamorphic conditions in the vicinity of the GTS are estimated at 400°C/2.5–3 kbar (Choukroune & Gapais 1983; Marquer et al. 1985). The following structural elements can be assigned to this greenschist facies metamorphism and associated ductile deformation: cleavage, mylonitization (including formation of quartz ribbons), mineral stretching lineation, extension fractures and quartz recrystallization. Within this ductile matrix feldspars and allanite (=orthite) show a brittle deformational behaviour.

The formation of brittle structures, which are common in the crystalline rocks at Grimsel, post-dates the ductile deformation and can be attributed to the post-metamorphic regional uplift that is still operative at present with rates of 1–2 mm yr$^{-1}$. In most cases, faults run along pre-existing mylonitic zones and are parallel to metamorphic cleavage (see inset in Fig. 2). The brittle structural elements include both cataclastic fault breccias and discrete fractures. Brittle deformation occurred at significantly lower temperatures and pressures than the preceding ductile deformation.

**TECHNICAL AND EXPERIMENTAL CONCEPTS**

**In situ impregnation**

In order to inject the water-saturated microporosity of the rock matrix, specific properties of the injection resins are required. These include mainly good wetting properties, good miscibility with the in situ pore-water, limited reactivity with pore-water, and low viscosity. Moreover, the resin needs to remain liquid for the duration of the experiment (weeks to months) and polymerize only after heating. A new acrylic resin was developed for the purpose of the Connected Porosity experiment and is described in Frieg et al. (1998). The experimental procedure included the following steps (details in Móri et al. 1999; see also Fig. 2).

1. Drilling of four boreholes (diameter 40 mm) for resin injection into the rock matrix adjacent to a fault.
(2) Drilling of two boreholes (diameter 16 mm) for temperature observation.

(3) Installation of a packer system in the injection boreholes, and injection of acrylic resin into the rock (20 bar overpressure for about 1 month).

(4) Removal of packer and resin from the boreholes, installation of a heating element, and heating (80 °C) for about 8 weeks to polymerize the resin. Temperature sensors were installed at distances of 20–30 cm from the injection boreholes to check whether the temperature exceeded 40 °C (the temperature required for resin polymerization).

(5) Overcoring of the boreholes (core diameter 200 mm).

Sample reference system

All samples (thin sections, cylinders and spheres) were orientated with respect to the macroscopic fabric elements (foliation, lineation) as shown in Fig. 3. As for all other fabric diagrams, the pole figures were plotted on a Schmidt net (lower hemisphere) with x orientated parallel to the lineation; the xy-plane is the foliation plane, and z is coaxial with the foliation pole.

U-stage fluorescence microscopy

The 3-D patterns of microcracks (orientations and frequencies of different types) were measured on standard thin sections using a fluorescence microscope equipped with a Universal-stage (U-stage). In order to record all microcrack orientations, the measurements for each sample were carried out in three mutually perpendicular sections. There is no evidence that a significant number of open cracks were produced during sample preparation, since the cracks show a consistent orientation in different thin sections. Moreover, the orientation and geometry of the microcracks are independent of the sample geometry.

Composite pole figures were computed from the three sections, and errors induced by the overlap of Schmidt net areas were statistically eliminated (for details see Vollbrecht et al. 1991).

During U-stage measurements the microcracks were classified according to their host grain, position (inter- or intragranular, interphase), and state (open, healed, sealed; Schild 1999). For this particular study, however, only open cracks impregnated in situ with acrylic resin and non-impregnated open cracks were considered, in order to discriminate in situ crack porosity from cracks formed during core relaxation and sample preparation. The microcrack fabrics were also quantified indirectly by using petrophysical methods (see below).
Porosity and hydraulic conductivity

Porosity was determined by buoyancy weighting at room temperature. For the investigation of permeability, three orthogonal cylindrical specimens were drilled along the directions of the macroscopic fabric elements (see Fig. 3). The measurements were performed up to a confining pressure of 20 MPa. A pressure transient method using argon gas as a flow medium was used. The gas pressure at the front of the sample was kept constant at 5 MPa. The pressure increase at the back of the samples, measured in a constant volume as a function of time, was used to calculate the permeability (for details see Nover et al. 1995). The permeability was then recalculated to hydraulic conductivity.

Seismic velocities

An experimental determination of the compressional-wave velocities ($V_P$) was performed to quantify the observed microcrack patterns indirectly. The wave velocity measurements ($V_P$) were carried out on spherical samples (diameter = 50 mm ± 0.01 mm) under dry and water-saturated conditions at atmospheric pressure. In addition, the complete patterns of $V_P$ were determined at confining pressures of up to 100 MPa (see Siegesmund et al. 1993). The samples were covered by an epoxy resin film to protect them against the pressure medium.

The measuring system in which the samples were mounted allows a rotation of 360° on the vertical axes and 75° around the horizontal axes. Within this system it is possible to measure the $P$-wave velocity in any direction with the same accuracy. Usually, 132 independent measuring directions were analysed. The $P$ waves were generated and measured by piezoceramic transducers with a 2 MHz resonance frequency. Three different approaches were introduced: (1) velocity measurements at selected confining pressure levels; (2) $V_P$ at atmospheric pressure under dry and saturated (sat) conditions; and (3) analyses of the $V_P$ pattern for comparable impregnated and non-impregnated rock samples.

RESULTS

Petrography and microstructure

All investigated samples show a comparable mineralogical composition. They are mainly composed of K-feldspar (12–24 vol. per cent), plagioclase (29–30 vol. per cent), quartz (27–28 vol. per cent) and biotite (7–11 vol. per cent), with accessory muscovite-sericite, apatite, sphenite, epidote, zircon, chlorite, calcite and opaques.

The foliation which is developed in all samples (see Fig. 4) is defined by compositional banding of alternating dark biotite layers and quartz-rich layers. The macroscopically visible lineation is represented by aggregates (rods) of K-feldspar and plagioclase. The K-feldspar, mainly microcline, exhibits rare twinning after the Ksnalad law, and shows different features of alteration and brittle deformation. The fractures are often filled withfeldspars and recrystallized quartz. Diffuse perthitic exsolution lamellae are visible by different degrees of sericitization. Oligoclase, twinned after the albite law, is also intensely sericitized. The grain size of K-feldspar varies between 100 μm and 1–2 cm, while that of plagioclase is smaller. Quartz occurs as aggregates of equigranular grains, and it is evident from relic structures that they originated by rotation recrystallization. The size of these equigranular quartz grains generally ranges between 100 and 200 μm. Larger grains (up to 1 cm) show undulatory extinction and subgrain boundaries.

The grain size of the mostly recrystallized biotite is approximately equal to that of quartz, while that of secondary muscovite/sericite is smaller. Sometimes the biotite is altered to chlorite, especially along cleavage cracks. The mica (001)-cleavage planes are orientated parallel to the foliation, whereas deviating orientations result from reorientation along phase boundaries (mainly K-feldspar and plagioclase). All samples show a great variation in grain size. The feldspars in particular show extensive features of brittle deformation. The microfractures within larger feldspars are often orientated normal to the lineation.

During greenschist-stage regional metamorphism and deformation, both feldspars behave in a brittle manner and occur as porphyroclasts. Quartz, on the other hand, is largely recrystallized into mica-type ribs and is deformed in a ductile style. The shear-zone sample SZ is a mylonite without macroscopic brittle discontinuities. It has a fine-grained, strongly recrystallized fabric.

Microcrack fabrics

Resin-impregnated pore spaces are related to an interconnected network composed of various crack types, as evident from fluorescence microscopy (Fig. 5). Most frequent are mica cleavage cracks (Figs 5a and b), intergranular cracks within quartz polycrystals (Figs 5c and d), and intragranular cracks in feldspars formed along cleavage planes (Figs 5e to h). More irregularly formed cracks in feldspars probably formed not parallel to crystallographic planes (Figs 5g and h). In addition, few interphase cracks between quartz and feldspars have been detected. The majority of cracks that were not resin-impregnated (and therefore represent artefacts) include a second set of mica cleavage cracks.

With only a few exceptions, the microcracks in mica (biotite and white mica) developed parallel to the (001)-cleavage planes, and a distinction between open and non-impregnated and impregnated cracks is only possible by fluorescence microscopy at high magnification. Mineralizations were not observed within cleavage cracks. Because alteration phenomena or coatings along open non-impregnated cleavage cracks are lacking, it is assumed that they formed mostly during core relaxation or sample preparation. However, it cannot be excluded that individual open cleavage cracks formed earlier but were isolated from the interconnected network. The corresponding crack pole figures are directly correlated with the mica textures and show comparable patterns for the three samples; that is, a concentration close to the foliation normal ($z$) and a tendency to form a girdle around the lineation ($x$; Figs 6a to c). The sample from the ductile shear zone (SZ) shows the strongest preferred orientation, which corresponds to the distinct mylonitic foliation and the related mica textures. The scattering of crack poles from the $z$- to the $x$-direction may be explained by lens-shaped feldspar porphyroclasts which control the shape orientation of the mica flakes. Accordingly, for sample SZ the stronger concentration of crack poles around $z$ can be related to increasing mylonitization, which causes a flattening and grain size reduction of the porphyroclasts.
Figure 4. Rock fabric of the samples (a) HS, (b) SZ and (c) LS, lineation strikes parallel to the longer edge. The foliation is visible from the parallel alignment of mica flakes and elongated quartz ribbons and feldspars. Especially in LS the cataclase of feldspars is obvious; the white feldspar is crossed by grey minerals of quartz.
The intergranular cracks in quartz polycrystals form a well-developed interconnected network which is almost completely impregnated, indicating that they existed in situ. Their lengths range approximately between 100 and 200 μm. Scanning electron microscopy reveals that these cracks display rather rough surfaces. In contrast, non-impregnated open intergranular cracks in quartz are comparatively rare, and show a maximum length of only 100 μm and smaller widths. In addition, few intragranular open cracks were observed. The crack pole figures for open interconnected in situ cracks (Figs 6g to i), which are controlled by the prevailing intergranular network, show for all three samples an orthogonal pattern with a weak tendency of girdles around the reference axes (x, y, z).

The population of impregnated in situ cracks in K-feldspar comprises one or two sets of cleavage cracks and, in addition, irregularly shaped cracks often forming an anastomosing network which usually cuts the cleavage cracks at high angles. The widths of these irregular cracks vary from section to section. For cleavage cracks, it is apparent that in situ opening and impregnation preferentially occurred on cracks that are inclined at a high angle to the foliation (xy-plane). In a few cases, impregnated cracks were observed along K-feldspar phase boundaries, too. The lengths of both cleavage cracks and irregular cracks vary considerably up to about some 100 μm, while the length of phase boundary cracks is strongly dependent on the grain size. In contrast, non-impregnated open cracks in K-feldspar are rare and are mostly represented by isolated intragranular cracks which often lack any connection with grain or phase boundaries (as indicated by 2-D observation in thin sections).

The impregnated in situ cracks in plagioclase display the same features as those in K-feldspar. In plagioclase, intragranular cracks are less frequent, while grain or phase boundary cracks are more frequent than in K-feldspar. The intragranular cracks are dominated by the irregular type described above. In contact zones between plagioclase and quartz polycrystals, the grain boundary cracks of both phases form a common interconnected network. Plagioclase grain boundaries are often decorated by mica flakes that also contain minute intragranular cracks which are connected with this network. As observed for K-feldspar, plagioclase also displays only a few non-impregnated open cracks. Most of them are very small isolated intragranular cracks with irregular curved shapes. Only a few non-impregnated open cracks parallel to twin lamellae have been observed.

Considering impregnated in situ cracks in K-feldspar and plagioclase together, the composite crack pole figures display orthogonal patterns which are comparable with those described for cracks in quartz (Figs 6d to f).
Porosity and hydraulic conductivity

The results of the porosity measurements are given in Fig. 7. As is known, the porosity is very low in crystalline rocks and covers the range between 1 and 1.17 per cent. Most of the porosity is found as crack porosity. The impregnated samples clearly show a distinctly lower porosity, between 0.55 and 0.59 per cent.

The pressure dependence of the permeability/hydraulic conductivity is shown in Fig. 8 and Table 1. The permeability decreases by 50 per cent for HS\textsuperscript{imp} and 1/3 for HS\textsubscript{ni} (where the subscript imp denotes impregnated and the subscript ni denotes non-impregnated) from 5 to 20 MPa (Fig. 8). The anisotropy of the permeability is obvious. Typically, the permeability is low normal to the foliation plane ($z$-direction), while larger

<table>
<thead>
<tr>
<th>Pressure (MPa)</th>
<th>Permeability ($\mu$m$^2$)</th>
<th>Conductivity (m s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HS\textsuperscript{imp}</td>
<td>1.0506</td>
<td>7.89E$-12$</td>
</tr>
<tr>
<td>HS\textsubscript{ni}</td>
<td>0.1189</td>
<td>8.93E$-13$</td>
</tr>
<tr>
<td>HS\textsuperscript{imp}</td>
<td>2.5241</td>
<td>1.90E$-11$</td>
</tr>
<tr>
<td>HS\textsubscript{ni}</td>
<td>0.8947</td>
<td>6.72E$-12$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pressure (MPa)</th>
<th>Permeability ($\mu$m$^2$)</th>
<th>Conductivity (m s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SZ\textsuperscript{imp}</td>
<td>0.7654</td>
<td>5.75E$-12$</td>
</tr>
<tr>
<td>SZ\textsubscript{ni}</td>
<td>0.1743</td>
<td>1.31E$-12$</td>
</tr>
<tr>
<td>SZ\textsuperscript{imp}</td>
<td>0.5783</td>
<td>4.35E$-12$</td>
</tr>
<tr>
<td>SZ\textsubscript{ni}</td>
<td>0.3487</td>
<td>2.62E$-12$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pressure (MPa)</th>
<th>Permeability ($\mu$m$^2$)</th>
<th>Conductivity (m s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LS\textsuperscript{imp}</td>
<td>0.7998</td>
<td>6.01E$-12$</td>
</tr>
<tr>
<td>LS\textsubscript{ni}</td>
<td>0.213</td>
<td>1.60E$-12$</td>
</tr>
<tr>
<td>LS\textsuperscript{imp}</td>
<td>0.5129</td>
<td>3.85E$-12$</td>
</tr>
<tr>
<td>LS\textsubscript{ni}</td>
<td>0.433</td>
<td>3.25E$-12$</td>
</tr>
</tbody>
</table>

Figure 5. (Continued.)
values are detected at higher pressures within the foliation plane, thus indicating that cracks remain open up to pressures of 20 MPa (Table 1).

Comparison of the permeability/hydraulic conductivity for the non-impregnated and the impregnated sample of HS shows lower permeabilities for the impregnated sample. The difference parallel to the lineation (x) is 1.5 μD at 5 MPa. With increasing pressure the difference decreases to 0.3 μD at 20 MPa, while in the direction normal to the foliation (z) it is 0.8 μD at 5 MPa and 0.2 μD at 20 MPa. For the other samples (SZ, LS), the permeability/hydraulic conductivity is generally lower (see Table 1). Furthermore, only in the z-direction is the permeability of the non-impregnated samples greater than that of the impregnated samples. The inverted difference in the x-direction could possibly be caused by larger crystals of feldspar within the cylindrical specimen. This would also affect the preferred orientation of microcracks, which may serve as pathways for the flow medium.

© 2001 RAS. GJI 146, 111–125
Seismic velocities

A typical experimental data set for $V_P$ as a function of pressure at room temperature is shown in Fig. 9. The $P$-wave velocities at atmospheric pressures are higher in saturated rocks than in those under dry conditions, owing to the limited compressibility of water-saturated pores and microcracks compared with air-filled ones (Figs 10 and 11). At higher confining pressures the curves are only slightly different (see Nur & Simmons 1969; Popp 1994). The influence of saturation depends on the matrix porosity. The results show that the $V_P$ values at atmospheric pressure significantly increase for saturated samples. For impregnated rocks the $V_P$ may be higher due to the higher Young’s modulus of the acrylic resin compared with water (see Figs 10d, e and 11d, e).

At low confining pressure, the samples show a pronounced directional dependence (Figs 10a and 11a). $V_{P\text{min}}$ can be observed normal to the foliation (3.22 km s$^{-1}$), whereas $V_{P\text{max}}$ is parallel to the lineation (4.85 km s$^{-1}$) for SZni. This sample also shows a remarkable pressure dependence. The velocities of elastic waves increase with confining pressure, a fact which is usually attributed to progressive microcrack closure (see Figs 10b and 11b). The maximum pressure-dependent increase of $V_P$ can be observed for the $z$-direction. In order to explain the low-pressure phenomena, $\Delta V_P$ (given by the difference between $V_P$ values measured at 100 and at 3 MPa) was calculated for SZni (Fig. 11c). As mentioned above, the corresponding microcrack pattern is dominated by cleavage cracks in muscovite and biotite and phase boundary cracks.

DISCUSSION

Microcrack fabric and texture

All samples show many different types of open in situ microcracks which can be optically discriminated by the injection of the fluorescent acrylic resin. For non-impregnated open cracks, however, it is still unclear whether they existed in situ but were not interconnected with the crack network, or whether they formed during core relaxation or later sample treatment. Microstructural features which are unequivocally indicative of an artificial crack origin, such as open crack segments in the prolongation of impregnated cracks, were not observed. It was possible to identify distinct sets of microcracks forming patterns which strongly depend on the size and shape parameters and textures of their host minerals. For this reason, the geometry of
the crack patterns is closely related to the bulk rock fabrics; that is, it varies significantly from the shear zone into the less deformed wall rocks. The relative frequency and orientation of the various microcrack sets are schematically illustrated for each sample in Fig. 12, which is based on the intensity and the position of maxima in the corresponding crack pole figures (Fig. 6). Because of the weak preferred orientation of the intragranular cracks in quartz and intergranular cracks in quartz polycrystals, only the cracks in feldspars and micas could be represented in this way.

From direct observation of thin sections, it is obvious that in all samples the open \textit{in situ} microcracks and intersections of different types build an interconnected network. The cleavage cracks, in particular, represent favoured planar pathways because of their smooth surface, while rough crack surfaces induce a lower hydraulic aperture. A second subordinate flow direction may be given by intersection lines between prominent crack sets.

Figure 9. Pressure-dependent seismic velocity of the non-impregnated sample SZ and the impregnated sample HS for two selected propagation directions.

Figure 10. Experimentally determined directional dependence of compressional wave velocities [km s$^{-1}$] for the impregnated samples HS, SZ and LS. (a) and (b) $V_p$ as a function of confining pressure; (c) $\Delta V_{P_{100-3}}$ between 100 and 3 MPa confining pressure; (d) and (g) $V_p$ under dry conditions; (e) and (h) $V_p$ under saturated conditions; and (f) and (i) $\Delta V_{P_{\text{sat}-\text{dry}}}$ as a measure of the microcrack-induced anisotropy. Schmidt net, lower hemisphere.

\copyright 2001 RAS, \textit{GJI} 146, 111–125
As mentioned above, the preferred orientation of the (001)-cleavage cracks of the micas is an indicator for the foliation strength. It is most clearly developed in the sample of the immediate shear zone (SZ) and the sample of the footwall (LS).

Porosity
In crystalline rocks, porosity determined in the laboratory yields consistent results, irrespective of the analytical method that is applied (e.g. Rasilainen et al. 1996). However, all laboratory methods suffer from the same systematic errors, namely the unquantified effects of stress relaxation, sampling procedures and sample preparation methods on the results.

Bossart & Mazurek (1991) used Hg-injection and water-loss measurements to measure the porosity of Grimsel granodiorite in the laboratory, and the values fall typically in the range 0.8-1.2 vol per cent. These values are relatively high for crystalline rocks that were not subjected to hydrothermal alteration. However, they can be explained by the abundance of microcrack systems and the substantial albitionization of magmatic plagioclase during greenschist-stage metamorphism. These values are consistent with the data presented in Fig. 7 for samples that were not impregnated in situ.

Samples that were resin-impregnated in situ should result in zero porosity in the laboratory. However, as Fig. 7 shows, this is not the case, and values of about 0.6 vol per cent are measured. This porosity is probably artificial; that is, created by unloading and sample treatment. Thus in situ matrix porosity, calculated as the difference between the values for impregnated and non-impregnated samples, is about 0.4-0.5 vol per cent, which is one-third to one-half of the porosity that is measured on conventional samples in the laboratory. Thus either the pore apertures in the natural microcrack system were enhanced, or new cracks were created due to unloading and sample treatment.

As discussed above, the design of the Connected Porosity experiment avoids a number of possible artefacts that affect porosity measurements in the rock matrix. Whereas the experiment represents a major step towards the characterization of in situ porosity, one uncertainty remains in that possible crack formation and related effects of porosity and pore connectivity by drilling the injection boreholes are as yet unquantified. Presently, there are no rigorous arguments that could quantify these effects. However, the similarity of matrix permeability derived from a large-scale ventilation test that characterizes the matrix well beyond the zone affected by the existence of the tunnel (see below) and the small-scale measurements presented here is a semi-quantitative argument that the possible crack formation during drilling of the injection boreholes does not have a substantial effect on matrix porosity and network structure.

Hydraulic conductivity/permeability
In order to estimate the in situ permeability/hydraulic conductivity, it is necessary to consider the in situ stress field. The maximum horizontal stress (σH) ranges between 18 and 45 MPa, and the minimum horizontal stress (σv) is almost 10 MPa less (Keusen et al. 1989). The lithostatic stress (σL) reaches 8 to 12 MPa, and the hydrostatic pressure (P) is 4 MPa. For average values, the effective stresses (σeff = σ - P) are estimated to be σH = 30 MPa, σv = 19 MPa and σL = 7 MPa. Hence the measurements at a confining pressure of 20 MPa and 30 MPa would give the best approximation of in situ conditions.
The measured hydraulic conductivity at a confining pressure of 20 MPa ranges between $4.9 \times 10^{-12}$ and $1.7 \times 10^{-12}$ m s$^{-1}$ in the z-direction for the non-impregnated sample HS. For SZ$\text{ni}$, the hydraulic conductivity is $1.1 \times 10^{-12}$ and $7.1 \times 10^{-13}$ m s$^{-1}$, and for LS$\text{ni}$, it is $5.6 \times 10^{-14}$ and $1.0 \times 10^{-12}$ m s$^{-1}$.

The dependence of hydraulic conductivity on confining pressure is relatively small. As shown in Fig. 8, the hydraulic conductivity of impregnated samples is less than half the value of non-impregnated samples. The measured values are of the order of $10^{-11}$ m s$^{-1}$. Kull et al. (1993) derived large-scale (dekameters along tunnel) matrix conductivities by measuring the evaporative water discharge from the unfractured tunnel surface, with typical values of $1 \sim 3 \times 10^{-11}$ m s$^{-1}$. Thus, the large-scale in situ measurements are quite consistent with the small-scale laboratory values presented here. It appears that unloading/sample preparation does not critically affect laboratory measurements. Therefore, the application of confining pressure for laboratory tests yields hydraulic conductivities similar to those measured in situ.

**Seismic velocities**

The rapid increase of $V_P$ at low pressures up to 100 MPa as shown in Fig. 9 is associated with the progressive closure of microcracks (e.g. Birch 1961a,b; Brace 1965). At higher pressures, the curves become nearly linear, indicating that intrinsic velocities are approached, reflecting the properties of a nearly crack-free rock volume. The assumption of isotropy is valid only for rocks with a random distribution of minerals, microcracks, pores, etc.—a very special case generally not fully realized in natural rocks. Most rocks are anisotropic when considering physical rock parameters as well as fabrics. Intrinsic elastic properties and anisotropy at higher pressures originate from the elastic properties of the constituent minerals and their crystallographic texture. The difference in velocities measured at high and low (below the crack-closing pressure) confining pressures should be mainly attributed to the contribution of microcracks or pores. Therefore, the spatial dependence of $V_P$ has been studied exclusively on spherical samples, allowing the determination in any direction.

The pressure dependence of the complete $V_P$ distribution and the observed dominant crack populations show a close relationship regarding the seismic anisotropy at low confining pressures. The most important observation is that, for saturated samples, the symmetry of the $V_P$ distribution does not change with pressure in the impregnated nor in the non-impregnated samples. At high pressures, for which most of the cracks are closed, the residual anisotropy is mainly controlled by the texture of biotite and muscovite. The significance and effective influence of the bulk microcrack fabrics on $V_P$ can be best demonstrated by the $\Delta V_P$-patterns. This is clearly indicated by the good correlation between crack pole figures and the corresponding $V_P$ stereograms (compare Figs 6 and 10). Therefore, the influence of the preferred orientations of biotite and muscovite is twofold: they control the intrinsic properties and the orientation of the most significant microcracks, namely open microcracks parallel to the (001)-cleavage planes. Moreover, from comparison of the $V_P$ characteristics of impregnated and non-impregnated samples and their saturation behaviour, it follows that the open in situ cracks are mostly interconnected.

To quantify the influence of the open in situ cracks on the $V_P$ pattern, $\Delta V_{\text{psd}}(\text{SZ}_\text{imp} - \text{SZ}_\text{ni})$ as the difference between $V_P\text{SZ}_\text{imp(dry)}$ and $V_P\text{SZ}_\text{ni(dry)}$ has to be calculated. In this case, the residual $\Delta V_P$ pattern only reflects the contributions of sealed cracks (decorated with the acrylic resin) because the effects of the intrinsic properties, isolated open cracks and cracks originated by stress release after core extraction as well as sample treatment are eliminated (Fig. 13). The $\Delta V_P$ diagram...
the impregnated and non-impregnated sample SZ under saturated conditions. Further explanation see text. in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in situ

in sit
ACKNOWLEDGMENTS

The authors thank NAGRA (Swiss National Cooperative for he disposal of radioactive waste) for supplying samples and background information, as well as financial support. MS thanks the University of Göttingen for a fellowship, and SS the German Science Foundation for a Heisenberg fellowship. A. Mörí (Geotechnical Institute, Bern, Switzerland), two anonymous reviewers and H.-J. Kümpel are acknowledged for useful comments.

REFERENCES


