The Role of Advective Fluid Flow and Diffusion during Localized, Solid-State Dehydration: Söndrum Stenhuggeriet, Halmstad, SW Sweden

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A localized dehydration zone, Söndrum stone quarry, Halmstad, SW Sweden, consists of a central, 1 m wide granitic pegmatoid dyke, on either side of which extends a 2–3 m wide dehydration zone (650–700 °C; 800 MPa; orthopyroxene–clinopyroxene–biotite–amphibole–garnet) overprinting a local migmatized granitic gneiss (amphibole–biotite–garnet). Whole-rock chemistry indicates that dehydration of the granitic gneiss was predominantly isochemical. Exceptions include [Y + heavy rare earth elements (HREE)], Ba, Sr, and F, which are markedly depleted throughout the dehydration zone. Systematic trends in the silicate and fluorapatite mineral chemistry across the dehydration zone include depletion in Fe, (Y + HREE), Na, K, F, and Cl, and enrichment in Mg, Mn, Ca, and Ti. Fluid inclusion chemistry is similar in all three zones and indicates the presence of a fluid containing CO2, NaCl, and H2O components. Water activities in the dehydration zone average 0.36, or \( X_{H_2O} = 0.25 \). All lines of evidence suggest that the formation of the dehydration zone was due to advective transport of a CO2-rich fluid with a minor NaCl brine component originating from a tectonic fracture. Fluid infiltration resulted in the localized partial breakdown of biotite and amphiboles to pyroxenes releasing Ti and Ca, which were partitioned into the remaining biotite and amphibole, as well as uniform depletion in (Y + HREE), Ba, Sr, Cl, and F. At some later stage, H2O-rich fluids (H2O activity > 0.8) gave rise to localized partial melting and the probable injection of a granitic melt into the tectonic fracture, which resulted in the biotite and amphibole recording a diffusion profile for F across the dehydration zone into the granitic gneiss as well as a diffusion profile in Fe, Mn, and Mg for all Fe–Mg silicate minerals within 100 cm of the pegmatoid dyke.

KEY WORDS: charnockite; fluids; CO2; brines; localized dehydration; Söndrum

INTRODUCTION

Solid-state, high-grade dehydration of hornblende- and biotite-bearing rock, by low H2O-activity fluids, to orthopyroxene ± clinopyroxene-bearing rocks on a localized scale (centimetres to metres) is a widespread phenomenon. The two principal dehydration reactions are

\[
\begin{align*}
&KC\text{a}_{2}(Fe, Mg)_4Al_3Si_6O_{22}(OH)_2 + 4 SiO_2 \\
&\text{hornblende quartz} \\
= & Ca(Fe, Mg)Si_2O_6 + 3 (Fe, Mg)SiO_3 + CaAl_2Si_2O_8 \\
&\text{clinopyroxene orthopyroxene plagioclase} \\
+ & KAlSi_3O_8 + H_2O \\
&K-feldspar
\end{align*}
\]

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conditions and

\[ K(Fe, Mg)_{3}AlSi_{3}O_{10}(OH)_{2} + 3 SiO_{2} = 3 (Fe, Mg)SiO_{3} \]

\[ + KAlSi_{3}O_{8} + H_{2}O. \]

K-feldspar

Also referred to, more generally, as ‘incipient’ or ‘arrested charnockitization’, localized, solid-state dehydration zones occur worldwide, most commonly in amphibolite-facies terranes (e.g. inner Bajnaföld, Tasiusarsuaq terrane, SW Greenland—McGregor & Friend, 1992; the Adirondacks, New York—McLelland et al., 1988; Kaapvaal craton, South Africa—van Reenen et al., 1990; Harris et al., 1993; Seward Peninsula, Alaska—Todd & Evans, 1994; Harlov & Förster, 2002a; the Bamble Sector, SE Norway—Knudsen & Lidwin, 1996; NJiland et al., 1998; the Rogaland area, SW Norway—Vander Auwera, 1993; the Quetico subprovince, Ontario, Canada—Pan & Fleet, 1996). However, they also can occur in granitoids (e.g. Natal, South Africa—van den Kerkhof & Grantham, 1999). Localized dehydration zones were first described, and have subsequently been extensively documented, in the Archean amphibolite-facies terranes of southern India (e.g. Friend, 1981; Hansen et al., 1984, 1987; Srikantappa et al., 1985; Kumar & Chacko, 1986; Santosh, 1996, 1992, 2003; Stähle et al., 1987; Harris & Bickle, 1989; Santosh et al., 1990, 1991a, 1991b; Yoshida et al., 1991; Jackson & Santosh, 1992; Harris et al., 1993; Raith & Srikantappa, 1993; Santosh & Wada, 1993a, 1993b; Santosh & Radhika, 1994; Yoshida & Santosh, 1994; Harley & Santosh, 1995; Rajesh et al., 1997; Sathish-Kumar & Santosh, 1998; Dobmeier & Raith, 2000; Kumar, 2004). In some cases, depending on PT conditions and \( \Delta H_{\text{R,O}} \), localized, CO\(_2\)-brine-induced dehydration can be complicated by apparent partial melting (e.g. Burton & O’Nions, 1990; Perchuk et al., 2000).

Sources for low H\(_2\)O activity fluids include magmatic dykes (e.g. Knudsen & Lidwin, 1996; van den Kerkhof & Grantham, 1999), volatile-rich rocks such as marbles (e.g. McLelland et al., 1988; Harlov & Förster, 2002a), or tectonic features such as cracks and/or shear zones, which act as conduits for fluid flow from a distant source (e.g. Hansen et al., 1987; Harris & Bickle, 1989; Radhika & Santosh, 1996). Fluid inclusion data indicate that CO\(_2\), coupled with a minor NaCl–KCl-rich brine fraction, are the principal components in the dehydrating fluid (e.g. Andersen et al., 1997; Tsunogae et al., 2002; Fonarev et al., 2003; Santosh & Tsunogae, 2003; Sarkar et al., 2003). Experimentally, fluids consisting only of NaCl and KCl brines are known to have relatively low wetting angles (see Watson & Brenan, 1987; Gilbert et al., 1998). This potentially would allow them to flow along grain boundaries over relatively large distances (e.g. on the kilometre scale; Newton et al., 1998). In contrast, CO\(_2\)-rich fluids along the CO\(_2\)-H\(_2\)O join (500–1300 MPa; 950–1200°C) are known to have large wetting angles such that they do not form an interconnected network along grain boundaries, but rather isolated pockets of fluid, primarily at three-grain junctions (Watson & Brenan, 1987; Brenan & Watson, 1988; Holness, 1997; Gibert et al., 1998). The inability of CO\(_2\) to flow easily along grain boundaries limits its ability to penetrate deep into the rock, resulting in dehydration zones extending from centimetres to metres only. This observation is supported by \( \delta^{18}O \) isotope studies of incipient charnockite formation in southern India, which suggest that high-grade, CO\(_2\)-rich fluids have a limited infiltration range of \( c. \) 2 m under granulite-grade conditions (Harris & Bickle, 1989). However, there are cases of CO\(_2\)-induced dehydration zones extending as much as 10 m (e.g. van den Kerkhof & Grantham, 1999).

In this study, a comprehensive investigation is made on samples collected along a 1260 cm traverse consisting of a central pegmatoid dyke surrounded by a localized dehydration zone in a migmatized granitic gneiss, all of which are exposed along one of the walls of the Söndrum stone quarry, Halmstad, SW Sweden. The obvious structural and geometric relationship between these three rock types makes this association ideally suited for the study of fluid-induced dehydration in high-grade rocks. Whole-rock and trace element chemistry is investigated, coupled with a detailed documentation of the petrographic fabric, mineralogy, and mineral chemistry of each sample. In addition, a detailed study is made of the fluid inclusions in specific samples along the traverse. Systematic changes in the mineral chemistry, whole-rock chemistry, and fluid inclusion data, both in the dehydration zone and the granitic gneiss, relative to the central pegmatoid dyke, are used to identify the nature of the dehydrating fluids during advectional fluid flow with regard to element depletion and enrichment. These profiles are also used to discern the role of later fluids in the diffusion of specific elements.

**GEOLOGICAL SETTING**

**Regional geology**

Rocks in the SW Swedish granulite region (SGR) consist predominantly of granitic to tonalitic orthogneisses (c. 1660–1700 Ma), with minor intermediate to mafic units (Fig. 1) (Larson et al., 1990; Johansson et al., 1993; Connelly et al., 1996; Söderlund et al., 1999, 2002). The SGR extends along the Kattegatt coast from Hallandsås to the Varberg–Torpa charnockite–granite complex (Fig. 1b). In general, orthopyroxene-clinopyroxene-bearing granitic bodies in the Swedish portion of the Baltic Shield tend to crop out primarily in the SGR.
Of these, the Varberg–Torpa charnockite–granite complex is the principal one. Although the location of some of the larger charnockite bodies is noted in Fig. 1b, a complete, systematic mapping of all known occurrences of orthopyroxene–clinopyroxene-bearing granitic bodies in the SGR has, so far, never been done.

Rocks in the SGR were affected by a major metamorphic event, c. 1420–1460 Ma (Söderlund et al., 1999),
during which migmatization and penetrative deformation of the granitic gneiss occurred throughout the region. At least two generations of granites, quartz monzonites, and syenites (c. 1400 Ma and 1200 Ma) have intruded the SGR. These include the Varberg–Torpa charnockite–granite complex and the Tjärnesjö granite at 1400–1370 Ma (Andersson et al., 1999). The last major tectonothermal event that affected the SGR was the Sveconorwegian orogeny at c. 950–970 Ma (Johansson et al., 2001). This event was characterized by metamorphic conditions that ranged from upper amphibolite to granulite grade (680–750°C and 850–1170 MPa) (Wang & Lindh, 1996), which in some places grades into eclogite-facies metamorphism (Möller, 1998). Subsequent cooling of the SGR during exhumation is compatible with 925–945 Ma U–Pb titanite (Connelly et al., 1996; Wang et al., 1996) and Ar–Ar hornblende geochronology indicating blocking temperatures of 500°C by 930 Ma (Page et al., 1996).

The Söndrum stone quarry

In a local stone quarry in the Söndrum area, two elongated, ‘fracture-controlled’ dehydration zones in the local migmatitic granitic gneiss are well exposed along two opposing walls (Fig. 1c). At present, they represent the only known examples of localized dehydration in the region surrounding Halmstad. The two zones differ distinctly with respect to lithology, mineralogy, and structure. The one along the eastern wall consists of a 0.5-m wide, dark green, orthopyroxene-absent zone rich in clinopyroxene and surrounded on either side by a 0.5-m wide, diffuse orthopyroxene- and clinopyroxene-bearing brown–green overprint of the host, red–pink granitic gneiss. The second zone, which is the focus of this study, occurs opposite on the western wall. It takes the form of a metre-wide pegmatoid dyke of granitic composition from which, on either side, extends an approximately 2.5–3-m wide margin consisting of an orthopyroxene- and clinopyroxene-bearing brown–green dehydration zone. Significantly, this dehydration zone again overprints the migmatization and bANDING of the granitic gneiss (Fig. 2a and b).

The pegmatoid dyke consists primarily of coarse K-feldspar, plagioclase, and quartz crystals as well as graphic intergrowths of feldspar and quartz (Fig. 2c). Clinopyroxene laths, up to 3 cm long, often with a thin black margin of amphibole, are present along with large amphibole crystals, also up to 3 cm long. In other parts of the pegmatoid dyke, the K-feldspar megacrysts have been recrystallized into a granular, relatively fine-grained mass of microcline.

Both the dehydration zone and the surrounding granitic gneiss have identical migmatized gneissic textures and, with the exception of orthopyroxene, clinopyroxene, biotite, and amphibole, have identical modal mineralogies (quartz–plagioclase–K-feldspar–garnet–magnetite–ilmenite–pyrrhotite–pyrite–chalcopyrite–graphite–fluorapatite–allanite–zircon). Gneissic bANDING is defined by the Fe–Mg silicate minerals. The one difference is that, in the dehydration zone, specific grains of hornblende and biotite are completely replaced by orthopyroxene and clinopyroxene. However, both the orthopyroxene and clinopyroxene show some sporadic, partial rehydration in the form of a scattering of relatively small biotite and amphibole crystals along the pyroxene grain rim. Large grains of allanite, surrounded by a spider’s web of radiating cracks (Fig. 2d), typically visible to the naked eye, are present both in the dehydration zone as well as in the surrounding granitic gneiss. The contact between the pegmatoid dyke and the dehydration zone is relatively sharp, in contrast to the contact between the dehydration zone and the amphibolite-facies gneiss, which is more patchy over c. 20 cm.

Zircon, from the dehydration zone and granitic gneiss, exhibits a pronounced oscillatory growth zoning, with U-poor cores and a series of U-rich rims. Both the core and inner rim of these zircons yield an ion microprobe age of c. 1670 Ma, which dates the regional crystallization of the granitic precursor that was migmatized to a granitic gneiss during regional metamorphism at c. 1420–1460 Ma (Söderlund et al., 2002). Ion-probe dating of unzoned zircons from the pegmatoid dyke yields an age of c. 1400 Ma, which dates the formation of the pegmatoid dyke. Zircons from the dehydration zone have an additional thick rim, which has also been dated at c. 1400 Ma, thus confirming that the formation of the pegmatoid dyke and dehydration zone were concurrent (Johansson, 1998; Rimsa et al., 2003). Very thin, outer rims on the zircons, from both the dehydration zone and granitic gneiss, were formed during the c. 970 Ma Sveconorwegian orogeny.

ANALYTICAL TECHNIQUES

Sample selection

The goal of this project was to undertake a systematic study of petrographic relationships and mineral chemistry across apparent metamorphic grade on a localized scale. This required selecting samples of the gneiss at a minimum of 15 cm intervals (and multiples thereof) along a traverse outwards from the approximate centre of the pegmatoid dyke across the dehydration zone deep into the granitic gneiss (see Fig. 2b). A total of 34 samples were studied. These included three from the pegmatoid dyke (first 0–40 cm), 19 from the dehydration zone (next 40–320 cm), and 12 from the granitic gneiss (last 320–1260 cm). The distance (in centimetres) of a particular sample from the approximate centre of the pegmatoid dyke is noted in the sample designation by the last number.
Whole-rock analysis
The samples were analysed at CNRS-CRPG, Service d’Analyse des Roches et des Minéraux (SARM) in Nancy, France. The analytical procedure has been described in detail by Carignan et al. (2001). Rock samples (3–4 kg) from each of the locations shown in Fig. 2b were powdered and fused in Pt crucibles along with LiBO$_2$ at 980°C in an automatic tunnel oven. Major and trace elements were measured using, respectively, inductively coupled plasma atomic emission spectrometry (ICP-AES) and flow injection inductively coupled plasma mass spectrometry (ICP-MS). A blank and five international geostandards were used (Govindaraju, 1994). Analysis for fluorine was performed using ion-selective electrodes. Total H$_2$O and CO$_2$ were determined by combustion IR detection (Förster et al., 1999). Whole-rock analyses of the pegmatoid, dehydration zone, and granitic gneiss are presented in Table A1, contained in Electronic Appendix 1, which may be downloaded from the Journal of Petrology website at http://www.petrology.oupjournals.org.
LA-ICP-MS (laser ablation–inductively coupled plasma mass spectrometry)

The REE abundances in garnet were determined at the Department of Geology, Gothenburg University, Sweden, using a CETAC LSX200 laser ablation sampler attached to a Hewlett Packard 4500 quadrupole ICP-MS system. Analytical procedures have been described by Bi et al. (2001). A 25 µm diameter UV laser was operated in 4 Hz pulsed mode for about 90 s on each spot, with 30 s of background counting, before activation of the laser. The time-resolved spectra were inspected, spikes caused by surface contamination or inclusions were avoided, and three time intervals were chosen for each spot to evaluate precision. 30Si was used as the internal standard and the NIST 612 glass as the concentration standard for the REE.

PIXE (proton-induced X-ray emission)

Nuclear microprobe analyses were undertaken at the Lund nuclear microprobe, which is linked to a 3 MeV single-ended accelerator (Johansson & Campbell, 1988). Analyses were carried out with either 2–55 or 2–82 MeV protons. For detection of X-rays a 50 mm², Si(Li) detector with a resolution of less than 155 eV was employed. The detector was equipped with a Mylar filter of either 425 or 775 µm thickness to suppress intense low-energy X-ray lines from major elements.

Electron microprobe (EMP) analysis

Electron microprobe (EMP) analyses of plagioclase and K-feldspar (Table A2a and b), biotite (Table A3), amphibole (Table A4), garnet (Table A5), clinopyroxene (Table A6), orthopyroxene (Table A7), ilmenite (Table A8), fluorapatite (Table A9), and allanite (Tables A10a and b) were carried out using the CAMECA SX50 and SX100 electron microprobes at the GeoForschungsZentrum Potsdam. (Tables A1–A10 are contained in Electronic Appendix 1 and may be downloaded from the Journal of Petrology website at http://www.petrology.oupjournals.org.) Unless otherwise specified, only mineral cores were analysed and mineral rims were avoided. This was done to obtain mineral compositions that are as close as possible to the peak metamorphic conditions present during formation of the dehydration zone. This assumes that these mineral cores were in equilibrium during the dehydration event. Mineral rims, which normally record post-peak metamorphic processes present during uplift and cooling, were avoided.

Analytical conditions for the silicates and oxides consisted of a 15 kV acceleration voltage, 20 nA beam current, and counting times of 10–30 s depending on the element. Beam spot size was 1 µm for garnet and the pyroxenes, 5 µm for biotite and amphibole, and 15 µm for feldspar, ilmenite,apatite, and allanite. Silicate and oxide standards were taken from both the CAMECA and Smithsonian standard sets (Jarosewich et al., 1980). The CAMECA PAP program was used for matrix correction (Pouchou & Pichoir, 1985). Reintegration of K-feldspar grains with exsolution lamellae of albite involved a systematic succession of 15 µm diameter electron beam spot analyses in a crosswise sweep such that the entire surface of the mineral grain was sampled. Individual analyses (up to 278 per grain) were then averaged to produce a composite analysis (Tables A2a and b). From two to 12 silicate and oxide grains, scattered evenly over the length and breadth of the thin section, were analysed per sample. To ensure that the relative differences observed are correct within microprobe error (+/-1 oxide wt %), silicate and oxide analyses per sample were obtained during one single measuring session including multiple checks of the calibration. The full EMP dataset, representing all individual silicate and oxide analyses, is contained in Electronic Appendix 2 and may be downloaded from the Journal of Petrology website at http://www.petrology.oupjournals.org.

Elements analysed for fluorapatite and allanite included P, Si, Y, La, Ce, Nd, Sm, Gd, Dy, Yb, Ca, Mn, Fe, Sr, Ba, Na, F, and Cl, with counting times of 20–120 s depending on the relative amount of the element. Technique and counting times for the [Y + rare earth elements (REE)] have been described by Harlov & Förster (2002b). Standards for the REE were taken from synthetic REE phosphates prepared by Jarosewich & Boatner (1991), with the remaining standards taken from the Durango fluorapatite and the Smithsonian oxide and silicate standard set (Jarosewich et al., 1980). From five to 15 fluorapatite or allanite grains were analysed per sample in a relatively even sampling over the entire breadth and length of the thin section. If possible, both the core and rim were sampled. No systematic difference was seen. The full EMP dataset, representing all individual fluorapatite and allanite analyses, is contained in Electronic Appendix 3 and may be downloaded from the Journal of Petrology website at http://www.petrology.oupjournals.org.

Microthermometry

Phase transitions in fluid inclusions were investigated by using a Linkam THMS 600 heating–freezing stage cooled with liquid nitrogen (Shepherd, 1981). Samples were prepared as unmounted, doubly polished sections, with an average thickness of 0·2 mm. The stage was calibrated using a set of synthetic fluid inclusion standards. For temperatures around −56·6°C (the melting point of CO2) and around 0°C, the accuracy is better than 0·5°C, whereas for temperatures between 200 and 600°C, the accuracy is better than 5°C.

WHOLE-ROCK CHEMISTRY

The amphibolite-facies granitic gneiss and dehydrated granitic gneiss show nearly identical major element
Compositions. Compared with the granitic gneiss, the dehydration zone is enriched in Ba and Sr and depleted in F (Table A1). Differences in \( \text{Y} + \text{REE} \) between the dehydration zone and the granitic gneiss are more pronounced (Fig. 3). Whereas the light REE (LREE) contents, up to Nd, show similar abundances, the heavy REE (HREE; including Y and Sm) are markedly depleted in the dehydration zone. In general, the pegmatoid dyke has a similar, but slightly more evolved major element composition than either the dehydration zone or the granitic gneiss. However, it is higher in Rb and Sr and lower in \( \text{Y} + \text{REE}, \text{Sc}, \text{and F} \) relative to both the dehydration zone and the granitic gneiss. The one exception is Eu, whose abundance remains unchanged across the entire traverse. The positive Eu anomaly of the feldspar-rich pegmatoid dyke indicates that the oxidation state of Eu must have been 2\(^+\) as opposed to 3\(^+\), thus allowing it to be strongly partitioned into the plagioclase where it substitutes for Ca. Water and CO\(_2\) do not show much variance across the traverse although there is some indication that CO\(_2\) might be slightly depleted in the dehydration zone compared with the granitic gneiss.

MINERAL TEXTURES AND CHEMISTRY

Across the traverse, starting at the centre of the pegmatoid dyke, certain trends in mineral populations and mineral chemistry are seen. All minerals, with the exception of orthopyroxene, clinopyroxene, and magnetite, are ubiquitous, showing the same modal mineralogy both in the dehydration zone as well as in the surrounding granitic gneiss, with the proviso that in the case of those samples containing orthopyroxene and clinopyroxene, modal amounts of hornblende and biotite are relatively less than in samples devoid of pyroxenes.

Feldspars

Feldspar–quartz textures and chemistry in the dehydration zone are identical to those in the surrounding granitic gneiss (Fig. 4a–d). Plagioclase (Table A2a) is enriched in albite \( (X_{ab} \approx 0.83–0.84) \). This albite component is reflected in the K-feldspar \( (X_{Kfs} \approx 0.2–0.3; \text{Table A2b}) \), which shows numerous exsolution lamellae of albite as well as diffusion of Na into the plagioclase along the grain boundary between K-feldspar and plagioclase (Fig. 4b and d). In addition, myrmekite-like lamellae of quartz occur in the plagioclase (Fig. 4e) as well as, more rarely, anti-perthite (Fig. 4f). Plots of plagioclase and K-feldspar compositions from the same thin section on the feldspar ternary indicate disequilibrium between the two phases.

Biotite

Various trends in the biotite chemistry along the traverse are shown in Fig. 5 (see Table A3). In general, per sample, the composition of independent biotite grains is identical, within EMP error, with biotite grains associated with the pyroxenes. Of special note is the Ti content, which is relatively higher in biotite from the dehydration zone compared with biotite in the granitic gneiss (Fig. 5a). The Ti content suddenly drops at the boundary between K-feldspar and plagioclase (Fig. 4b and d). In addition, myrmekite-like lamellae of quartz occur in the plagioclase (Fig. 4e) as well as, more rarely, anti-perthite (Fig. 4f). Plots of plagioclase and K-feldspar compositions from the same thin section on the feldspar ternary indicate disequilibrium between the two phases.

Fig. 3. Whole-rock, chondrite-normalized \( \text{Y} + \text{REE} \) distribution patterns for the pegmatoid dyke, the dehydration zone, and the original granitic gneiss. Chondrite values are taken from Anders & Grevesse (1989).
A sharp increase in the immediate vicinity of the pegmatoid dyke, followed by a gentle dip, and then a gradual increase moving outwards into the granitic gneiss.

The logarithms of the HF and HCl fugacities relative to H₂O, for a fluid in equilibrium with biotite, were estimated using the calibrations of Munoz (1992), which incorporate the updated experimental data of Zhu & Sverjensky (1991, 1992). Fluorine and Cl contents in biotite from the dehydration zone, especially in the vicinity of the pegmatoid dyke, display contrasting trends (Fig. 5d–f). Chlorine is uniformly depleted throughout the dehydration zone, whereas F abundances, although showing their greatest depletion at the contact with the pegmatoid dyke, display a gradual increase along the

Fig. 4. BSE images showing a variety of typical K-feldspar–albite–plagioclase–quartz textures in both the dehydration zone (a, b, e) and the granitic gneiss (c, d, f). (b) is an enlargement of the area outlined by the black box in (a); (d) is an enlargement of the area outlined by the black box in (c).
traverse, evening out at 800 cm with the apparent mean F content in biotite from the granitic gneiss.

**Amphibole**

EMP analysis of the amphiboles indicates that they belong to a group of sodic–calcic amphiboles termed taramite, ideally Na(CaNa)Fe$_3$$^{2+}$AlFe$_3$$^{3+}$Si$_2$Al$_2$O$_{22}$(OH)$_2$ according to IMA-approved nomenclature (Leake et al., 1997). Amphibole analyses, given in Table A4, suggest that Ti probably substitutes for Al, whereas K and Na are positioned on the Ca site. Amphibole mineral chemistry along the traverse mirrors that of the biotite (Fig. 6).

**Fig. 5.** Plot of biotite chemistry as a function of distance (cm) along the traverse outwards from the centre of the pegmatoid dyke (0 cm) for TiO$_2$ (a), FeO (b), MnO (c), $f$$_{H_2O}$ relative to $f$$_{HF}$ (d), $f$$_{H_2O}$ relative to $f$$_{HCl}$ (e), and $f$$_{HF}$ relative to $f$$_{HCl}$ (f). Dotted line designates the approximate boundary between the dehydration zone and the granitic gneiss. The first symbol in each plot designates the boundary with the pegmatoid dyke. The logarithms of HF and HCl fugacities relative to H$_2$O for a fluid in equilibrium with biotite were estimated using the calibrations of Munoz (1992), which incorporate the updated experimental data of Zhu & Sverjensky (1991, 1992). Also shown in (d) and (e) are insets of plots of F and Cl, respectively, vs distance (cm).
Fig. 6. Plot of amphibole chemistry as a function of distance (cm) along the traverse outwards from the centre of the pegmatoid dyke for TiO$_2$ (a), FeO (b), MnO (c), CaO (d), Na$_2$O (e), K$_2$O (f), F (g), and Cl (h). Dotted line designates the approximate boundary between the dehydration zone and the granitic gneiss. The first symbol in the plot designates the boundary with the pegmatoid dyke.
Here, amphibole in the dehydration zone has higher Ti and lower Fe contents than in the granitic gneiss (Fig. 6a and b). Manganese (Fig. 6c), as well as Mg (not plotted), shows a sharp increase in the immediate vicinity of the pegmatoid dyke, followed by a shallow dip, and then gradually increases to a constant value deep within the granitic gneiss. Whereas the Ca content in the amphibole is uniformly higher in the dehydration zone than in the granitic gneiss, both Na and K show a sharp depletion in amphibole in the immediate vicinity of the pegmatoid dyke, followed by an intermediate ‘plateau’ value throughout the rest of the dehydration zone, and a gradual increase in content moving along the traverse into the granitic gneiss (Fig. 6d–f). Both F and Cl chemistry mimic that observed in the biotite showing marked depletion in the dehydration zone (Fig. 6g and h). Moving into the granitic gneiss, F displays the same gradual increase seen in the biotite, although more pronounced, again evening out to what is presumably the mean F content in amphibole in the granitic gneiss. In contrast, Cl shows a sudden jump at the boundary with the granitic gneiss, whereas afterwards it approximates a constant value.

**Garnet**

Garnet occurs as relatively small (5–100 μm), anhedral grains with numerous vermicular inclusions of quartz and feldspar. They are commonly in association with the pyroxenes, biotite, and amphibole. There is no change in garnet textures across the dehydration zone into the granitic gneiss. Garnet chemistry across the traverse is relatively consistent, with the exception of Mn, Fe, and (Y + HREE) (Figs 7 and 8; Table A5). Manganese and Mg (not plotted) again show a sharp increase in the immediate vicinity of the pegmatoid dyke, whereas Fe mirrors this increase by showing a relatively sharp decrease. LA-ICP-MS and PIXE analysis of garnet indicate that garnet from the dehydration zone, and rare garnet in the pegmatoid dyke, is depleted in (Y + HREE) compared with garnet from the granitic gneiss (Fig. 8).

**Pyroxenes**

Pyroxenes occur only in the dehydration zone, in addition to rare clinopyroxene in the pegmatoid dyke. Going outwards from the boundary with the pegmatoid dyke, only clinopyroxene is found in the first 80–85 cm. Orthopyroxene appears at this point and is the predominant pyroxene for the next 85–290 cm, with clinopyroxene still present, though sporadically. A second clinopyroxene zone, in which orthopyroxene is again absent, starts at around 290–295 cm and continues to the patchy boundary with the granitic gneiss at about 320–325 cm. Both orthopyroxene and clinopyroxene show no specific relationship with the surrounding minerals, with the exception of random retrograde grains of biotite and amphibole scattered along the pyroxene grain rims.

Clinopyroxene chemistry mirrors that of the garnet, although less dramatically (Fig. 7; Table A6). Manganese and Mg (not plotted) show a moderate increase in the immediate vicinity of the pegmatoid dyke, whereas Fe shows a more dramatic decrease. Iron strongly increases across the width of the dehydration zone as a function of distance from the pegmatoid dyke.

Orthopyroxene chemistry is relatively constant across the width of the dehydration zone, with the exception of samples SD9-120, SD15-210, and SD17-240, which contain two orthopyroxene populations (Table A7). The two populations are distinguished by a difference of about 0.05 in $X_{Mg}$, with some indication that the population richer in Fe is also richer in Mn. Possible faint trends in Mn and Fe, similar to those seen in the clinopyroxene, are also observed in the orthopyroxene, but are within

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**Fig. 7.** Plot of garnet and clinopyroxene chemistry as a function of distance along the traverse outwards from the centre of the pegmatoid dyke for $X_{Fe} = Fe/(Fe + Mg + Mn)$ (a) and $X_{Mn} = Mn/(Fe + Mg + Mn)$ (b), both in a.p.f.u. Dotted line designates the approximate boundary between the dehydration zone and the granitic gneiss. The first symbol in the plot designates the boundary with the pegmatoid dyke.
The orthopyroxene is also characterized by extremely low Al contents (mean Al₂O₃ content is 0.4 oxide wt %; Table A7).

Fe–Ti oxides, sulphides, and graphite

Oxide minerals also show indications of certain systematic changes along the traverse. Within the first 100 cm of the dehydration zone outwards from the pegmatoid dyke, ilmenite is the only Fe–Ti oxide. Chemically, the ilmenite is more enriched in Mn in the vicinity (~100 cm) of the pegmatoid dyke than further away (Table A8). It contains a moderate hematite component (4–8 oxide wt % Fe₂O₃) and, subsequently, no hematite exsolution lamellae. Modal amounts of ilmenite per sample are constant throughout the traverse and show no particular relationship with respect to the biotite.

Magnetite first appears at ~100 cm. A minority of the magnetite grains contain trellis-like exsolution lamellae of ilmenite. These tend to be very fine and numerous in magnetite grains from the dehydration zone (Fig. 9a) whereas in the granitic gneiss, these lamellae tend to coarsen (Fig. 9b). Compositioanally, the magnetite is near end-member, with relatively minor amounts (<1 oxide wt %) of Cr, Mn, Ti, and Mg. Ilmenite exsolution lamellae in the magnetite–ilmenite grains are enriched in Mn compared with individual ilmenite grains (Table A8). Reintegration of the magnetite–ilmenite grains indicates an original ulvöspinel component of about 20 ± 3%, or X₁₃₃ = 0.15 ± 0.03 (n = 20).

Graphite, chalcopyrite, pyrrhotite, and late-stage pyrite occur sporadically in both the dehydration zone...
as well as in the granitic gneiss. Molybdenite is a rare accessory phase in the pegmatoid dyke.

**Fluorapatite**

Fluorapatite is ubiquitous throughout the dehydration zone and the granitic gneiss and does not contain monazite or xenotime inclusions or rim grains (Table A9; Fig. 10). LREE abundances are relatively constant across the traverse (Fig. 10a). In contrast, mean Y abundances in the fluorapatite (within EMP error) are lower in the dehydration zone than in the granitic gneiss (Fig. 10b) reflecting what is seen in garnet. Similar to both biotite and amphibole, fluorapatite in the dehydration zone has a relatively lower F content compared with fluorapatite in the granitic gneiss, although the change from lower to higher F is abrupt at the boundary between the two zones (Fig. 10c). In contrast, Cl shows more variation, with some indications of enrichment in the immediate vicinity of the pegmatoid dyke and depletion through the remaining dehydration zone compared with fluorapatite in the granitic gneiss, where it suddenly assumes a higher, constant value (Fig. 10d).

**Allanite**

Allanite forms two morphological populations, both with a similar mineral chemistry, which show no change either in morphology or in relative abundances across the dehydration zone into the granitic gneiss. Population-I consists of small (5–20 μm), relatively homogeneous, anhedral to semi-euhedral grains, which occur in all samples along the traverse (Table A10a). Population-I allanite is commonly associated with fluorapatite, either as discrete grains along the fluorapatite rim or as inclusions. More rarely, allanite is associated with zircon. Typically, the larger Population-II allanite shows faint zoning under back-scattered electron (BSE) imaging, with the brighter rims showing some enrichment in (Y + REE) compared with the darker cores. Population-II consists of large, anhedral, inhomogeneous, mottled grains surrounded by long radiating cracks enriched in LREE, which are visible both in hand sample (Fig. 2d).
and thin section (Table A10b; Fig. 11a). Population-II allanite is generally surrounded by numerous grains of zircon and ilmenite (Fig. 11b).

Over the entire traverse, allanite displays variable degrees of alteration. This alteration is manifested by analytical totals below 100% and the presence of some F. Deviations in analytical totals are probably related to the presence of OH and some trivalent Fe. Locally, some of the allanite has undergone more extensive alteration to parisite-(Ce), Ca(Ce,La,Nd)2(CO3)3F2, which shows up more brightly in the BSE image than unaltered allanite. Parisite-(Ce) is intimately intergrown with precursor allanite, hampering reliable analyses of pure material.

ESTIMATION OF TEMPERATURE, PRESSURE, OXYGEN FUGACITY, AND \( H_2O \) ACTIVITY

Because of the ubiquity of garnet throughout the traverse, temperatures could be estimated using a variety of Fe–Mg exchange thermometers involving garnet and a series of Fe–Mg silicate minerals. These included garnet–orthopyroxene (Lee & Ganguly, 1988), garnet–clinopyroxene (Berman et al., 1995), garnet–biotite (Holdaway et al., 1997), and garnet–hornblende (Graham & Powell, 1984). Temperatures were also estimated using the Fe–Mn garnet–ilmenite exchange thermometer of Pownceby et al. (1991). In the estimation of either temperature or pressure, only mineral core analyses were used. Because of the low Al content in the orthopyroxenes (0.4 oxide wt% \( Al_2O_3 \); Table A7), correction for retrograde Fe–Mg exchange between orthopyroxene and garnet (Pattison et al., 2003) adjusted temperatures upwards, on average, only 10°C, well within both analytical and statistical error of the uncorrected Fe–Mg orthopyroxene–garnet temperatures. Disequilibrium between the plagioclase and K-feldspar ruled out the use of two-feldspar thermometry (e.g. Fuhrman & Lindsley, 1988). Pressures were estimated using the assemblage garnet–orthopyroxene–plagioclase–quartz (Harlov, 2000a) utilizing mineral thermochemical data from Berman (1988). Fe–Mg–Mn–Ca mixing in garnet is assumed to approximate the model of Berman (1990). As Fe–Mg mixing in garnet is presumed to closely approximate ideality, Fe–Mg experimental \( K_{Fe} \) exchange data (e.g. Lee & Ganguly, 1988) then require that Fe–Mg mixing in orthopyroxene must approximate ideality as...
well. Again, because of the low Al$_2$O$_3$ content, effects of Al mixing in orthopyroxene were assumed to be negligible. Ca–Na–K mixing in plagioclase was approximated using the plagioclase mixing model of Newton (1983).

In the dehydration zone, utilizing the mineral chemistry documented in Tables A3–A8, garnet–orthopyroxene thermometry indicates a mean temperature of $624 \pm 23^\circ C$ ($n = 16$), whereas garnet–clinopyroxene thermometry gives a mean temperature of $567 \pm 67^\circ C$ ($n = 31$). Starting at the boundary with the pegmatoid dyke, no gradation in temperature is seen across the traverse for any of the five thermometers utilized, i.e. outside their published error bars (see Table 1).

Assuming that the garnet–orthopyroxene, garnet–clinopyroxene, and garnet–biotite thermometers have the highest blocking temperatures (Berman et al., 1995; Holdaway et al., 1997; Pattison et al., 2003) would suggest

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### Table 1: Temperatures and HF and HCl fugacities relative to H$_2$O

<table>
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<th>Sample</th>
<th>Zone</th>
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<th>Gt–Cpx $T_y$</th>
<th>Gt–Bt $T_z$</th>
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<th>log($f_{H_2O}/f_{HCl}$)</th>
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†Garnet–clinopyroxene thermometer, Berman et al. (1995).
‡Garnet–biotite thermometer, Holdaway et al. (1997).
an approximate temperature for the c. 1400 Ma dehydration event of 650–700°C. This temperature range is also supported by the relatively low Al₂O₃ content in the orthopyroxene, i.e. 0.4 oxide wt % (e.g. Aranovich & Berman, 1997; see also Nijland et al., 1998; Morishita et al., 2003). Subsequently, garnet–orthopyroxene–plagioclase–quartz barometry indicates a mean pressure range of 750 ± 30 to 820 ± 30 MPa (n = 13) or a mean pressure of 785 MPa.

Preservation of element traverses in the mineral chemistry, coupled with the low Al content in the orthopyroxene, argues against overprinting of the mineral cores in the c. 1400 Ma dehydration zone or surrounding granitic gneiss during the c. 900–1000 Ma Sveconorwegian–Grenvillian orogeny (680–750°C; 850–1050 MPa—Söndrum area; Wang & Lindh, 1996). This particular PT range is reflected in orthopyroxene from mafic rocks in the SGR, whose Al₂O₃ content typically range from 1 to 2 wt % (Wang & Lindh, 1996), more than twice that measured for orthopyroxene from the dehydration zone (Table A7). Local evidence for this Sveconorwegian overprint is seen in a nearby granulite-facies metadolerite dyke, where the orthopyroxene has an Al₂O₃ content of 1:3 wt % (Fig. 1c; Table A7). Here, garnet–orthopyroxene and garnet–clinopyroxene thermometry indicate a temperature of 720°C and a subsequent pressure (garnet–orthopyroxene–plagioclase–quartz) of 770 MPa; both in good agreement with the PT range determined by Wang & Lindh (1996).

Oxygen fugacity was estimated using the assemblage ferroelite–magnetite–quartz in those samples containing orthopyroxene (see Harlov, 1992, 2000b). For purposes of internal consistency, Fe–Mg mixing in orthopyroxene was approximated using the mixing model of Sack & Ghiorso (1989), whereas the composition of the magnetite is assumed to represent a near end-member composition. Assuming a mean temperature and pressure of 700°C and 800 MPa, respectively, results in an H₂O activity of 0.36 and subsequent X₃H₂O of 0.25 when the H₂O activity is estimated for individual mineral analyses per sample, despite some scatter, no obvious gradient in the H₂O activity is seen across the dehydration zone (Table 1). If X₃H₂O is calculated as a function of temperature at 800 MPa for reaction (2), using these averaged mineral compositions, this equilibrium can be plotted vs temperature in T–X₃H₂O–X₃CO₂ space (Fig. 12). Also plotted is the granite melting curve at 800 MPa approximated from the experimental data of Ebadi & Johannes (1991). At 700°C and 800 MPa, an X₃H₂O of 0.25 plots well into the stability field for granite, implying that partial melting could not have taken place during the dehydration event and thus supporting the contention that conversion of amphibole and biotite to pyroxene occurred under solid-state conditions.

**FLUID INCLUSIONS**

Fluid inclusions from eight samples along the traverse were studied (see Fig. 2b for sampling sites). These included one from the pegmatoid dyke (SD1-0), five from the dehydration zone (SD3-30, SD4-45, SD8-105, SD15-210, SD22-315), and two from the granitic gneiss.
(SD26-375, SD34A-495). Fluid inclusions were found only in quartz, where they occur as isolated inclusions, in clusters, or in a planar arrangement. The fluid inclusions are classified as (1) aqueous inclusions and (2) CO2-bearing inclusions. Members of these groups occur in all samples throughout the traverse, with relative abundances varying with respect to the different lithologies.

Aqueous inclusions
Two distinct population groups can be distinguished among the aqueous inclusions. These include low-salinity aqueous inclusions and saline aqueous inclusions with a maximum of 18 wt % equivalent NaCl. Solid phases in the fluid inclusions are rare and correspond to accidentally trapped phases or secondary carbonate.

Low-salinity aqueous inclusions
The low-salinity aqueous inclusions are by far the most abundant and occur mostly in healed fractures. This indicates the late introduction of almost pure water into the rock system. These inclusions are often monophase at room temperature, with many inclusions losing their bubbles on freezing. They have densities close to that of ice (0.917 g/cm³) and show homogenization temperatures (Tₜₕ) normally above about 150°C, though a few show Tₜₕ of more than 200°C (Fig. 13). The highest Tₜₕ for pure water is about 320°C. Although monophase inclusions are abundant in most samples, only a few of them produce bubbles on freezing and show subsequent homogenization always below 150°C (Fig. 13).

So-called ‘stretched’ fluids in monophase inclusions remain in tension until bubble nucleation. Ice melting in ‘stretched’ water without a bubble (S + L → L) may take place above 0°C, with the bubble often appearing at or just after ice melting. ‘Non-stretched’ monophase inclusions (higher density) melt at lower temperatures than ‘normal’; that is, along the melting (S + L) line. As a consequence, melting temperatures as low as −7°C are sometimes observed for apparently pure water inclusions.

Saline aqueous inclusions
Saline aqueous inclusions are more abundant in samples with higher-density, carbonic, and aqueous–carbonic inclusions. The saline aqueous inclusions occur in clusters, or sometimes in short trails, which locally, on the micron scale, have the same salinity as the inclusions in the clusters. This suggests the local shearing and retrapping of fluids along new-formed fractures. Larger saline aqueous inclusions in the same healed fracture may have a somewhat higher salinity compared with the smaller inclusions. Furthermore, larger inclusions usually show a lower density, whereas smaller inclusions show a higher density. These observable facts can be explained by the retrapping of immiscible fluids.

Eutectic melting temperatures (Tₑ) are typically about −22°C, indicating that the brines are NaCl-dominated. High-salinity inclusions (XNaCl = 15–18 wt %) have Tₑ <130°C. The high-salinity inclusions also have a lower Tₑ compared with pure water (Fig. 14).

CO2-bearing inclusions
Two groups of CO2-bearing fluid inclusions can be distinguished. These include aqueous–carbonic inclusions (H₂O–NaCl–CO₂) and essentially pure carbonic inclusions.

Aqueous–carbonic inclusions
Although rare, aqueous–carbonic inclusions are present in most of the samples. Some inclusions with ‘primary’ characteristics occur in the dehydration zone (e.g. SD15-210). These are relatively large (10–20 μm) inclusions in clusters, with regular morphologies and water fill degrees of about 0.5 (Fig. 15a). However, in other samples, the degree of fill varies between 0.1 and 0.7. The inclusions occur in relatively non-disturbed micro-domains, between sharp trails of pure water inclusions. The systematic higher density [up to about 0.9 g/cm³ (Tₑ[L = −3°C)] for CO₂ in the aqueous–carbonic inclusions is obvious compared with the non-aqueous carbonic inclusions in the same sample.

Rare observations of ice melting between −4 and −6°C correspond to salinities of 7–9 wt % NaCl. Clathrate hydrate melting at 6–8°C for salinities of 4–8 wt % NaCl was observed in a few of the fluid inclusions.

Carbonic inclusions
Carbonic inclusions often occur in clusters with disordered textures (Fig. 15b). Other inclusions occur in sharp trails (Fig. 15c) or in planar arrangement (open trails). These CO₂ inclusions are younger than H₂O inclusions, which occur in clusters with an irregular morphology. Fluid inclusion morphologies vary from irregular to rounded and negative crystal shapes, and occur primarily in the trails. These textures are indicative of post peak-metamorphic re-equilibration and re-trapping. The inclusions sometimes contain small amounts of visible water (degree of fill <0.1). The carbonic inclusions show extreme density variation, from apparent ‘empty’ inclusions (dark appearance in transmitted light and without visible phase transitions) to high-density fluid inclusions of c. 0.90 g/cm³. Only a few carbonic inclusions show bubbles on cooling, with homogenization occurring mostly in the gas phase. However, the vast majority of the ‘measurable’ inclusions show an approximate critical homogenization; that is, at or just below 31°C, corresponding to a density of about 0.47 g/cm³. This also implies an essentially pure CO₂ composition. Accordingly, the melting temperature is always close to that of
Fig. 13. Microthermometry data for fluid inclusions along the traverse covering the pegmatoid dyke, the dehydration zone, and the granitic gneiss. Horizontal dashed lines signify divisions between the pegmatoid dyke (P), the dehydration zone (D), and the granitic gneiss (G). Frequency distributions are shown for the melting temperature of CO\(_2\) (\(T_{m\text{ CO}_2}\)), homogenization temperature of CO\(_2\) (\(T_{h\text{ CO}_2}\)), ice melting temperatures (\(T_{m \text{ ice}}\)), and homogenization temperatures of aqueous inclusions (\(T_{h \text{ H}_2\text{O}}\)). Pie-diagrams show the estimated relative abundance of carbonic (CO\(_2\)), saline aqueous (\(\text{H}_2\text{O}–\text{NaCl}\)), and pure water inclusions (\(\text{H}_2\text{O}\)). The homogenization temperature for CO\(_2\) (i.e. \(T_{h\text{ CO}_2}\)) may be either to the liquid phase (positive values) or to the vapour phase (negative values). White numbers in the histograms indicate measurements, which are not graphically represented. Vertical dotted lines are as follows: A, \(T_{m \text{ CO}_2}\) for pure CO\(_2\) (\(-35.6^\circ\text{C}\)); B, critical \(T_{h\text{ CO}_2}\) for pure CO\(_2\) (\(31^\circ\text{C}\)); C, \(T_{m \text{ ice}}\) for \(\text{H}_2\text{O}\) (0°C); D, the boundary between monophase (left) and two-phase inclusions (right); E, critical \(T_{h\text{ H}_2\text{O}}\) for pure \(\text{H}_2\text{O}\) (37°C). \(\text{H}_2\text{O}\) melting in the histogram is represented as triple-point melting \(\text{S} + \text{L} + \text{V} \rightarrow \text{L} + \text{V}\) or as metastable melting of ‘stretched’ fluids \(\text{S} + \text{L} \rightarrow \text{L}\).
pure CO₂ (−56·6°C). High CO₂ density fluid inclusions are found sporadically in almost all the samples (Fig. 13). However, the best preserved are found in samples that contain abundant high-salinity fluid inclusions, such as those in the granitic gneiss (e.g. SD34A-495 in Fig. 13), and to a lesser extent, those in the central part of the dehydration zone.

**Major fluid types in the different zones**

The pegmatoid dyke shows the highest abundance of CO₂ inclusions (Fig. 13), which is reflected in the whole-rock analyses (Table A1). The density of these inclusions is generally very low. In addition to the carbonic inclusions, aqueous inclusions also occur but in separate domains. Mixed aqueous–carbonic inclusions are rare.

In the dehydration zone, the relative abundances of aqueous and carbonic inclusions are about equal. The greatest percentage of high-salinity aqueous inclusions occurs about 0–4 m from the contact with the pegmatoid dyke (Figs 13 and 16). On the micrometre scale, the carbonic and high-salinity aqueous inclusions tend to be associated in the same domains. In contrast, pure H₂O inclusions occur in trails, mostly in different domains.

The granitic gneiss samples contain well-developed, aqueous and carbonic inclusions. Mixed H₂O–CO₂ inclusions are also rare here. The sample from deep within the granitic gneiss (SD34A-495) contains relatively many aqueous inclusions of high salinity, which are associated with CO₂. On the other hand, only almost pure water inclusions are found in clusters or in healed fractures in the sample of granitic gneiss that borders the contact with the dehydration zone (SD26-375) (Fig. 13). Here, CO₂ inclusions are almost absent. Some parallel trails of low-density, or ‘empty’ CO₂ inclusions, indicate strong deformation and/or shearing in this zone.

**INTERPRETATION OF MINERAL TEXTURES AND CHEMISTRY**

Feldspar–quartz mineral textures and chemistry in the dehydration zone are identical to those in the surrounding granitic gneiss, implying that the fluids responsible for the partial conversion of biotite and amphibole to orthopyroxene and clinopyroxene via reactions (1) and (2) had no obvious effect on either (Fig. 4a and c; Table A2). Enrichment in Na along plagioclase grain rims in contact with K-feldspar (Fig. 4b and d), apparent myrmekitic intergrowths of quartz and plagioclase (Fig. 4e), as well as mesoperthite (Fig. 4f), all point to some sort of interaction with either a fluid or a fluid-rich melt, probably with an NaCl brine component. This probably occurred during the regional, 1420–1460 Ma migmatization event, as such textures are also found in the granitic gneiss throughout the SGR.
One feature common to the Fe–Mg silicate minerals and ilmenite is the strong partitioning of Mn and Mg into, and Fe out of, these minerals in the first 100–150 cm of the dehydration zone (Figs 5b, c; 6b, c; and 7). Such a phenomenon cannot be interpreted as having resulted from an imposed high oxidation state (e.g. Harlov et al., 1997), as magnetite has been removed and ilmenite has lost Fe. Rather, fluids in the first 100–150 cm of the dehydration zone must have had a high affinity for Fe, influenced by the fluid-rich granitic melt responsible for the pegmatoid dyke, which acted as chemical sink for Fe, such that the Fe was then removed from the general locality. This is supported by the fact that the pegmatoid dyke is low in Fe relative to the granitic gneiss (Table A1). This chemical affinity for Fe was apparently strong enough to influence the chemistry of Fe–Mg silicate minerals c. 100–150 cm into the surrounding dehydration zone, such that the chemical potentials favoured the partitioning of Fe into the CO₂–NaCl–H₂O grain boundary fluids present, which subsequently aided the mobility of Fe towards the granitic melt.

Both biotite and amphibole show relatively uniform enrichment in Ti in the dehydration zone (Figs 5a and 6a), which would appear to indicate that the dehydration event itself was characterized by advective transport. Titanium enrichment in biotite is common in both localized and well as regional dehydration zones; that is, regions that have undergone granulite-facies metamorphism to orthopyroxene-bearing rocks (e.g. Hansen et al., 2002; Harlov & Förster, 2002a). In either case, Ti enrichment appears to be related to reactions (1) and (2), which free up Ti to be absorbed by the remaining biotite and amphibole, thereby stabilizing both minerals against conversion to pyroxenes. This would also explain why enrichment of biotite and amphibole in Ti is directly correlated with the formation of orthopyroxene and clinopyroxene. With respect to the amphiboles, enrichment in Ti certainly cannot be ascribed as a function of increasing temperature across the traverse (e.g. Ernst & Liu, 1998).

Correlation between Ti in biotite and the metamorphic ‘grade’, as it relates to the presence or absence of

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Fig. 15. (a) (SD15-210) A domain with primary inclusions surrounded by strongly modified inclusions and fluid inclusion trails. The inclusions, which are out of focus, are outlined. The three inclusions indicated have water-filling degrees of 0–3–0–5. The CO₂ homogenization (to liquid) is +8°C for the three inclusions; Tm_ice for inclusion 1 is −4–5°C, Tm_clathrate is +8–9°C. (b) (SD15-210) Two decrepitated clusters of CO₂ inclusions with irregular shapes and smaller aqueous inclusions showing bubbles. (c) (SD8-105) Sharp trails of pure CO₂ inclusions. Scale bar represents 25 μm in all photographs.
orthopyroxene, agrees well with experiment (e.g. Patiño-Douce, 1993). Patiño-Douce (1993) demonstrated experimentally that the solubility of Ti in biotite increases with temperature in a strongly non-linear fashion such that up to amphibolite-facies conditions, the effect of Ti on the stability of biotite is very small. However, upon initiation of the dehydration reactions associated with granulite-facies metamorphism and the lower H$_2$O activities that characterize it, Ti in biotite increases markedly, rendering the biotite relatively refractory. Recently, Ti enrichment in biotite, as a function of increasing metamorphic grade, has been empirically calibrated as a geothermometer for high-grade metapelites using extensive natural biotite data (Henry et al., 2005).

In a similar manner, Ca enrichment in the remaining amphiboles in the dehydration zone (Fig. 6d) is also an indicator of advective transport. In the most likely scenario, a portion of this Ca, released during amphibole breakdown, would be taken up by the formation of clinopyroxene. However, because of the paucity of clinopyroxene across a large portion of the dehydration zone, the majority was apparently partitioned back into the remaining amphibole, subsequently forcing out Na and, to a lesser extent, K, such that charge balance was maintained (Fig. 6e and f). Sodium and K were apparently either flushed out of the dehydration zone or, more probably, partitioned into the feldspars, whose high Na and K content allows them to be treated as an infinite reservoir with respect to the absorption of such relatively minor amounts of Na and K. Steep Na and K depletion in amphiboles in the first 100 cm out from the pegmatoid dyke (Fig. 6e and f) appears to mirror the equally dramatic depletion in Fe seen in the Fe–Mg silicates in general. This suggests that they might be coupled in some way, perhaps as a result of increased mobility of a brine component along grain boundaries during either the formation or the crystallization of the granitic melt responsible for the pegmatoid dyke.

**Biotite, amphibole, and fluorapatite halogen chemistry**

In the dehydration zone, all three halogen-bearing minerals (i.e. biotite, amphibole, and fluorapatite) show depletion in F and Cl (Figs 5d–f; 6g, h; and 10c, d). Depletion in Cl is limited to the dehydration zone, with a sharp cut-off at the boundary with the granitic gneiss, thus suggesting that its depletion was also a product of advective transport. This indicates that the Cl was partitioned into the low H$_2$O activity fluid responsible for the dehydration event and subsequently moved out of the dehydration zone via advective transport. What is apparently only a partial depletion in Cl in the first 100 cm out from the pegmatoid dyke, relative to the rest of the dehydration zone, could be explained as being due to the expulsion of Cl during the last phases of crystallization of the granitic melt responsible for the pegmatoid dyke. Such a phenomenon is common in crystallizing, deep-seated melts under similar, granulite-facies conditions (e.g. Markl & Piazolo, 1998).

In contrast, whereas F is uniformly depleted in fluorapatite, with a relatively sharp cut-off at the boundary with the granitic gneiss again indicative of advective transport, the F component in biotite and amphibole shows a more...
gradual transition across this boundary. This behaviour indicates that F depletion in the silicate minerals continued deep into the granitic gneiss, without conversion of biotite and amphibole to pyroxenes, or depletion of garnet and fluorapatite in HREE (Figs 8, 10a and b). Such a convex curve documenting depletion in F in either biotite or amphibole as a function of distance from an apparent sink for F, represented by the central fluid-rich granitic melt responsible for the pegmatoid dyke, essentially mirrors that of a concentration gradient following Fick’s Second Law or

$$\frac{\partial c}{\partial t} = D \frac{\partial^2 c}{\partial x^2}$$

(3)

where $c$ is concentration, $t$ is time, $D$ is the diffusion coefficient, and $x$ is the direction of the concentration gradient. Here, the shape of the F curve indicates that $(\partial^2 c/\partial x^2) < 0$, which would imply that F diffused along the concentration gradient towards the granitic melt.

This observation results in two inferences: (1) a continuous fluid medium existed along grain boundaries throughout both the dehydration zone and the surrounding granitic gneiss and (2) F was partitioned from the biotite and amphibole into this fluid and subsequently diffused along grain boundaries towards the granitic melt. This would imply that biotite and amphibole were in disequilibrium with the fluid in their immediate neighbourhood such that F was preferentially and continuously in the process of being partitioned from either mineral into this fluid. The pattern in biotite and amphibole F concentrations currently seen along the traverse would then represent a mirror of the concentration gradient existing in the fluid along grain boundaries at the point, when the granitic melt had completely crystallized to the pegmatoid dyke. These patterns would imply that the granitic melt acted as a natural sink for F. If true, the low F abundances in the pegmatoid dyke relative to the surrounding rock seen today (Table A1) would further imply that the granitic melt–tectonic fracture must have been an open system continually replenished by F-poor fluids such that F, diffusing in from the surrounding rock, was continually being diluted in this fluid-rich melt and removed, presumably to higher levels up along the rock column.

Trace element mineral chemistry

Despite having a sufficiently large $(Y + $REE$)$ budget [i.e. $(Y + $REE$)_{2}O_{3}$ of 0.5-1.0 wt %; see Harlov & Förster (2002b)], monazite and xenotime inclusions or rim grains associated with fluorapatite did not form during dehydration of the amphibolite-facies gneiss. In addition, fluorapatite shows no depletion in LREE nor does Population-I or -II allanite, with the exception of rare low-grade paratite-(Ce), show any alteration. In contrast, measurable depletion of Y in fluorapatite (Fig. 10b) from the dehydration zone, coupled with depletion of $(Y + $REE$)$ in the garnet (Fig. 8) as well as in the whole-rock analysis (Table A1; Fig. 3), indicates that these elements were highly mobile in the same fluids responsible for the partial conversion of biotite and amphibole to pyroxenes. Both experimentally and in nature, $(Y + $REE$)$ are known to preferentially complex with the F, whereas LREE preferentially complex with Cl (Haas et al., 1995; Pan & Fleet, 1996). The most likely scenario would be leaching of $(Y + $REE$)$ from the fluorapatite and garnet to complex with the F released during the partial conversion of biotite and amphibole to orthopyroxene and clinopyroxene as well as F partitioned out of the remaining biotite and amphibole and out of the fluorapatite. As a consequence, migration of F from the dehydration zone into the fluid-rich granitic melt responsible for the pegmatoid dyke would then deplete the dehydration zone in $(Y + $REE$)$. Lastly, lack of monazite and xenotime associated with fluorapatite, coupled with Cl depletion, puts constraints on the fluid to one containing CO$_2$ and NaCl components, but not dominated by CaCl$_2$ or KCl (Harlov & Förster, 2003), which agrees both with the fluid inclusion data as well as with the feldspar mineral chemistry.

INTERPRETATION OF FLUID INCLUSION DATA

Fluid compositions and densities, combined with textural information, delimit the trapping conditions and the relative timing for each of the fluid types during rock evolution. Hence, the pure H$_2$O fluids, typically trapped in healed fractures and evidently late, must be unrelated to the dehydration event. On the other hand, the high-salinity brines are clearly earlier than pure H$_2$O. These early fluids can be interpreted as (1) the original magmatic fluid inherent to the granitic gneiss, which originated during the migmatization event (ca. 1420–1460 Ma), and was probably collected by an uptake of water in the melt during migmatization of the granitic gneiss, and/or (2) brines produced by the unmixing of aqueous–carbonic fluids in the dehydration zone.

The construction of isochores allows for the possibility of determining the $PT$ conditions present during fluid trapping, and subsequent comparison with $PT$ estimates obtained from mineral thermobarometry. Isochores have been calculated from microthermometry data using the FLUID package of Bakker (2003). Figure 17 shows the isochores of selected fluid inclusions, together with a selection of metamorphic $PT$ conditions.

Identification of the dehydration fluid

The poorly preserved H$_2$O–CO$_2$–NaCl inclusions (see Fig. 15a) are identified as the dehydrating fluid. This
interpretation is supported both by the observation that CO₂-rich fluids are dominant in the pegmatoid dyke as well as by the position of the corresponding isochore, which demonstrates that the possible trapping conditions for H₂O–CO₂–NaCl inclusions with a fill degree of around 0.5 concur with the estimated PT conditions (650–700°C and 750–820 MPa) present during the dehydration event [e.g. the H₂O–CO₂–NaCl isochore in Fig. 17]. At these temperatures and pressures, the aqueous–carbonic fluid inclusions are homogeneous and would represent one possible means of transport for CO₂, namely as a part of a supercritical fluid mixture. This would explain the high CO₂ density currently seen in the carbonic fraction. In the dehydration zone, the H₂O–CO₂–NaCl fluids are relatively well preserved, suggesting near-isochoric uplift in the fluid miscibility field followed by fluid immiscibility towards lower temperatures. The latter conditions have resulted in the formation of the closely associated carbonic and brine inclusions.

Fig. 17. Representative isochores for the main fluid inclusion compositions: carbonic, aqueous (water or brine), and aqueous–carbonic inclusions. The shaded boxes indicate the metamorphic conditions during the dehydration event as revealed by the mineral chemistry. The larger box shows the maximum ranges implicit to the accuracy of the different mineral geothermobarometers. The retrograde uplift path is in part characterized by isothermal decompression (lithostatic). The earliest recognizable fluid is characterized by an H₂O–CO₂–NaCl composition. An example of such fluid inclusions is shown in Fig. 15a. Near-isothermal uplift resulted in the decrepitation of fluid inclusions under conditions of aqueous–carbonic immiscibility [e.g. Fig. 15b]. The latest part of the uplift path must have graded towards the 30°/km lithostatic gradient. Here, evidence was found for the admixing of pure water at temperatures below about 500°C. Implosion decrepitation resulted in progressive higher densities of the aqueous inclusions, whereas the CO₂-rich inclusions must have lost the water as a result of partial leakage along microfractures [e.g. Fig. 15c]. Geothermal gradients are given for 30°/km (lithostatic and hydrostatic) and for 50°/km (lithostatic). Miscibility limits are shown for salt-free H₂O–CO₂ and for H₂O–CO₂–NaCl (6 wt % NaCl relative to the NaCl subsystem and 21 mol % CO₂; Krüger & Diamond, 2001). (I) represents the peak of metamorphism characterized by local dehydration owing to the introduction of H₂O–CO₂–NaCl fluids from the initial tectonic fracture responsible for the dehydration zone in the granitic gneiss. Brines originating from the magmatic stage were probably already present (and in part preserved) in the granitic gneiss, but these fluids were immiscible with the dehydration fluid. (II) represents a more or less isochoric uplift, which left the trapped brines and dehydration fluids largely unchanged. (III) represents an almost isothermal uplift (dropping pressures) resulting in the decrepitation of fluid inclusions, almost directly followed by fluid unmixing and retrapping represented by (IV). During this stage, the salinity of the aqueous fluids was reduced and the density of CO₂ lowered. However, the higher-salinity and carbonic inclusions remained in close association. During final cooling, almost pure water was introduced into the system and in part disturbed the original contrasting fluid compositions. Retrapped water inclusions show progressive higher density (monophase). Carbonic inclusions with small amounts of water may have reduced their density as a result of water leakage.
As the source of the dehydration fluid is obviously the pegmatoid dyke, it is strange that aqueous–carbonic fluid inclusions are apparently rare here. This near absence would suggest that these fluids must have been expelled from the pegmatoid dyke before the end of its magmatic stage. However, the PT conditions were not sufficient to induce melting of the surrounding rock. In such a situation, the reactive potential of the crystallizing pegmatitic magma to cause intense alteration of the enclosing rock was even higher (see Figs 5, 6, 7, 10, and 16).

Relations between the different fluid types

The high-salinity brines (particularly those preserved in the granitic gneiss) must pre-date fluid infiltration. However, under the peak-metamorphic conditions, the possible mixing of brines and the dehydration fluid (homogeneous H$_2$O + CO$_2$) can be excluded. This is because of the presumed high salinity of the brines in the original migmatized granitic gneiss. The carbonic and brine inclusions are best preserved in the central part of the dehydration zone and, in spite of later modifications, reflect the fluid that controlled the dehydration. The pure CO$_2$ inclusions could have been derived from the original fluid responsible for the dehydration event, either by unmixing or by water leakage. However, unmixing is considered to be more likely, as this would explain the progressive density decrease related to isothermal uplift. Unmixing more efficiently reduces the CO$_2$ density in a progressive density decrease related to isothermal uplift. Considered to be more likely, as this would explain the unmixing or by water leakage. However, unmixing is fluid responsible for the dehydration event, either by mixing or by water leakage. However, unmixing is considered to be more likely, as this would explain the progressive density decrease related to isothermal uplift. Unmixing more efficiently reduces the CO$_2$ density in a progressive density decrease related to isothermal uplift. Considered to be more likely, as this would explain the unmixing or by water leakage. However, unmixing is fluid responsible for the dehydration event, either by mixing or by water leakage. However, unmixing is...
of K, Na, Ca, Sr, Rb, Ba, and the LREE (see previous discussion). At the same time, in an open system, a brine component could also be responsible for the depletion in metal cations such as Fe, Mg, Ti, and Zn as a result of complexing with Cl. With the exception of the Burton & O’Nions (1990) study, what is most intriguing is that, in each example including the present study, Y and the HREE (where tabulated) have lower abundances in the orthopyroxene ± clinopyroxene-bearing dehydrated rock than in the original amphibolite-facies gneiss. As already noted above, Y and the HREE prefer to complex with F as opposed to Cl. This would suggest that a F component was present in each of the dehydrating fluids. Besides an external source, F could also have come from the breakdown of biotite and amphibole during the dehydration event.

**LOCALIZED, FLUID-INDUCED, SOLID-STATE DEHYDRATION AT SÖNDRUN**

Low H₂O activities \(a_{\text{H}_2\text{O}} = 0.36; X_{\text{H}_2\text{O}} = 0.25\), as defined by orthopyroxene–biotite–K-feldspar–quartz, indicate that the fluids responsible for the dehydration event must have been present before the \(H_2O\)-rich melt responsible for the pegmatoid dyke. A \(CO_2\)-rich fluid with an \(H_2O\) activity of \(0.36\) could not have caused partial melting of the granitic gneiss at \(700^\circ C\), either \textit{in situ} or elsewhere. Rather, the fluid responsible must have had an \(H_2O\) activity of at least \(0.8\), which would lie well within the biotite–quartz stability field as opposed to orthopyroxene–K-feldspar (see Fig. 12). In addition, the very low solubility of \(CO_2\) in felsic melts (e.g. Ebadi & Johannes, 1991), coupled with the small volume of granitic melt represented by the pegmatoid dyke, would argue against such a melt being the principal source of \(CO_2\). In the dehydration zone, mineral textures and whole-rock chemistry give no evidence for partial melting nor that the dehydration zone might represent a residue from which \(H_2O\)-rich melts could have been extracted. Lastly, the relatively sharp contact between the pegmatoid dyke and the dehydration zone (Fig. 2a) suggests that the granitic melt responsible for it originated elsewhere and was later injected along the tectonic fracture after the actual dehydration event had occurred. \textit{In situ} localized partial melting would have produced a more gradual transition between the pegmatoid dyke and the dehydration zone.

Fluid inclusion data from all three zones suggest that whatever fluids were present during the dehydration event probably consisted of three phases (i.e. \(CO_2\), NaCl, and \(H_2O\)), although their exact proportions are not discernible. Experimental data on \(CO_2\)-NaCl–\(H_2O\) fluids (Johnson, 1991; Gibert \textit{et al.}, 1998; Shmulovich & Graham, 2004) suggest that, at \(700^\circ C\) and \(800\) MPa, the fluid would consist of two phases, \(CO_2\) and an NaCl brine. The limited extent of the dehydration zone (Holness, 1997; Gibert \textit{et al.}, 1998), coupled with the fluid inclusion data, would imply that \(CO_2\) was the principal component relative to NaCl brines. The presence of an NaCl brine, in fact, could be simply residual, dating from the migmatization of the original granitic gneiss. Evidence that an NaCl-rich brine was probably present during migmatization can be seen in the albite rims along plagioclase grain boundaries, albite rim intergrowths into the K-feldspar, the 20% albite component in the K-feldspar, and the 80% albite component in the plagioclase, as well as the fact that halogen-bearing minerals in the granitic gneiss tend to be more Cl enriched than the same minerals in the dehydration zone.

The two separate events (i.e. first dehydration of the surrounding granitic gneiss followed by injection of the granitic melts responsible for the pegmatoid dyke) would presumably have produced different effects with respect to the mineral chemistry, whole-rock data, and fluid inclusion data currently documented across the dehydration zone into the granitic gneiss (Figs 3, 5–8, and 10). During formation of the dehydration zone, fluid flow would have been purely advective as the \(CO_2\)-rich fluid moved out from the central tectonic fracture into the surrounding granitic gneiss, lowering the \(H_2O\) activity, and subsequently causing the partial, solid-state breakdown of biotite and amphibole to orthopyroxene and minor clinopyroxene. At the same time, the released Ca was apparently partitioned back into the amphibole (Fig. 6d). More importantly, the released Ti was partitioned back into the remaining biotite and amphiboles, which acted to stabilize them against further conversion to pyroxenes (Figs 5a and 6a). Fluorine shows some evidence of advective depletion in fluorapatite (Fig. 10c), whereas Cl shows advective depletion in fluorapatite as well as biotite and amphibole (Figs 5e, 6h, and 10d). Both halogens would have been released into the grain boundary fluids, and then apparently partitioned into the mobile fluids present in the tectonic fracture such that they were removed from the locality. The most likely explanation as to why a uniform depletion is seen in \((Y + HREE)\) across the dehydration zone (see Table A1; Figs 3, 8, and 10b) could be complexing with F (see the previous discussion). However, uniform depletion in Cl in all halogen-bearing minerals would suggest that Cl was evidently much more mobile. Yet the fact that the allanite in the dehydration zone appears to be unaltered compared with the allanite in the granitic gneiss, as well as the lack of depletion in LREE in other minerals, despite the affinity of LREE to complex with Cl (see the previous discussion), would imply that (1) there was considerably more F available in the dehydrating fluid than that supplied by the fluorapatite or (2) the affinity of the
Later emplacement of the granitic melt responsible for the pegmatoid dyke was the chief vehicle for causing the diffusion of various elements both throughout the dehydration zone as well as in the granitic gneiss. This is seen in the biotite and amphibole mineral chemistry, which indicate that the influence of the granitic melt was felt deep within the granitic gneiss, primarily in its behaviour as a strong chemical sink for F (Figs 5d and 6g). This influence was facilitated probably by NaCl brine-rich fluids along grain boundaries, which presumably were pervasive throughout the dehydration zone as well as regionally throughout the surrounding granitic gneiss. In addition, their presence would have also allowed for the high mobility of Fe and (Y + HREE) along grain boundaries towards the chemical potential sink represented by the granitic melt.

Although not easily discernible, the presence of clinopyroxene zones in the dehydration zone, both at the contact with the pegmatoid dyke as well as at the contact with the granitic gneiss, suggests that mechanical dispersion could have also played a role (see Fetter, 1985). The clinopyroxene zone, at the boundary between the dehydration zone and the pegmatoid dyke, could be the product of higher H₂O activities as a result of the proximity of the granitic melt as well as H₂O expelled by the crystallizing melt during formation of the pegmatoid dyke. In the case of the clinopyroxene zone at the boundary between the dehydration zone and the granitic gneiss, increased H₂O activities could be the result of an H₂O front pushed outward by the CO₂-rich dehydrating fluid, in addition to be mixing at the boundary with the more H₂O-rich fluids from the granitic gneiss. This scenario appears to be supported by the fluid inclusion data (see Fig. 16). The H₂O in such a front would be supplied by the breakdown of biotite and amphibole to pyroxenes. In either zone, formation of only clinopyroxene apparently reflects the H₂O activities present during crystallization of the pegmatoid dyke at presumably 650–700°C (ayH₂O > 0.8; see Fig. 12), where coexisting clinopyroxene and amphibole are also found, and water activities are lower than those in the granitic gneiss, where no clinopyroxene is found. In addition, either zone would appear to reflect higher Ca activities in the fluid as well.

It is interesting to note that the second elongated ‘fracture controlled’ dehydration zone along the eastern quarry wall, opposite the studied one in the Søndrum quarry, probably represents a continuation of the Søndrum dehydration zone–pegmatoid dyke complex, except that this dehydration zone consists only of a 0.5 m wide clinopyroxene-rich zone with a surrounding 0.5 m wide, diffuse, orthopyroxene- and clinopyroxene-bearing zone overprinting the granitic gneiss. This suggests the separation of two fluids flowing up the tectonic fracture marked by the clinopyroxene-rich zone, with the more CO₂-rich fluid partitioning off to also dehydrate the surrounding granitic gneiss. If this eastern dehydration zone indeed is a continuation of the western Søndrum dehydration zone–pegmatoid dyke complex, it is also an indication of what the western dehydration zone may have looked like before emplacement of the granitic melts responsible for the pegmatoid dyke; that is, a more H₂O-rich clinopyroxene zone surrounded by a relatively more H₂O-poor orthopyroxene-bearing zone. It would also imply that in three dimensions, one possible shape for the pegmatoid dyke–dehydration zone complex at Søndrum could be in the form of an elongated ellipsoid whose centre consists of the pegmatoid dyke surrounded by concentric rings consisting of an inner clinopyroxene zone, an orthopyroxene zone, and an outer clinopyroxene zone.

The source of the fluid responsible for the dehydration zone, and ultimately for the pegmatoid dyke, is unknown. Regionally, there are no known, nearby granite bodies of the same age (∼1400 Ma), which could have served as a fluid source. One possibility could be that the CO₂-rich fluids were forced up along some tectonic fracture, shear zone, or zone of weakness from below. Gravity maps of the Søndrum area suggest the presence of a body, or perhaps bodies, denser than the local granitic gneiss some kilometres below the present-day erosional surface (Graversen, 1998). Presuming that one of these bodies is a mafic intrusion concurrent with emplacement of the pegmatoid dyke, during crystallization it could have served as a source of the fluids. Other than H₂O, the principal component in the fluids given off by mafic magmas during crystallization would be CO₂ as well as brines such as NaCl [see discussion by Newton et al. (1998)]. The scenario described above would suggest that this fluid, at first, was primarily CO₂, which allowed for the formation of the dehydration zone, followed by a much more H₂O-rich fluid, perhaps during the later stages of crystallization of the mafic body, which resulted in the granitic melt responsible the pegmatoid dyke. The proposal that the granitic melt acted as a chemical sink for Cl, Fe, and (Y + HREE) relative to the surrounding rock would make sense only if it was an open system, acting as a conduit for the migration of fluid or melts away from this locality into the upper crust along the tectonic fracture. Curiously enough, the Varberg–Torpa charnockite–granite complex is also ∼1400 Ma (Åhäll et al., 1997; Christoffel et al., 1999). This suggests that there could be some relationship between the formation of this complex and the fluids or melts responsible for the pegmatoid dyke and surrounding dehydration zone at Søndrum, perhaps in the form of some deep-seated regional magmatic event at this time.

In summary, the pegmatoid dyke, the dehydration zone, the clinopyroxene zones, and the surrounding
granitic gneiss at Söndrum record the superposition of two physical processes, namely advective flow and diffusion. Advective flow, in the form of CO₂-rich fluids, along what was initially a tectonic fracture of some sort, came first and resulted in the formation of the dehydration zone, and is limited to the margins of the dehydration zone. Advection is reflected in the partial conversion of biotite and amphibole to pyroxenes, coupled with the stabilization of the remaining biotite and amphibole through enrichment in Ti (Figs 5c and 6a). It is also reflected in the uniform enrichment, or the uniform depletion, of a particular element in a particular mineral, such as the uniform depletion of Cl in biotite, amphibole and fluorapatite (Figs 5e, 6h, and 10d) or the uniform enrichment of Ca in amphibole (Fig. 6d). In contrast, diffusion came later, with emplacement of the granitic melt responsible for the pegmatoid dyke. It is not limited to the dehydration zone, but extends far into the granitic gneiss, as is seen in the concentration gradient of F in both biotite and amphibole along the traverse (Figs 5d and 6g). Here, the element diffusion profile in either mineral approximates Fick's Second Law. It represents the element concentrations present in the original interconnected grain boundary fluid, now mirrored in the mineral chemistry, with which the fluid is presumed to have been in local equilibrium. Whereas advective flow and diffusion can overlap in a manner independent of each other, as witnessed by the contrasting behaviours of F and Cl in amphibole or biotite, they can also be influenced by each other such that a profile that is primarily advective also shows some influence from diffusion. This is particularly seen in the behaviour of Na and K in amphibole, or the behaviour of Fe, Mg, and Mn in each of the Fe–Mg silicate minerals (Figs 5–7).

Comparison of the results and conclusions of this study with the many previous studies of localized, solid-state dehydration indicate that, no matter what the fluid source, solid-state breakdown of biotite and amphibole to pyroxenes is a function of low H₂O activity fluid infiltration of the rock, the fluids already present along grain boundaries, and, just as importantly, the chemistry of the Fe–Mg silicate minerals present (i.e. Fe/Mg ratio). All three act in unison to determine the balance and direction of the chemical potentials both in the dehydration zone and beyond into the surrounding, non-dehydrated rock. This interaction not only manifests itself in the lowering of H₂O activities beneath a certain threshold so as to allow for pyroxene formation, but also leaves behind additional chemical fingerprints in the metal cation, halogen, and (Y + REE) chemistry of the silicate and phosphate minerals, which are themselves mirrors of both the chemistry of the fluids present during the dehydration event as well as the extent of their total interaction with the rock at large.

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SUPPLEMENTARY DATA
Supplementary data for this paper are available at Journal of Petrology online.

REFERENCES


