Shear zones channelize fluid flow in the Earth's crust. However, little is known about deep crustal fluid migration and how fluids are channelized and distributed in a deforming lower crustal shear zone. This study investigates the deformation mechanisms, fluid-rock interaction and development of porosity in a monzonite ultramylonite from Lofoten, northern Norway. The rock was deformed and transformed into an ultramylonite under lower crustal conditions (T=700-730° C, P=0.65-0.8 GPa). The ultramylonite consists of feldspathic layers and domains of amphibole + quartz + calcite, which result from hydration reactions of magmatic clinopyroxene. The average grain size in both domains is <25 m. Microstructural observations and EBSD analysis are consistent with diffusion creep as the dominant deformation mechanism in both domains. Festoons of isolated quartz grains define C'-type shear bands in feldspathic layers. These quartz grains do not show a crystallographic preferred orientation. The alignment of quartz grains is parallel to the preferred elongation of pores in the ultramylonites, as evidenced from synchrotron X-ray microtomography. Such C'-type shear bands are interpreted as creep cavitation bands resulting from diffusion creep deformation associated with grain boundary sliding. Mass-balance calculation indicates a 2% volume increase during the protolith-ultramylonite transformation, which is consistent with synkinematic formation of creep cavities producing dilatancy. Thus, this study presents evidence that creep cavitation bands may control deep crustal porosity and fluid flow. Nucleation of new phases in creep cavitation bands inhibits grain growth and enhances the activity of grain-size sensitive creep, thereby stabilising strain localization in the polymineralic ultramylonites.
Creep cavitation bands control porosity and fluid flow in lower crustal shear zones

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ABSTRACT

Shear zones channelize fluid flow in the Earth’s crust. However, little is known about deep crustal fluid migration and how fluids are channelized and distributed in a deforming lower crustal shear zone. This study investigates the deformation mechanisms, fluid-rock interaction and development of porosity in a monzonite ultramylonite from Lofoten, northern Norway. The rock was deformed and transformed into an ultramylonite under lower crustal conditions (T=700-730°C, P=0.65-0.8 GPa). The ultramylonite consists of feldspathic layers and domains of amphibole + quartz + calcite, which result from hydration reactions of magmatic clinopyroxene. The average grain size in both domains is <25 μm. Microstructural observations and EBSD analysis are consistent with diffusion creep as the dominant deformation mechanism in both domains. Festoons of isolated quartz grains define C’-type shear bands in feldspathic layers. These quartz grains do not show a crystallographic preferred orientation. The alignment of quartz grains is parallel to the preferred elongation of pores in the ultramylonites, as evidenced from synchrotron X-ray microtomography. Such C’-type shear bands are interpreted as creep...
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grain growth and enhances the activity of grain-size sensitive creep, thereby stabilising strain
localization in the polymineralic ultramylonites.

INTRODUCTION

Many studies document that metamorphic reactions and viscous deformation in the lower
crust are triggered by grain-size reduction and fluid infiltration (e.g., Austrheim, 1987; Rutter
and Brodie 1992; Getsinger et al., 2013). A very fine grain size of reaction products may activate
grain size sensitive creep, which leads to a marked weakening of the rock and to strain
localization (Rutter and Brodie 1992; Pearce et al., 2011). The feedback between grain size
reduction, fluid flow, and the activity of different deformation mechanisms is critical for the
understanding of the rheology of shear zones and the processes leading to strain localization.

Deformation-enhanced fluid flow and development of synkinematic porosity in mid-
crustal shear zones rocks has been a subject of numerous studies (e.g. Mancktelow et al., 1998;
Fusseis et al., 2009). Fluid transfer has been linked to syndeformational dynamic porosity
generated by creep cavitation during viscous grain boundary sliding (e.g., the granular fluid
pump model: Fusseis et al., 2009). Fluid infiltration results in shear zones being preferential
conduits for fluid flow even at deeper crustal conditions (Austrheim, 1987; Mancktelow, 2006).
However, little is known about the fluid flow in the lower crust, and, more specifically, about the
processes that control formation and distribution of syndeformational porosity.
High strain torsion experiments on synthetic anorthite aggregates deforming by grain boundary sliding have highlighted the development of creep cavitation bands (Rybacki et al., 2008; 2010). The bands developed with a C’-type shear band orientation, presumably from growth and coalescence of individual pores originally formed at triple junctions and dilatant sites resulting from the operation of grain boundary sliding. However, observational evidence of similar creep cavitation bands in natural ultramylonites from the lower crust is currently lacking, thus questioning the extrapolation of such experimental results to natural conditions.

To investigate the relationships between deformation mechanisms and the formation and distribution of porosity in lower crustal shear zones, we have analysed a feldspar-rich ultramylonite deformed at T>700° C. Our results provide evidence for the formation of creep cavitation bands during grain-size sensitive creep and have important implications for the understanding of high-temperature creep and synkinematic fluid flow in the lower crust.

SAMPLES AND METHODS

We sampled a shear zone in the Anorthosite-Monzonite-Charnockite-Granite intrusive suite of Lofoten, northern Norway (Corfu, 2004, and references therein). The shear zone is hosted in monzonite and shows a mylonite to ultramylonite transition from the shear zone boundary to the shear zone centre (see Fig. DR1; sample location in UTM coordinates relative to WGS84: zone 33W, 0505656E, 7594514N). The transition is marked by an extreme grain size reduction of perthites and clinopyroxene. Grain size reduction occurred by fracturing and neocrystallization in perthites and by hydration reactions in clinopyroxene, forming amphibole + quartz + calcite. Plagioclase-amphibole geothermobarometry and Ti-in-amphibole geothermometry yield P, T conditions of deformation of 700-730° C, 0.65-0.8 GPa (Menegon et al., 2013).
We used a combination of detailed microstructural analysis, synchrotron X-ray microtomography and mass-balance calculations to characterize deformation microstructures and the associated synkinematic porosity in the ultramylonite. Electron backscatter diffraction (EBSD) was used to quantify the crystallographic preferred orientation (CPO) of feldspars, amphibole and quartz. Analytical methods are described in detail in the Data Repository.

RESULTS

Microstructure and EBSD analysis

The ultramylonite displays a compositional banding between feldspathic layers and domains of pyroxene-derived reaction products (amphibole, quartz and calcite) (Fig. 1A). The feldspathic layers originate from the neocrystallization of perthites, and may locally contain also quartz, amphibole and biotite (Fig. 1B). The rare clinopyroxene porphyroclasts preserved in the ultramylonite show the localization of reaction products along intragranular fractures (Fig. 1C). Calcite is a synkinematic reaction product, typically at triple junctions and dilatant sites (Fig. 1D). In both feldspathic and pyroxene-derived domains the average grain size is < 25 μm.

The EBSD phase map of a feldspathic layer shown in Fig. 2A is dominated by the bi-phase mixture of plagioclase and K-feldspar deriving from the recrystallization of original perthites. The EDS compositional map of the Si content of the same area is shown in Fig. 2B. Quartz occurs as isolated grains along discrete C’-bands inclined at 10-20° to the trace of the ultramylonite foliation, consistent with the sinistral sense of shear (Fig. 2A, 2B). The CPO of quartz, K-feldspar and plagioclase and does not show a clear relationship of crystallographic planes and axes with the kinematic framework of the ultramylonite (Fig. 2C). The long axis of quartz grains are preferentially oriented either at 0-40° or at 160-180° to the trace of the ultramylonite foliation, measured anticlockwise (Fig. 2D). An additional EBSD map of a
feldspathic layer containing festoons of quartz grains in a C’-band orientation is shown in the supplementary material (Fig. DR2).

Amphibole CPO in an elongate domain of reaction products shows clusters of [001] axes oriented at a low angle to the stretching lineation. Poles to the (100) and (010) planes are preferentially distributed along a girdle subparallel to the YZ plane (Fig. 2E). In the same domain, quartz c-axis CPO is weak and characterized by some clustering at a low angle to the foliation plane. One cluster occurs near the centre of the pole figure (Fig. 2E).

Porosity distribution and orientation

We used synchrotron X-ray microtomography to analyse the distribution and orientation of pores in two feldspathic layers that were micro-drilled from the ultramylonite sample (Fig. 3A; see GSA Data Repository for details on data acquisition). The absorption microtomographic data resolve the different materials in the sample well and clearly highlight the pores, which attenuate the least and appear darkest (yellow in Fig. 3A; movies DR_ Lu-1_light_pores_slcmigration and DR_ Lu-3_light_pores_slcmigration in the Data Repository).

Low-aspect-ratio-pores can easily be distinguished from cracks that might have formed along grain boundaries during exhumation and cooling of the rocks (movie Lu-1_pores_slcmigration_1). The latter were excluded from the following analyses. Although pore diameters can vary from about the resolution limit (1.5 μm diameter) to about 20 μm, they generally are significantly smaller than the grains themselves. The pores decorate phase and grain boundaries between feldspars and quartz, hornblende and/or biotite and often form festoons or clusters of more than two pores.

Pores were segmented by binary thresholding and analysed for their orientations. Orientation of each pore was defined as the orientation of the best ellipsoid fit to the pore’s
shape. To avoid a bias in the orientation data, the analysis was limited to pores with volumes between 34 μm$^3$ (125 voxels) and 4119 μm$^3$ (1.5x10$^4$ voxels). In figure 3 we show the results from one feldspathic layer (data-set Lu-3_light); similar results were obtained from the second feldspathic layer (data-set Lu-1_light: see Fig. DR3). The pole figure shows the orientation of the long axis of the best-fit ellipsoid of the pores. These are referenced to the trace of the mylonitic foliation (a kinematic framework defined by the long axis of the best ellipsoid fit to the biotite grains, Fig. 3B). The diagram reveals that the pores have a preferred orientation, with their long axes oriented at 20-30° to the trace of the mylonitic foliation, in a C’-band orientation (Fig. 3B).

**Mass-balance calculations**

Whole-rock chemistry and total carbon (TC) analysis was performed to assess element mobility and volume changes during the protolith-to-ultramylonite transformation using the ‘normalized Gresens’ method (Potdevin and Marquer, 1987). Results of the whole-rock chemistry analysis are reported in Table DR1.

The total carbon (TC) content of the ultramylonite is 2.75 times higher than the original content in the protolith (0.139 Vs. 0.037 wt%). Assuming that the original monzonite contained minute amounts of carbonate material, we used TC as a reference to calculate the volume change associated with the shear zone formation. The TC increase reflects CO$_2$ infiltration during shearing, consistent with the synkinematic growth of calcite in the ultramylonite (Figs. 1C, 1D). A 2.75 times increase of TC implies a volume increase of 2.3%. Detailed information on the mass-balance results is compiled in the Data Repository.

**DISCUSSION**

The CPO of all phases, the grain size and shape, and phase mixing indicate that grain size sensitive creep was the dominant deformation mechanism in both, the feldspathic layers and the
pyroxene-derived layers in the ultramylonite. The feldspar CPO data is not interpretable in terms
of intracrystalline plasticity. In the layers composed of reaction products, the similar grain sizes
and shapes of calcite, quartz and hornblende indicate the dominance of grain size sensitive creep
(Getinger et al., 2013). At the given temperatures, calcite is expected to be substantially weaker
than quartz and hornblende if deforming by dislocation creep (e.g., Renner et al., 2007).
However, in the microstructures calcite never appears less competent, and all three phases show
similar aspect ratios and grain sizes (Fig 1D). The distribution of hornblende [001] axes near the
instantaneous stretching axis for a sinistral sense of shear (Fig. 2C) is consistent with a shape
fabric attained by oriented growth and/or rigid body rotation during deformation accommodated
by diffusion creep and grain boundary sliding, which is a common feature in amphibole
deforming at lower crustal conditions (e.g., Berger and Stünnitz, 1996; Getinger and Hirth,
2014). Quartz c-axes show weak maxima oriented similar to hornblende [001] axes. We likewise
interpret this weak CPO as the result of preferential synkinematic growth of quartz grains with
their c-axis parallel to the elongation direction during diffusion creep (e.g. Hippertt, 1994;
Hippertt and Egydio-Silva, 1996; Kilian et al., 2011).

The occurrence of quartz grains in a C’-band orientation in the feldspathic layers is
interpreted as the result of creep cavitation, which is referred to as the coalescence of
intergranular pores originally formed at grain triple junctions and grain boundaries (Zavada et
al., 2007; Rybacki et al., 2008, 2010; Delle Piane et al., 2009). Creep cavitation takes place
during grain boundary sliding, and dilating creep cavities form local sites of low stress that
attract grain boundary fluids (Fusseis et al., 2009). Our mass-balance calculations indicate (1) a
volume increase of 2.3%, and (2) fluid infiltration during the protolith-ultramylonite
transformation. Hence, positive volume change accompanied by fluid infiltration can explain the
precipitation of new phases from intragranular aqueous fluids collected in cavitation bands. Volume increase is a consequence of dilatancy at grain boundaries (Schmocker et al., 2003; Fusseis et al., 2009).

Our interpretation is supported by the similar orientation of the preferred elongation of pores in the feldspathic layers and the orientation of the C’ bands (Figs. 2A-B and 3). The orientation of pores is not related to specific phase boundaries but to the kinematic framework of the shear zone. Thus, we interpret the final porosity imaged by X-ray microtomography as representative of the porosity and fluid flow at an instant during deformation. The preferred distribution of pores and isolated quartz grains in a C’-type shear band orientation is a syn-deformational feature reflecting the local dilatancy in a dynamically evolving microstructure during diffusion creep deformation (Schmocker et al., 2003; Rybacki et al. 2008, 2010). Grain boundary sliding, creep cavitation, and heterogeneous nucleation form pores or new grains in low stress sites (Ree 1994; Kassner and Hayes, 2003; Kilian et al., 2011) (Fig. 4). Dilatancy has an initial form normal to the extension direction and only after some extension and further opening of the porosity pore shape attains a stable orientation along C’-bands (Fig. 4).

The precipitation of quartz along dilatant grain boundaries requires material transport, most likely in a grain boundary fluid film. The interpretation is that quartz is dissolved from the pyroxene-derived reaction products (Fig. 1C) and precipitates locally in dilatant sites in the feldspathic layers. Dissolution, transport, grain rotation, and precipitation are intimately related processes during diffusion creep of geological material (e.g. Fusseis et al., 2009; Kilian et al., 2011), and may result in a dynamically evolving microstructure and distribution of porosity.

There is a certain degree of similarity between shape of pores and quartz grains. About 50% of the quartz grains contained in the festoons in Figs. 2A-B are preferentially elongated at 0-40°
to the trace of the foliation, similar to the elongation of pores (Figs. 2D, 3B). It could indicate that shape of quartz grains is determined by the cavitation process (Fig. 4). However, considering the shape modifications that quartz grains can undergo after precipitation (i.e. dissolution, grain rotation), this is a speculation.

**IMPLICATIONS AND CONCLUSIONS**

We conclude that the orientations of pores, quartz bands and phase boundaries along C’-type shear bands in the ultramylonite are evidence of creep cavitation during lower crustal deformation accommodated by diffusion creep, grain boundary sliding and heterogeneous nucleation. The microstructures presented in this paper share many similarities with the creep cavitation bands reported from experimental deformation of synthetic anorthite aggregates (Rybacki et al., 2008; 2010). Creep cavitation bands can be identified by the occurrence of pores and isolated grains of different phases. However, if the same phases precipitate, this will result in overgrowths on existing grains, thereby rendering the identification of dilatancy and cavitation bands difficult.

Strain localization in lower crustal rocks is typically associated with grain size reduction, hydration reactions and phase mixing (e.g., Rutter and Brodie, 1992; Pearce et al., 2011; Getsinger et al., 2013). Phase mixing by heterogeneous nucleation during grain size sensitive creep critically relies on synkinematic porosity (e.g., Hiraga et al., 2013). Creep cavitation can be a major contributor to porosity in lower crustal shear zones, and hence control fluid flow.

Nucleation of new phases in cavitation bands inhibits grain growth and enhances the activity of grain-size sensitive creep, thereby maintaining strain localized in the polymineralic ultramylonite (e.g., Herwegh et al., 2011). Thus, our findings provide a key component for the understanding
of strain localization in the lower crust and of the mechanisms by which fluid flow can be
channelized within lower crustal shear zones.

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**FIGURE CAPTIONS**

Figure 1. Microstructure of the ultramylonite in thin sections cut normal to foliation and parallel to stretching lineation. Abbreviations: FL=feldspathic layer, PDRP=pyroxene-derived reaction products, Pl=plagioclase, Kfs=K-feldspar, Qtz=quartz, Hbl=hornblende, Cpx=clinopyroxene, Cc=calcite, Bt=biotite. A: Light micrograph of the layered microstructure of the ultramylonite. Parallel polarizer. B: SEM backscattered electron image of a feldspathic layer. C: SEM backscattered electron image of pyroxene-derived reaction products. Note the clinopyroxene porphyroclast with reaction products along intracrystalline fractures. D: Close-up of pyroxene-derived reaction products. SEM backscattered electron image.
Figure 2. Results of EBSD and EDS analysis of the ultramylonite. All plots are upper hemisphere projections contoured with 15° half-width and 5° cluster size using Channel 5 (Oxford Instruments). Inset in C shows the kinematic framework of the sample ($L_s$=stretching lineation). Data is plotted as one point per grain ($N$ = number of plotted grains). Maxima are expressed as multiples of the uniform distribution. Mean angular deviation number for all datasets is $< 0.9$. Shear sense is sinistral. A: Processed EBSD phase map from a feldspathic layer. The map is superposed to the band contrast map. Grey areas are non-indexed points. Mineral abbreviations: Kfs=K-feldspar, Pl=plagioclase, Qtz=quartz, Cc=calcite, Bt=biotite, Grt=garnet. B: EDS-derived compositional map of Si content of the same area shown in A. C: Pole figures of the crystallographic orientation data of quartz, K-feldspar and plagioclase from the area shown in A. D: Rose diagram to show the orientation of the long axis of quartz grains included in A. Only grains with aspect ratio $> 1.3$ are considered ($N=50$). E: Pole figures of the crystallographic orientation data of hornblende and quartz from a layer of pyroxene-derived reaction products.

Figure 3. Synchrotron x-ray microtomographic data. S is the trace of the ultramylonite foliation, $C'$ the trace of $C'$-bands, $L_s$ the stretching lineation (red dot). A: Slice through microtomographic data Lu-3_light, showing 3D objects in 2D. Grey values correspond to x-ray absorption. Red=biotite, yellow=pores. The grey-scale image is the backside of a thin migrating box through the 3D data-set, in which pores and biotite are highlighted. As the box moves through the volume, pores and biotite disappear out of the box at the front and enter the slice at the back, through the greyscale image (see movie DR_Lu-3_light_pores_slcmigration in the Data Repository). The inset shows the trends of preferred orientation of pores and biotite. B: Pole figure illustrating the long axes of pores preferentially oriented at 20-30° to the foliation (top) and the preferred orientation of biotite long axis aligned in the foliation plane (bottom). Data is
plotted as one-point-per-pore (or biotite) and as contoured pole figures. Equal area lower hemisphere stereoplots. Contouring is up to 10 times MUD.

Figure 4. Schematic drawing of cavitation during grain boundary sliding in shear (after Pilling and Ridley 1989). Elongation orientation of the pores will depend on the extent of dilatancy.

1GSA Data Repository item 2014xxx, xxxxxxxx, is available online at www.geosociety.org/pubs/ft2014.htm, or on request from editing@geosociety.org or Documents Secretary, GSA, P.O. Box 9140, Boulder, CO 80301, USA.
Figure 3

Click here to download Figure: Fig3_microtomography_new.pdf

Fig. 3
shear plane

starting material

nucleation of pores

progressive coalescence of pores in a C’-band orientation
SUPPLEMENTARY MATERIALS

Methods

Light- and Scanning Electron Microscopy. The petrography and microstructure of the ultramylonite have been investigated with polarized light- and scanning electron microscopy on polished thin sections cut perpendicular to the foliation and parallel to the stretching lineation. SEM backscatter electron images were collected with a Jeol-840 SEM at the Department of Medical Biology, University of Tromsø, and with a Philips XL-30 FEG-ESEM at the Department of Geological Sciences, Stockholm University. The same thin sections were used for electron backscattered diffraction (EBSD). The grain size and aspect ratio of individual grains were measured on grain boundary maps obtained from manually digitizing SEM-BSE and EBSD images. The 2D size of the grains was calculated as the diameter of the circle with an area equivalent to that of the grain using the freeware Image SXM software (http://www.ImageSXM.org.uk).

EBSD and EDS Analysis. EBSD and EDS analysis were carried out on a Jeol LV6610 SEM equipped with an Oxford Instruments Nordlys Nano EBSD detector and with an Oxford Instruments SDD X-Max 80 mm² EDS detector at the Electron Microscopy Centre of Plymouth University. Additional EBSD analysis was conducted on a Philips XL-30 FEG-ESEM equipped with a HKL Technology (Oxford Instruments) Nordlys detector at the Department of Geological Sciences, Stockholm University. Thin sections were chemically polished carbon coated (for EBSD analysis in Plymouth) or left uncoated (for the EBSD analysis in Stockholm) during the acquisition of the electron backscatter patterns over gridded areas of varying sizes. Step sizes of 1, 2 and 3 μm were used in the 3 EBSD datasets presented in this study. Working conditions during the pattern acquisition were 20 or 25 keV accelerating voltage and either low vacuum (0.3-0.4 torr: Stockholm) or high vacuum (Plymouth). EBSD patterns were indexed and processed with the Channel 5 analysis suite from HKL Technology (Oxford Instruments). A match unit for oligoclase was created with the Twist component of the Channel 5 suite using the cell parameters for An16 (spatial group C-1) reported in Phillips et al. (1971) and served as theoretical model to index plagioclase. Noise reduction on the raw EBSD data was performed following the procedure tested by Prior et al. (2002) and Bestmann and Prior (2003). Crystallographic data were plotted on pole figures (upper hemispheres) using one point per grain.
X-Ray Microtomography. Microtomographic samples with a diameter of 1 mm were extracted from the ultramylonite sample shown in Fig. DR1B using a rock drill. These samples were scanned at the microtomography beamline 2BM of the Advanced Photon Source (USA). A double multilayer monochromator of 1.5% band-width provided 27 KeV X-rays; radiographic projections were collected in transmission mode by a CCD camera behind the sample in the hutch configuration. The sample detector distance was 70 mm. During each scan, 1440 projections were collected through rotating the samples in steps of 0.125° over 180°. The acquisition time for each data set was about 25 min. From the radiographic projections, three-dimensional absorption microtomography datasets were reconstructed using filtered back-projection.

Two microtomographic data-sets (Lu-1_light and Lu-3_light) were cropped to a volume of 1000x1000x750 voxels, which corresponds to 650x650x488 µm. From these subvolumes, pores, which are the least-attenuating, hence darkest phase in the data, were segmented by binary thresholding. From the same raw data, micas were segmented using the same algorithm. Biotite was chosen because it defines the mylonitic foliation in the sample. From the segmented mica data artefacts (mostly phase contrast ‘shadows’) had to be manually removed. Binary data of both the pores and the micas were sequentially loaded into Blob3D (Ketcham, 2005) for analysis. Blob3D recognises face-connected voxels of the same kind as clusters (or ‘blobs’), which allows determining for each cluster the volume, shape, location and orientation (given in direction cosine of the inscribed eigenvectors), amongst other parameters. For our orientation analysis, all clusters smaller than 34 µm^3 (125 voxels) and larger than 4120 µm^3 (15000 voxels) were discarded. The former would have introduced artefacts due to the limited possibilities to arrange a small number of voxels in a pore cluster, and the latter would have very complex shapes, producing meaningless results. We furthermore discarded pore clusters with aspect ratios ≤ 1.7 and mica clusters with aspect ratios < 3. The orientation values of the longest eigenvector were converted into dip direction/dip angle values using the formulation given in Groshong (2006).

For each of the two datasets (Lu-1_light and Lu-3_light), two orientation datasets exist – one for the mica grains and one for the pores. The longest eigenvectors of both mica data-sets cluster around well-defined maxima. We assume these maxima to define the orientation of the longest diameter of the finite strain ellipsoid in each sample. As can be seen in the supplementary movies and also Fig. 3A, the mica furthermore define a mylonitic foliation. We used these two orientations as a kinematic framework. Because this framework does not
spatially coincide with the Cartesian coordinate axes of the microtomography data, we rotated the maximum defined by the longest eigenvectors of the mica data into a horizontal E-W orientation and the pole to the foliation in a horizontal N-S orientation using Stereo32 (Fig. 3A and DR2). The exact same rotations were then applied to the longest eigenvectors of the Lu-3_light and Lu-1_light pore populations, which yield the stereo plots shown in Fig. 3B and DR2. This allowed us to assess the orientation of the pores in a kinematic context.

**Whole-Rock Chemical Analysis.** Whole-rock chemical analysis of major elements was performed by wavelength dispersive X-Ray fluorescence (WD-XRF) analysis with a Bruker S8 Tiger XRF spectrometer at the Department of Geology, University of Tromsø. Powder samples were mixed and diluted at 1:7 with Li$_2$B$_4$O$_7$ flux, and melted into fused beads. Loss on ignition (LOI) was determined from weight lost after ignition at 1050°C for 1.5 h. Total Carbon (TC) was measured with a LECO CS-200 at the Department of Geology, University of Tromsø. The LECO CS-200 uses infrared absorption to measure the quantity of carbon dioxide generated by combustion of the sample in an induction furnace in a pure oxygen environment. Accuracy of the measurements is ± 2 ppm.

**Supplementary Information on the Mass-Balance Calculations**

The two samples were collected along a continuous strain gradient in the field at a relative distance of 1 meter, so that we can safely conclude that the ultramylonite derives from (micro)structural and mineralogical modifications of the monzonite. The mass-balance calculations were carried out following the method designed by Potdevin and Marquer (1987), which is referred to as the ‘normalized Gresens’ method. The method uses the following equation to derive mass gain or loss of a chemical component n during modification of rock A to rock B in relationship to the initial amount of the component n in rock A:

$$\Delta X_n = F_v \left( \frac{\rho_B}{\rho_A} \right) \left( \frac{X_{nB}}{X_{nA}} \right) - 1$$  \hspace{1cm} (1)

$\Delta X_n$ represents the gain or loss of chemical component n related to its initial content in rock A, $F_v$ is the volume factor ($F_v = V$ modified rock B/$V$ initial rock A), $X_{nA}$ is the weight % of the component n in the initial rock A, $X_{nB}$ is the weight % of the component n in the modified rock B, $\rho_A$ is the density of the initial rock A, and $\rho_B$ is the density of the modified rock B. Our calculations refer to the transformation protolith $\rightarrow$ ultramylonite (sample LST29F $\rightarrow$ sample LST29B). The main differences are in the LOI and TC content, consistent with fluid infiltration during shear zone formation. The very minor difference between the compositions
of the monzonite and ultramylonite (Table DR1) indicate that, apart from the fluid infiltration, there has been no major chemical change during deformation. Our whole-rock chemical composition data are consistent with the average composition of the Raftsund mangerite (Griffin et al. 1978). Thus, we are confident that primary heterogeneities of the protolith do not represent a limitation to our analysis.

The density of the samples has been measured with a pycnometer using pulverized material at the Department of Geology, University of Tromsø, following routine procedures outlined in Hutchinson (1975). Weight measurements were repeated 5 times for each sample, and were reproducible with an accuracy of ± 0.004 grams. The densities of the protolith (monzonite) and of the ultramylonite are 2.711 g/cm³ and 2.654 g/cm³, respectively.

Setting ΔXn for TC = 2.75 and solving (1) for Fv, we obtain Fv = 1.023. This means a 2.3% volume increase. Setting Fv = 1.023 and solving (1) for ΔXn, we derive the following gains or losses of chemical components:

<table>
<thead>
<tr>
<th>Component</th>
<th>Fv</th>
<th>ΔXn</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>1.023</td>
<td>0.0150</td>
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REFERENCES CITED


**Table DR1.** Whole-rock chemical composition (major elements) of the protolith (LST29F) and of the ultramylonite (LST29B) samples used for the mass-balance calculations.

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**Figure DR1.** Hand specimens of the monzonite protolith (A) and of the mylonite-ultramylonite transition (B).
**Figure DR2.** A: EBSD-derived phase map of part of the same feldspathic layer shown in Fig. 2A of the paper. Note the festoons of quartz grains with a C'-band orientation for a sinistral sense of shear. B: Pole figures of K-feldspar, plagioclase and quartz grains in A.

**Figure DR3.** Synchrotron x-ray microtomographic data Lu-1_light (compare with Fig. 3B). S is the trace of the ultramylonite foliation, Z is the pole to the foliation, X is parallel to the stretching lineation. Pole figure illustrating the long axes of pores preferentially oriented at 20-30° to the foliation (top) and the preferred orientation of biotite long axis aligned in the
foliation plane (bottom). Equal area lower hemisphere stereoplots. Contouring is up to 10 times MUD.

**Movie DR_Lu-1_pores_slemigration.** Migrating slice through the microtomographic data-set Lu-1_light (volume of 650x650x488 μm). Grey values correspond to x-ray absorption. See text for further details.

**Movie DR_Lu-3_pores_slemigration.** Thick migrating slice through the microtomographic data-set Lu-3_light (volume of 650x650x488 μm). Grey values correspond to x-ray absorption. Red=biotite, yellow=pores. The greyscale image is the backside of a thin migrating box through the 3D data-set, in which pores and biotite are highlighted. As the box moves through the volume, pores and biotite disappear out of the box at the front and enter the slice at the back, through the greyscale image. See text for further details.
Movie file 1 (supplemental file)
Click here to download Movie File: DR_Lu-1_light_pores_slcmigration.mpg
Movie File 2 (supplemental file)

Click here to download Movie File: DR_Lu-3_light_pores_slcmigration.mpg