Original paper

The Sklené garnet peridotite: petrology, geochemistry, and structure of a mantle-derived boudin in Moldanubian granulite

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The Moldanubian Sklené garnet peridotite, which preserves mineralogical, geochemical, and structural evidence for a protracted mantle evolution, is a small boudin (~1 m) enclosed by granulite of the Gföhl Assemblage. The peridotite has a porphyroclastic texture, in which large garnet and smaller olivine and pyroxene porphyroclasts (Stage 1) reside in a fine-grained, recrystallized matrix of olivine, pyroxene, and pargasitic amphibole (Stage 2). Garnet is surrounded by kelyphite, consisting of an outer zone of pargasite and an inner zone of spinel and pyroxene (Stage 3).

Lattice-preferred orientations (LPO) for olivine, orthopyroxene, and clinopyroxene were measured by electron backscatter diffraction. The LPO of olivine (mostly Stage 2) has an [010] maximum parallel to the Z fabric direction and [100] and [001] girdles in the X–Y fabric plane. Both orthopyroxene and clinopyroxene have [010] maxima parallel to Z and [001] girdles in the X–Y plane.

The Sklené peridotite is slightly depleted in incompatible major and minor elements relative to primitive mantle. Whole-rock REE contents are 0.3 to 0.8 times that of primitive mantle, with a convex upward pattern, and Rb, Th, U, Pb, and Sr show positive anomalies in an extended element plot. Among the highly siderophile elements, Os, Ir, Ru, and Pt are unfractionated, but Pd and Re are significantly depleted. A Sm-Nd garnet-whole rock-clinopyroxene isochron yields a cooling age of 338 ± 13 Ma, and $\epsilon_{Nd}$ and ($^{87}\text{Sr}/^{86}\text{Sr})_i$ values for clinopyroxene (at 335 Ma) are +4.5 and 0.7035, respectively.

By comparison with other Moldanubian garnet peridotites in the Czech Republic, the Sklené peridotite likely represents a fragment of subcontinental lithospheric mantle. It experienced early partial fusion and depletion of incompatible elements at moderate pressure in the spinel stability field, followed by transport to high-pressure conditions at 1230 °C and 50 kbar, where the Stage 1 garnet-bearing assemblage was stabilized. Subsequently, the depleted peridotite was metasomatized by transient, primitive basaltic melts, with the addition of Rb, Th, U, Pb, and Sr showing positive anomalies in an extended element plot. Among the highly siderophile elements, Os, Ir, Ru, and Pt were unfractionated, but Pd and Re were significantly depleted. A Sm-Nd garnet-whole rock-clinopyroxene isochron yields a cooling age of 338 ± 13 Ma, and $\epsilon_{Nd}$ and ($^{87}\text{Sr}/^{86}\text{Sr})_i$ values for clinopyroxene (at 335 Ma) are +4.5 and 0.7035, respectively.

Keywords: peridotite, LPO, geochemistry, radiogenic isotopes, PT conditions, Moldanubian

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1. Introduction

Fragments of mantle-derived peridotite are a distinctive and widespread feature of convergent orogenic belts around the world, including the Caledonian, Variscan, and Alpine belts in Europe. Although such peridotites are subordinate in volume to surrounding crustal rocks, they commonly provide a wealth of information on properties of the mantle and processes related to incorporation of peridotites into the crust.

The focus of this investigation, the Sklené nad Oslavou peridotite, is a Variscan garnet peridotite boudin enclosed by granulite of the Moldanubian Gföhl Assemblage. Despite its small size (0.7×1.0 m), the Sklené garnet peridotite deserves detailed examination because of its polymetamorphism, distinctive texture, interesting olivine...
lattice-preferred orientation (LPO), and remarkable lack of serpen- 

tinization, features which taken together are different from 

those commonly exhibited by other Czech garnet peri-

dotites. Some preliminary data on the Sklené mineral and 

rock compositions were published previously in an over-

view of Czech garnet peridotites (Medaris et al. 2005), and 

these data are included here for the sake of completeness 

and convenience of the reader. In this detailed investiga-

tion, we present new data on the mineralogy, petrofabrics, 

geochemistry (elemental and isotopic), Sm-Nd cooling age, 

and Re-Os model depletion age of the boudin, followed by 

an interpretation of its evolution from a mantle source to its 

eventual incorporation into a crustal milieu.

2. Geological setting

2.1. Peridotites in the Moldanubian Zone

Numerous bodies of mantle-derived peridotite occur in 

various tectonostratigraphic units in the Moldanubian 

Zone of the Bohemian Massif (Fig. 1a), including the 

Monotonous and Varied groups, the Gföhl Unit and the 

Kutná Hora–Svaratka Terrane, which includes elements 

of the Gföhl Unit (both shown in green in Fig. 1a), the 

Teplá–Barrandian block, and the Moravo-Silesian Terr-

ane (Machart 1984). Most of these ultramafic bodies are 

spinel peridotite and include meta-ophiolite in the 

Teplá–Barrandian block and Moravo-Silesian Terrane 

(Cháb 1973; Jelínek et al. 1984; Kastl and Tonika 1984; 


The Gföhl Unit and its correlatives in the Kutná Hora– 

Svaratka Terrane, however, are unique in containing garnet 

peridotite in addition to spinel peridotite. Elsewhere in 

the European Variscides, garnet peridotite also occurs in 

the Saxothuringian Zone of the Bohemian Massif 

(Schmädicke and Evans 1997) and in Moldanubian 
tectonic units in other Variscan massifs, including the 

Schwarzwald (Kalt et al. 1995; Kalt and Altherr 1996), 

Vosges (Altherr and Kalt 1996), and Massif Central (Las-

nier 1971; Gardien et al. 1990).

The Gföhl Unit is an assemblage of high-grade ortho-

gneiss, paragneiss, migmatite, felsic kyanite-garnet 

granulite, and scattered peridotite bodies, and amphibolite 
occurs at the base of the complex. This lithologic associa-
tion is thought to have originated in a Devonian back-arc 

basin related to the subduction of Saxothuringia beneath 

Teplá–Barrandia and Moldanubia (Carswell 1991; Schul-

mann et al. 2005). Carboniferous closure of the thermally 

weakened back-arc basin, due to convergence and ev-
tual collision of Brunia, led to crustal thickening and 

~340 Ma high pressure-temperature metamorphism in 
dereper crustal levels (the Gföhl Assemblage). Metamor-

phism was followed soon thereafter by rapid exhumation,

the evidence for which is the presence of Gföhl detritus in 

326 to 345 Ma Viséan sediments of the Culm foreland 

basin (Hartley and Otava 2001).

Peridotites in the Gföhl Unit display diverse character-

istics and have been divided into three groups, based on 

their texture, mineralogy, elemental and isotopic chemical 

composition, PT conditions, and calculated cooling rates 
(Medaris et al. 1990, 2005). Type I, e.g. the Mohelno 

peridotite, is surrounded by granulite and consists pre-
dominantly of Mg–Cr spinel peridotite, in which garnet 
occurs only in the margins of the peridotite body within a 
few meters of granulite or in closely associated peridotite 
boudins in granulite. The Mohelno spinel peridotite is 
devoid of eclogite lenses. Type II has been identified in 
the Horní Bory quarry and is represented by a group of 
Mg–Cr to Fe–Ti spinel-garnet peridotite boudins, associ-
ated with abundant garnet pyroxenite, which are enclosed by 
felsic granulite. Type III, e.g. the Nové Dvory peridot-

tite, is hosted by migmatitic gneiss, consists solely of 
Mg–Cr garnet peridotite, and contains prominent lenses of 
eclogite. Type I spinel peridotite is interpreted as frag-
ments of suboceanic lithosphere or asthenosphere, and 
Type III garnet peridotite is most likely derived from 
subcontinental lithosphere (Medaris et al. 2005). Type 
II Fe–Ti peridotite has been modelled as the product of 
reaction between Mg–Cr peridotite and percolating, SiO₂-
undersaturated basaltic melt in a mantle wedge above a 
Variscan subduction zone (Ackerman et al. 2009).

2.1.1. The Sklené peridotite boudin and host granulite

The Sklené garnet peridotite is a solitary, 0.7×1.0 m bou-
din enclosed by felsic granulite (Fiala 1966) of the Bory 
granulite, which is an ~4×9 km body surrounded by a 
variety of high-grade gneisses and migmatites (Fig. 1b). 
The boudin was previously exposed in a railroad cut, but it 
is now completely obscured by colluvium and vegetation. 
The Sklené peridotite and host granulite are likely related to 
the suite of ultramafic boudins and associated felsic 
granulite in the Horní Bory quarry (Ackerman et al. 2009), 
which is located ~2 km WSW of the Sklené locality (Fig. 
1b). On a regional scale, the Bory granulite and associated 
gneisses, which are part of the Gföhl Unit, represent the 
lower levels of orogenically thickened Moldanubian crust 
and now constitute the uppermost structural level of the 
Moldanubian Zone (Schulmann et al. 2005, 2009).

The host felsic granulite consists of quartz, plagioclase, 
alkali feldspar, garnet, and kyanite, and subordinate, more 
mafic varieties of granulite also contain clinopyroxene 
and orthopyroxene (Misař et al. 1984). Retrogression of 
granulite is widespread, yielding garnet–sillimanite or 
garnet–biotite–cordierite assemblages (Kotková et al. 
2003). The garnet peridotite boudin is separated from
granulite by a reaction zone ~2 cm thick, which consists of an inner zone (~1 cm) predominantly of talc, an intermediate zone (~3 mm) of talc, anthophyllite, and hornblende, and an outer zone (~7 mm) of phlogopite. This sequence of minerals resulted from reaction between peridotite and felsic granulite under amphibolite-facies conditions.

3. Petrography

3.1. Texture

The Sklené peridotite has a pronounced inequigranular texture, in which large, rounded, subequant grains of garnet (≤ 7 mm) are surrounded by a much finer-grained, extensively recrystallized matrix of olivine, orthopyroxene, clinopyroxene, and subordinate amphibole (Figs 2–4). Additionally, pentlandite, (Fe,Ni)₉S₈, and heazlewoodite, Ni₃S₂, occur as small, irregular grains at silicate boundaries or as rounded blebs within silicates. The matrix consists of subordinate, larger porphyroclastic grains (~0.5 mm to 1.0 mm) of olivine and pyroxene, which are surrounded by much finer grained (~0.1 mm), recrystallized olivine and pyroxene with predominantly straight to gently curved grain boundaries and 120° triple junctions (Figs 2b, 3b, 4b, 5 and 6). The porphyroclasts have axial ratios of <1:1.5 and are most commonly randomly oriented. However, it is possible to define a weak foliation and lineation in hand specimen, which we subsequently denote as the fabric reference frame, where X is parallel to the lineation and Z is the pole to the foliation. We infer that this reference frame reflects the finite strain ellipsoid within the peridotite. The finer grains have axial ratios of around 1:2, decreasing to <1:1.5 at distances greater than ~350 µm from the garnet porphyroclasts (Fig. 5). Most of these finer grains have their longest axis parallel to fabric X and
Fig. 2 Photomicrographs of the Sklené garnet peridotite, illustrating large rounded garnet grains in a fine-grained, extensively recrystallized matrix of olivine, orthopyroxene, clinopyroxene, and subordinate pargasitic amphibole. Thin section is cut parallel to the X–Z fabric plane. a – Plane polarized light. b – Crossed polarizers. c – Composite kelyphite rim on garnet. Pre- to syn-kinematic pargasite in pressure shadows has a delta-shape, indicating non-coaxial shear. Plane polarized light. d – Oriented, acicular grains of orthopyroxene and amphibole in garnet. Partly crossed polarizers.

Fig. 3 Photomicrographs of the Sklené garnet peridotite. Thin section is cut parallel to the Y–Z fabric plane. a – Plane polarized light. b – Crossed polarizers. c – Pre- to syn-kinematic pargasite in a sigma-shaped pressure shadow of garnet, indicating non-coaxial shear. Sense of shear indicated by the half arrows. Plane polarized light. d – The same view, crossed polarizers.

Fig. 4 Photomicrographs of the Sklené garnet peridotite. Thin section is cut parallel to the X–Y fabric plane. a – Plane polarized light. b – Crossed polarizers. c – Composite kelyphite rim on garnet; note the train of pre- to syn-kinematic pale brownish pargasite grains extending from the kelyphite into the matrix. Plane polarized light. d – The same view, crossed polarizers.
their shortest axis parallel to fabric \( Z \). Within 350 \( \mu m \) of the garnet porphyroclasts, the fine grains are inclined to the main foliation of the rock (the \( X-Y \) plane), and these local foliations wrap around the porphyroclasts (Fig. 5), where they exhibit sigma-shaped pressure shadows (Fig. 3c). Using the linear intercept method (Smith and Guttmann 1953) in \( X-Z \) sections, we estimate that the grain size is approximately 40 \( \mu m \), which can be corrected to a 3-dimensional grain size by multiplication by a factor of 1.75 (e.g. van der Wal et al. 1993), yielding an estimated average recrystallized grain size of 70 \( \mu m \).

Garnet grains are surrounded by thin rims of kelyphite, which consist of an outer zone of monomineralic pargasitic amphibole and an inner zone of extremely fine-grained, intergrown spinel and orthopyroxene (Fig. 7). The inner kelyphite zone is confined to the margins of garnet, but the kelyphitic amphibole is drawn out into the matrix (Fig. 2c), parallel to the \( X \) fabric direction (Figs 2c and 4c), and some kelyphitic tails define a delta geometry (Figs 2c, 4c). Small amounts of very fine-grained amphibole are also dispersed in the recrystallized matrix. Significantly, spinel and garnet are absent from the matrix.

Individual grains of olivine, orthopyroxene, or clinopyroxene occur rarely as inclusions in garnet, as seen in Fig. 4b. Some, but not all, garnet grains contain clusters of tiny, oriented, acicular grains of orthopyroxene and amphibole (Fig. 2d). Such clusters are not uniformly distributed within garnet grains, but occur instead in small domains near grain margins.

### 3.2. Mineral compositions

Minerals in the Sklené peridotite are typical of those in other mantle-derived Mg–Cr garnet peridotites, including Mg-rich olivine and orthopyroxene, Cr-diopside, and pyrope-rich garnet. Compositions of the constituent minerals were determined by electron probe microanalysis, the analytical details of which are given in the Appendix. Because of the large uncertainty in calculating \( \text{Fe}^{2+} \) and \( \text{Fe}^{3+} \) by stoichiometry in silicate minerals with low \( \text{Fe}/\text{Mg} \) ratios, as is the case here, all \( \text{Fe} \) has been taken as \( \text{FeO} \), which is a convention commonly followed in studies of Mg–Cr peridotites.

The composition of olivine is relatively uniform on both the intragrain and intergrain scales. Olivine porphyroclasts (Table 1) have a mean Mg\# [100×Mg/(Mg + Fe)] of 89.5 and contain 0.40 wt. % NiO.

Garnet (Table 2) is relatively uniform in composition, except within ~25 micrometers of grain margins, where there is a marked decrease in Mg\# from 81.5 to 74.7 (illustrated by the bright band at the garnet margin in Fig. 7b). Garnet is pyrope-rich, and the molar proportions of almandine : spessartine : pyrope : grossular vary little from the cores, 16.3 : 0.6 : 73.3 : 9.9, to intermediate
Fig. 6 Photomicrographs of the recrystallized matrix in the Skleně peridotite. Thin section is cut parallel to the X–Z fabric plane; crossed polarizers. a – Two relatively coarse, porphyroclastic olivine grains in the recrystallized matrix. b – Several relatively coarse, porphyroclastic orthopyroxene grains in the recrystallized matrix.

zones, 16.6 : 0.6 : 72.9 : 9.9, of the large grains. Garnet contains ~0.9 wt. % Cr$_2$O$_3$, except for the cores of a few grains, where Cr$_2$O$_3$ reaches 2.2 wt. %, and Mg# attains a maximum value of 82.1.

The composition of orthopyroxene (Table 3) is similar for most elements among the cores of larger grains, their rims, and matrix grains, all having a Mg# of 90.6–90.8 and CaO contents of 0.35–0.37 wt. %. However, there is a slight, but significant, increase in Al$_2$O$_3$ from the cores of larger grains, 1.02 wt. %, to rims, 1.24 wt. %, to matrix grains, 1.45 wt. %, which is accompanied by an increase in Cr$_2$O$_3$ from 0.10 to 0.14 to 0.17 wt. %. Compared to exterior grains, the tiny, acicular grains of orthopyroxene in garnet (Fig. 2d) have a higher Mg#, 92.3, and apparently higher Al$_2$O$_3$ content, 3.3 wt. %, reflecting the chemical influence of the enclosing garnet, i.e., exchange of Fe and Mg between orthopyroxene and garnet during cooling in the former parameter and fluorescence effects of surrounding garnet in the latter.

The compositional variation in clinopyroxene (Table 3) is similar to that in orthopyroxene. The cores and rims of larger grains and matrix grains have comparable Mg# numbers, 92.0–92.6, and CaO contents, 20.27–20.43 wt. %, whereas Al$_2$O$_3$ increases from the cores of larger grains to rims to matrix grains, from 3.62 to 3.84 to 4.00 wt. %, as does Cr$_2$O$_3$, from 0.64 to 0.80 to 0.81 wt. %.

Amphibole (Table 4) in kelyphite is pargasite, with 6.3 atoms of Si per formula unit, 1.66 atoms of Ca in the B-
Moldanubian garnet peridotite

Tab. 1 Electron microprobe analysis: olivine porphyroclasts

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<th></th>
<th>mean</th>
<th>1 S.D.**</th>
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<tr>
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<td>cations per 4 oxygen atoms</td>
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<td>Fe</td>
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<tr>
<td>Ni</td>
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* Total Fe as FeO
** n – number of analyses; S.D. – standard deviation

Tab. 2 Electron microprobe analyses of garnet

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<td>8.014</td>
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</table>

* Total Fe as FeO
** hi-Cr – high chromium; lo-Cr – low chromium; Alm – almandine; SpS – spessartine; Prp – pyrope; Grs – grossular

3.3. Mode
The modal composition of the Sklené peridotite was calculated by a linear least squares method (Wright and Doherty 1970), utilizing its constituent mineral compositions (Tables 1–3) and bulk rock composition (Table 5). Ignoring the small amounts of pargasite and spinel in the sample, the Sklené peridotite is a lherzolite, which contains 55 % olivine, 13 % garnet, 22 % orthopyroxene, and 10 % clinopyroxene.

4. Petrofabrics
The petrofabrics (lattice-preferred orientation, LPO) and microstructures of olivine, orthopyroxene and clinopyroxene in the matrix of the Sklené peridotite were characterized using electron backscatter diffraction, as described in the Appendix. Representative microstructural maps and all measured LPOs are illustrated in Figs 5 and 8. Crystallographic orientations of both porphyroclasts and recrystallized grains are included in these diagrams, but since the latter are volumetrically dominant and only one point per grain is represented on the fabric diagrams, the fine grains contribute the majority of data to the measured petrofabrics.

The LPOs of pyroxenes and olivine were measured in two thin sections, in the X–Z and X–Y fabric planes. The resulting LPOs of all phases are relatively weak, reflected in M-indices (Skemer et al. 2005) < 0.1 and maximum m.u.d. (multiples of uniform density) < 3. In both pyroxenes, where there is a resolvable LPO, the normals to (100) in clinopyroxene and [100] in orthopyroxene are randomly distributed, both phases have axial concentrations of [010] about the Z fabric axis, and [001] are distributed in weak girdles in the X–Y fabric plane (Fig. 8b, d). Olivine has girdles of [100] and [001] in

site, 0.58 atoms of Na + K in the A-site, and an Mg# of 90.0. The scarce amphibole in the matrix is also pargasitic, but with a slightly lower Al₂O₃ content, 12.45 wt. %, higher CaO content, 11.52 wt. %, and higher Mg#, 91.5. The tiny, acicular grains of amphibole in garnet (Fig. 2d) are also pargasitic, but with slightly higher Al₂O₃, 17.07 wt. %, and Mg#, 92.8, compared to exterior amphibole, which reflects the chemical influence of encasing garnet, as also observed for orthopyroxene.

Spinel (Table 4), as noted above, occurs only as an extremely fine-grained intergrowth with orthopyroxene in inner kelyphite rims (Fig. 5). The spinel is Mg-rich and Cr-poor, having a Mg# of 72 and a Cr# [100×Cr/(Cr + Al)] of 11. Such a composition is typical of spinel, which occurs in kelyphite in garnet peridotites and has formed through the reaction, garnet + olivine → spinel + orthopyroxene.
Tab. 3 Electron microprobe analyses of pyroxene

<table>
<thead>
<tr>
<th></th>
<th>Orthopyroxene</th>
<th>Clinopyroxene</th>
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<tr>
<td></td>
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<td>p.clast matrix</td>
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<tr>
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<td>grains</td>
</tr>
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<td>n = 2</td>
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</tr>
<tr>
<td>Al₂O₃</td>
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</tr>
<tr>
<td>Cr₂O₃</td>
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</tr>
<tr>
<td>FeO*</td>
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<td>6.44</td>
</tr>
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<td>0.11</td>
</tr>
<tr>
<td>MgO</td>
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<td>34.97</td>
</tr>
<tr>
<td>CaO</td>
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<td>0.35</td>
</tr>
<tr>
<td>Na₂O</td>
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<td>0.01</td>
</tr>
<tr>
<td>Sum</td>
<td>100.17</td>
<td>100.03</td>
</tr>
</tbody>
</table>

Cations per 6 oxygen atoms

| Si            | 1.961         | 1.958         | 1.953         | 1.909         | 1.974         | 1.961         | 1.956         |
| Ti            | 0.003         | 0.001         | 0.002         | 0.134         | 0.008         | 0.009         | 0.008         |
| Al            | 0.041         | 0.050         | 0.059         | 0.003         | 0.153         | 0.163         | 0.170         |
| Cr            | 0.003         | 0.004         | 0.004         | 0.003         | 0.018         | 0.023         | 0.023         |
| Fe            | 0.183         | 0.186         | 0.186         | 0.151         | 0.074         | 0.069         | 0.072         |
| Mn            | 0.003         | 0.003         | 0.003         | 0.001         | 0.002         | 0.002         | 0.002         |
| Mg            | 1.807         | 1.799         | 1.792         | 1.811         | 0.856         | 0.856         | 0.859         |
| Ca            | 0.013         | 0.013         | 0.014         | 0.009         | 0.782         | 0.790         | 0.783         |
| Na            | 0.001         | 0.000         | 0.001         | 0.001         | 0.126         | 0.132         | 0.130         |
| Sum           | 4.015         | 4.014         | 4.014         | 4.021         | 3.995         | 4.004         | 4.004         |

| Ca            | 0.7           | 0.6           | 0.7           | 0.5           | 45.7          | 46.1          | 45.7          |
| Mg            | 90.2          | 90.1          | 90.0          | 91.9          | 50.0          | 49.9          | 50.1          |
| Fe            | 9.1           | 9.3           | 9.3           | 7.7           | 4.3           | 4.0           | 4.2           |
| Mg#           | 90.8          | 90.6          | 90.6          | 92.5          | 92.0          | 92.6          | 92.2          |

* Total Fe as FeO
** p.clast – porphyroclast

the X–Y fabric plane and axial concentrations of [010] parallel to the Z fabric axis (Fig. 8e–f).

5. Geochemistry

Major and minor elements in the Sklené peridotite were determined by wet chemical analysis, and trace elements in whole-rock and mineral separates (clinopyroxene and garnet) were obtained by Inductively Coupled Plasma – Mass Spectrometry (ICP-MS). The Sr and Nd isotopes were measured by Thermal Ionization Mass Spectrometry (TIMS). Rhenium, Ir, Ru, Pt, and Pd were analyzed using multi-collector ICP-MS, with concentrations determined by isotopic dilution. Osmium concentrations and isotopic composition were measured by Negative Thermal Ionization Mass Spectrometry (N-TIMS). Details for each of these analytical techniques are given in the Appendix.

5.1. Major and minor elements

Consistent with its lherzolitic mode, the Sklené peridotite has a modest concentration of MgO, 37.7 wt. %, and is only slightly depleted in incompatible major and minor elements, containing 3.31 wt. % Al₂O₃, 0.18 TiO₂, 2.74 CaO, and 0.27 Na₂O (Table 5). In oxide variation diagrams (Fig. 9) the Sklené sample plots closer to the composition of primitive mantle (McDonough and Sun 1995) than do most analyzed Czech peridotites, which display a pronounced depletion trend. Although the Sklené peridotite is located geographically closest to the Type II peridotite suite in the Horní Bory quarry, its major-element composition lies between the major element compositional fields for Type I and Type III peridotites (Fig. 9).

5.2. Trace elements

5.2.1. Whole rock

The rare earth element (REE) contents of the Sklené peridotite are depleted with respect to primitive mantle (McDonough and Sun 1995), with normalized values ranging from ~0.3 to 0.8 (Fig. 10a, Table 5). The light REE (LREE) are slightly depleted compared to the
Table 4: Electron microprobe analyses of amphibole and spinel

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<tr>
<th></th>
<th>Amphibole</th>
<th>Spinel</th>
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<tr>
<td></td>
<td>lamellae</td>
<td>kelyphite</td>
</tr>
<tr>
<td></td>
<td>in Grt</td>
<td></td>
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<td>97.38</td>
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Cations per 23 oxygen atoms

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<th>Cr</th>
<th>Fe</th>
<th>Mn</th>
<th>Mg</th>
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<tbody>
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<td>0.716</td>
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<tr>
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<td>0.000</td>
<td>0.052</td>
<td>0.051</td>
<td>0.022</td>
<td>0.001</td>
<td>0.021</td>
</tr>
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<td>Sum</td>
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<td>Mg#</td>
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<tr>
<td>Cr#</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>15.794</td>
<td>15.816</td>
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<td></td>
<td></td>
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<tr>
<td>Mg#</td>
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<td>90.0</td>
<td>91.5</td>
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</tbody>
</table>

5.2.2. Clinopyroxene and garnet

The normalized REE patterns for clinopyroxene and garnet are typical for those in garnet lherzolite, i.e., LREE enriched and slightly convex upward in clinopyroxene, and HREE enriched and LREE depleted in garnet, reflecting the preferential accommodation of LREE in clinopyroxene and HREE in garnet. The present REE compositions of the Sklené garnet and clinopyroxene most likely represent high-temperature, subsolidus re-equilibration between the two phases, rather than their original compositions resulting from partial fusion and melt extraction. Compared to clinopyroxene in Type II-Mg peridotite, the Sklené clinopyroxene has a slightly higher Ce₃/Yb₃ ratio, i.e., 6.8 vs. 3.8 (Fig. 12a).

With respect to normalized, extended trace elements, the Sklené clinopyroxene exhibits a pronounced negative Nb anomaly, a marked positive Pb anomaly, and slight negative anomalies for Zr, Hf, and Ti (Fig. 12b). Similar anomalies also occur in clinopyroxene from Type II-Mg peridotite (Fig. 12b). The Sklené garnet also shows

heavy REE (HREE), with a La₃/Yb₃ ratio of 0.71, but the most distinctive feature of the normalized REE pattern is its pronounced convex upward shape, reflecting the relatively higher normalized values of the middle REE (Fig. 10a). The REE patterns may be compared in a plot of Ce₃/Sm₃ vs. Sm₃/Yb₃ (Fig. 11), in which different configurations of REE patterns are separated into four fields, i.e. LREE depleted, LREE enriched, convex upward, and convex downward. Among Czech garnet peridotites, Type II-Fe peridotites, which display convex upward REE patterns, are the group most similar to the Sklené sample with respect to REE pattern, although one sample of Type III peridotite plots close to the Sklené sample.

In a plot of normalized, extended trace elements, the Sklené peridotite exhibits pronounced positive anomalies for Sr and U and elevated values for Rb and Th (Fig. 10b). Such features also occur in Type II-Mg peridotites (Fig. 10b) and in Type II-Fe peridotites (not figured). However, numerous samples of Type II-Mg and -Fe peridotites exhibit negative anomalies for Zr and Hf, which are not seen in the Sklené sample.
Tab. 5 Major-element composition of the Sklené garnet lherzolite and trace-element compositions of rock, clinopyroxene, and garnet

<table>
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<th>wt. %</th>
<th>rock ppm</th>
<th>cpx ppm</th>
<th>grt ppm</th>
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<td>SiO₂</td>
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<td>0.08</td>
<td>0.18</td>
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<tr>
<td>TiO₂</td>
<td>0.18</td>
<td>29</td>
<td>220</td>
<td>1</td>
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<tr>
<td>Al₂O₃</td>
<td>3.31</td>
<td>2.50</td>
<td>3.15</td>
<td>13.25</td>
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<tr>
<td>Cr₂O₃</td>
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<td>6.58</td>
<td>12.25</td>
<td>22.84</td>
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<td>0.07</td>
<td>0.23</td>
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<td>FeO*</td>
<td>8.93</td>
<td>0.03</td>
<td>0.07</td>
<td>0.01</td>
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<tr>
<td>MnO</td>
<td>0.13</td>
<td>20</td>
<td>0.25</td>
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</tr>
<tr>
<td>MgO</td>
<td>37.70</td>
<td>0.225</td>
<td>1.78</td>
<td>0.03</td>
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<tr>
<td>CaO</td>
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<td>0.669</td>
<td>5.46</td>
<td>0.10</td>
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<td>Na₂O</td>
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<td>LOI**</td>
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<td>0.126</td>
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<td>0.418</td>
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Tab. 6 Rb–Sr and Sm–Nd isotopic data for the Sklené garnet lherzolite

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<tr>
<th>Sample</th>
<th>Rb (ppm)</th>
<th>Sr (ppm)</th>
<th>87Rb/86Sr</th>
<th>87Sr/86Sr (i)</th>
<th>Sm (ppm)</th>
<th>Nd (ppm)</th>
<th>147Sm/144Nd</th>
<th>143Nd/144Nd</th>
<th>εNd (i)</th>
<th>Age (Ma)</th>
<th>TDM (Ga)</th>
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<td>clinopyroxene</td>
<td>0.01</td>
<td>232</td>
<td>0.001</td>
<td>0.7035</td>
<td>0.7035</td>
<td>1.313</td>
<td>4.522</td>
<td>0.1755</td>
<td>+4.5</td>
<td>338 ± 13</td>
<td>0.70</td>
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<td>garnet</td>
<td>n.a.</td>
<td>n.a.</td>
<td>n.a.</td>
<td>n.a.</td>
<td>0.629</td>
<td>0.588</td>
<td>0.6464</td>
<td>0.513874</td>
<td>+5.2</td>
<td>338 ± 13</td>
<td>0.65</td>
</tr>
<tr>
<td>bulk rock</td>
<td>n.a.</td>
<td>n.a.</td>
<td>n.a.</td>
<td>n.a.</td>
<td>0.265</td>
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<td>0.512933</td>
<td>+4.7</td>
<td>338 ± 13</td>
<td>0.69</td>
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5.3. Sr-Nd geochemistry

5.3.1. Sm-Nd geochronology

The Sklené peridotite yields a well-defined Sm–Nd isochron age of 338 ± 13 Ma for clinopyroxene, garnet, and whole rock, with an initial 143Nd/144Nd ratio of 0.512443 and corresponding εNd value of −3.8 (Fig. 13, Table 6). This age is the same within error of the mean Sm–Nd age, 336 ± 7 Ma, for nine samples of peridotite-hosted pyroxenite and eclogite elsewhere in the Gföhl Assemblage (Medaris et al. 2006a). Such ages are interpreted to represent the time of isotopic closure for these high temperature minerals. These closure ages may be related to the rapid exhumation of the Gföhl Unit, which cooled at a rate of 100 to 150 ºC/m.y., as determined by diffusional modelling of compositionally zoned minerals in the Spačice eclogite (Medaris et al. 2006b) and 40Ar/39Ar cooling ages of 329–341 Ma for hornblende and 326–329 Ma for muscovite in units overridden by the Gföhl Unit (Dallmeyer et al. 1992).

5.3.2. Sr and Nd isotopes

The Sklené clinopyroxene has an initial εNd value of +4.5 and an initial 87Sr/86Sr ratio of 0.7035 (both calculated at 335 Ma) (Table 6). These values are comparable to those for some samples of Type III peridotite, and they

Tab. 7 HSE and Re–Os isotopic data for the Sklené garnet lherzolite

<table>
<thead>
<tr>
<th>Sample</th>
<th>Re (ppb)</th>
<th>Os (ppb)</th>
<th>Ir (ppb)</th>
<th>Ru (ppb)</th>
<th>Pt (ppb)</th>
<th>Pd (ppb)</th>
<th>187Re/188Os</th>
<th>187Os/188Os</th>
<th>γOs</th>
<th>Tvs (Ma)</th>
<th>TDM (Ga)</th>
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<tr>
<td>CS-SK-1 WR</td>
<td>0.152</td>
<td>3.68</td>
<td>3.34</td>
<td>6.76</td>
<td>5.72</td>
<td>1.78</td>
<td>0.1997</td>
<td>0.12425</td>
<td>-1.3</td>
<td>811</td>
<td>0.9</td>
</tr>
</tbody>
</table>

γOs values calculated back to 335 Ma. Tvs represents Re depletion age from PUM reservoir (see Shirey and Walker 1998 for details on calculations and Meisel et al. 1996 for PUM values)
5.4. Highly siderophile element geochemistry

Highly siderophile elements (HSE) in the Sklené peridotite, when normalized to the primitive upper mantle (PUM) of Becker et al. (2006), exhibit an unfractionated pattern for the Iridium group PGE (I-PGE: Os, Ir, Ru) and Pt, but a significant depletion in Pd and Re (Fig. 15, Table 7). This pattern reflects the presence of pentlandite and late-stage heazlewoodite in the Sklené peridotite and their influence on HSE contents. Pentlandite commonly exhibits an unfractionated I-PGE pattern, but significant depletions in Pt and Pd (Luguet et al. 2001). In general, the Sklené HSE pattern is similar to that in Type II–Mg peridotites from Horní Bory, except for a lower Pd content. In contrast, the HSE pattern in Type II Fe peridotites from Horní Bory is quite different, showing marked depletion in the I-PGE and a highly fractionated pattern for Pt, Pd, and Re (Fig. 15), similar to that of basalts and eclogites.

The $^{187}$Os/$^{188}$Os isotopic ratio of the Sklené lherzolite is 0.1242 ($\gamma$Os = −3.1). This value is similar to those for many samples of European subcontinental lithospheric mantle (Meisel et al. 2001; Pearson et al. 2004; Reisberg et al. 2005), but lower compared to PUM (0.1290; Meisel et al. 1996) and estimates for convecting mantle (0.1281; Walker et al. 2002).

6. Evolution of the Sklené peridotite

6.1. Pressure-temperature history

The texture of the Sklené peridotite reveals three stages of mineralogical development, including an early garnetiferous stage (1), an intermediate stage (2) of matrix recrystallization and deformation accompanied by growth of amphibole in kelyphite and matrix, and a final stage (3) of spinel–orthopyroxene kelyphite development.
The conventional method for estimating pressure–temperature (PT) conditions for garnet peridotite is to apply the olivine–garnet Fe–Mg exchange geothermometer and the Al-in-orthopyroxene barometer to the core compositions of garnet, olivine, and orthopyroxene. The rationale for this method is that Fe–Mg diffusion is relatively slow in garnet, Al diffusion is slow in orthopyroxene, and olivine, because it is the predominant phase in peridotite, will undergo the least compositional change during recrystallization and/or cooling (Brenker and Brey 1997). Note, however, that combination of an exchange reaction and a net transfer reaction will yield meaningful results only if the garnet and orthopyroxene compositions equilibrated at the same time and both were preserved from the PT conditions of interest.

The PT conditions for Stage 1 of the Sklené peridotite were estimated by applying the O’Neill and Wood (1979) calibration of the olivine–garnet Fe–Mg exchange geothermometer (corrected by O’Neill 1980) and two calibrations of the Al-in-orthopyroxene geobarometer (Nickel and Green 1985; Brey and Köhler 1990) to the core compositions of garnet, coarser grains of matrix olivine, and coarser grains of matrix orthopyroxene. Results from the two different geobarometer calibrations are similar, 1229 °C, 49.1 kbar (Brey and Köhler 1990) and 1242 °C, 52.2 kbar (Nickel and Green 1985), and such results are comparable to those for some Czech Type III garnet peridotites (Fig. 16).

The Stage 1 PT estimate for the Sklené peridotite, which lies in a subsolidus regime, is just within the stability field for majoritic garnet (Fig. 16), raising the possibility that the scarce clusters of tiny, acicular grains of orthopyroxene and pargasite in some garnet grains (Fig. 2d) may have originated by exsolution from a majoritic garnet. Similar features have been described in several ultrahigh pressure garnet peridotites elsewhere, e.g. pyroxene lamellae in Norwegian garnet peridotites have been ascribed to pyroxene exsolution from supersilicic garnet (Van Roermund et al. 2001), and pargasite lamellae in Tibetan garnet peridotites are proposed to have originated by amphibole exsolution from supersilicic, OH-bearing garnet (Song et al. 2005). Contents of H₂O ranging from <1 to 290 ppm have been reported in garnet from garnet peridotite xenoliths in kimberlite (Beran and Libowitzky 2006), and if the Sklené garnet contained 200 ppm H₂O, it could account for 0.9 wt. % exsolved
Fig. 10 Whole-rock REE (a) and extended trace element (b) compositions of the Sklené peridotite, normalized to primitive mantle (McDonough and Sun 1995). The compositional fields for Czech Type II Mg peridotites are shown for comparison.

Fig. 11 Ratios of CeN/SmN vs. SmN/YbN for the Sklené peridotite compared to those for other types of Czech peridotites, all normalized to primitive mantle (McDonough and Sun 1995).

Note, however, that the clusters of pyroxene and pargasite in the Sklené garnet are uncommon, occur near the margins of garnet grains, and consist of minerals that are compositionally similar to those in adjacent kelyphite. Alternatively therefore, it is possible that the clusters of orthopyroxene and pargasite in garnet are related to the introduction of H₂O into peridotite from the surrounding granulite, development of kelyphite around garnet, and incipient, topotaxial growth of kelyphite minerals in garnet, rather than exsolution from a supersilicic, OH-bearing garnet. Regardless of the source of H₂O for the growth of pargasite within garnet, such growth must have occurred in the stability field for amphibole at less than ~29 kbar (Fig. 16), rather than in the majorite field.

Stage 2 in the Sklené sample is represented by the recrystallized matrix, which consists of olivine, orthopyroxene, clinopyroxene, and subordinate pargasite, grains of which are scattered through the matrix and, locally, strung out from kelyphitic rims into the foliation plane of the matrix (Figs 2c and 4c). The presence of pargasite constrains the pressure of Stage 2 to be less than ~29 kbar (Fig. 16), based on the stability field of pargasite in hydrous peridotite (Schmidt and Poli 1998; Fumagalli and Poli 2005). The temperature for Stage 2 has been estimated by the application of three two-pyroxene geothermometers (Bertrand and Mercier 1985; Brey and Köhler 1990; Taylor 1998) to the mean compositions of fine-grained matrix pyroxenes. All three geothermometers yield similar results, the averages of which range from 900 ± 39 °C at 15 kbar to 919 ± 34 °C at 25 kbar. Based on these constraints and the absence of spinel in the recrystallized matrix, the PT conditions for Stage 2 are judged to be ~915 °C and ~24 kbar (Fig. 16). This pres-
Fig. 12 The REE (a) and extended trace element (b) compositions of clinopyroxene and garnet in the Sklené peridotite, normalized to primitive mantle (McDonough and Sun 1995). The compositional fields for clinopyroxene in Czech Type II Mg peridotites are shown for comparison.

6.2. Mantle geochemical processes

Many samples of mantle peridotite commonly record the effects of two geochemical processes: 1) initial depletion of incompatible elements by partial melting, and 2) subsequent modification of certain elements and isotopic ratios by reaction of peridotite with infiltrating melts or fluids (e.g., basaltic melts, carbonatitic melts, or fluids, either H₂O-rich or CO₂-rich).

Although the Skleně peridotite is a relatively fertile lherzolite, which contains 10% clinopyroxene, its Al₂O₃, CaO, Na₂O, TiO₂, and REE contents, when compared to those of primitive mantle (Table 5; Figs 9 and 10a), indicate that the Skleně peridotite experienced a small amount of partial fusion and depletion in incompatible elements. A quantitative calculation of the degree of depletion, based on the Yb and Y contents of clinopyroxene in equilibrium with melt (Norman 1998), is pre-
Moldanubian garnet peridotite

Fig. 14 $\varepsilon_{\text{Nd}}$ vs. ($^{87}\text{Sr}/^{86}\text{Sr}$) for Sklené clinopyroxene compared to clinopyroxene in other types of Czech peridotites and the array defined by clinopyroxene in peridotite-hosted pyroxenites and eclogites (Ackerman et al. 2009, and references therein). Mantle components, DMM, EM1, and EM2, from Harte (1988). All data calculated at 335 Ma.

Fig. 15 The HSE pattern of Sklené peridotite, normalized to primitive upper mantle (PUM; Becker et al. 2006). The HSE compositions of Type II Mg and Type II Fe peridotites are shown for comparison (Ackerman, unpublished data).

Fig. 16 Pressure–temperature (PT) estimates for the three textural and mineralogical stages of the Sklené peridotite. PT estimates for garnetiferous assemblages in other types of Czech garnet peridotites are shown for comparison. Mineralogical stability fields from Fumagalli and Poli (2005) and Poli and Schmidt (1998); upper pressure limit of spinel calculated from O’Neill (1981); majorite isopleth from Gasparik (2003); anhydrous and hydrous (200 ppm H$_2$O) solidi for fertile lherzolite from Green and Falloon (2005).

cluded, however, because of the likely modification of initial clinopyroxene composition by high-temperature, subsolidus re-equilibration with garnet. Qualitatively, the low normalized values, ~0.5, for the HREE in the Sklené peridotite (Fig. 10a) indicate that partial fusion and melt extraction may have occurred in the spinel stability field, prior to stabilization of garnet at high pressures, otherwise the HREE contents in the peridotite residuum would have remained near a value of 1.0 in the presence of garnet during low to moderate degrees of partial fusion.

Compared to primitive mantle, the Sklené peridotite has a lower Mg#, 88.3 vs. 89.3, and higher contents of (and positive anomalies for) Th, U, and Sr (Fig. 10b). Clinopyroxene and garnet also have positive anomalies for Th and U and pronounced positive anomalies for Pb (Fig. 12b), and clinopyroxene has a marked negative Nb anomaly (Fig. 12b). With respect to the HSE in peridotite,
Fig. 17 Deformation mechanism maps for (a) dry olivine and (b) wet olivine (200H/10^6Si) at 915 °C and 2400 MPa, as experienced during Stage 2 deformation. The maps were constructed using constitutive equations described by Warren and Hirth (2006). The estimated mean matrix grain size, 70 µm, is indicated by the vertical dashed line. disGBS = dislocation-accommodated grain boundary sliding.

the normalized content of Re is greater than that for Pd (Fig. 15), despite Re being less compatible than Pd (e.g., Pearson et al. 2004). All of these features indicate that the Skleně peridotite experienced some degree of cryptic metasomatism, with addition of Fe, Th, U, Pb, Sr, and Re, coupled with removal of Nb from clinopyroxene.

Although it appears that the Skleně peridotite has been enriched in Sr, there is no evidence for this in the 87Sr/86Sr ratio of its clinopyroxene, which is only slightly higher, 0.7035, than that for depleted MORB mantle (DMM; Fig. 14). We suggest that the metasomatic Sr was relatively unradiogenic, having been derived from an isotopically primitive or depleted mantle source, thereby having little effect on the 87Sr/86Sr composition of the Skleně clinopyroxene. The Skleně Os isotopic composition is also relatively unradiogenic (γOs = −1.3), indicating little or no addition of radiogenic Os.

In summary, the geochemical characteristics of the Skleně peridotite are consistent with some degree of cryptic metasomatism, most likely produced by transient, primitive mantle-derived melts of basaltic composition. In contrast to the neighbouring Horní Bory ultramafic suite (Ackerman et al. 2009), there is no evidence for subduction-related metasomatism and melt-rock reaction in the Skleně peridotite.

Although partitioning of Re between crystals and silicate melt is dependent on oxygen fugacity, Re behaves as a moderately to strongly incompatible element in the range of oxygen fugacities commonly assigned to the mantle, e.g. Δ log fO2 (QFM) = −2 to +2 (Mallman and O’Neill 2007). Because Os is a compatible element, Re/Os ratios in peridotite residua will decrease during partial fusion, which offers a method for estimating the age of melt extraction, provided that neither Re nor Os of different isotopic compositions were subsequently added to the system. The model Re-depletion age (Tpd; Table 7; Walker et al. 1989) for the Skleně peridotite is 0.9 Ga. However, because of the likely metasomatic addition of Re, this age is probably a minimum estimate for the time of melt extraction. The Tpd age for Skleně is within the large range (0.3–2.0 Ga) of those for several other European orogenic peridotites (e.g., Ronda, Lherz, and Lanzo; Becker et al. 2006), which show evidence for mantle refertilization, but slightly lower than those (1.0–1.5 Ga) for most Lower Austrian peridotites (Becker et al. 2001).

6.3. Deformation and recrystallization

The textural characteristics of the matrix of the Skleně peridotite developed during Stage 2 deformation, coeval with or postdating formation of the pargasitic kelyphite, which has been drawn out into the matrix. The PT conditions for deformation are therefore ~900 °C and ~24 kbar, corresponding to a depth of ~80 km, assuming a 30 km thick crust with ρ = 2750 kg/m³ underlain by mantle with ρ = 3200 kg/m³. The presence of recognizable LPOs in olivine, orthopyroxene and clinopyroxene indicate that all these phases deformed in part by dislocation-accommodated creep.

The petrofabrics are relatively weak, reflected in their low M-indices (cf., Skemer et al. 2005). This
pattern could have several possible causes, including: 1) the peridotites have experienced insufficient strain for strong fabrics to form; 2) initially strong fabrics have been weakened, due either to post-deformational static annealing (e.g., Wightman et al. 2006) or activation of grain size sensitive deformation mechanisms (Skemer and Karato 2008); or 3) the type of kinematics does not produce strong petrofabrics.

The average matrix grain size, 70 µm, is consistent with the operation of grain size sensitive creep for reasonable geological strain rates (<10^{-10} s^{-1}; Pfiffner and Ramsay 1982) for both dry and wet olivine (200 H/10^6 Si), as illustrated in Fig. 17. The relatively polygonal matrix microstructures, including straight grain boundaries and 120° triple junctions, are consistent with either low total strain or post-deformational annealing, but the straight grain boundaries would also facilitate grain boundary sliding. One possible interpretation of these various microstructural and textural data is a history of changing deformation mechanisms during progressive deformation. In this scenario, initial grain size reduction and formation of a porphyroclast and matrix microstructure occurred by dislocation creep. Once grain size was reduced sufficiently by this process, however, the rock began to deform by grain size sensitive creep (cf. Warren and Hirth 2006; Skemer and Karato 2008; Toy et al. in print).

The observed pyroxene LPOs (Fig. 8) are consistent with glide on the (010) plane during the initial dislocation creep phase, which is not typically reported for pyroxenes (except by Toy et al. in print, and in omphacite in natural eclogites; Bascou et al. 2002). The girdles of [001] in the X–Y plane are similar to S-type petrofabrics of Helmstedt et al. (1972), which may form due to glide parallel to this crystallographic axis during a flattening-dominated deformation, producing an oblate strain ellipsoid (Ulrich and Mainprice 2001).

Olivine LPOs (Fig. 8) are consistent with glide on (010) parallel to [100] and [001]. Such LPOs differ from typical olivine patterns diagnostic of (010)[100] and (010) [001] glide during dislocation creep, which would have concentrations of [100] and [001], respectively, parallel to fabric X. Sundberg and Cooper (2008) also reported similar LPOs in fine-grained olivine–orthopyroxene aggregates experimentally deformed by diffusion creep in general shear. However, we think it is most likely that the observed LPOs were developed during an initial stage of dislocation creep, because of the porphyroclast and matrix microstructure and matrix grain elongation. Consequently, we suggest that the observed olivine LPOs were developed in a flattening-dominated deformation regime.

(010)[100] is the slip system most commonly reported to have been activated in olivine under upper mantle conditions in naturally-deformed peridotites (e.g. van der Wal and Vissers 1993; Dijkstra et al. 2001, 2002) and in experimental studies under relatively low differential stresses and low water contents (e.g. Carter and Ave’Lallemand 1970; Jung and Karato 2001). With higher water contents, we might expect olivine LPO diagnostic of glide on the system (010)[001] (Type B fabric of Jung and Karato 2001). During Stage 2 deformation of the Sklené peridotite, water contents must have been high enough to stabilize pargasite, suggesting that conditions were consistent with those necessary to develop a Type B fabric. The other alternative is that the observed fabrics reflect the kinematics of deformation; numerical simulations of Tommasi et al. (1998) illustrated that different LPO geometries are expected for different magnitudes and types of deformation. Thus, the LPOs for Sklené olivine, which are consistent with both Type A and Type B fabrics, could be the result of either: 1) water content causing activation of different slip systems in a simple shear deformation; 2) three-dimensional kinematics; or 3) some combination of the two.

At lower temperatures, glide in olivine is expected to occur on the system {0kl} [100] (Carter and Ave’Lallemand 1970). The absence of LPOs diagnostic of operation of this slip system, and the presence of static Stage 3 kelyphite rims formed at ~900 °C and 17–21 kbar, indicate that the Sklené peridotite was not substantially deformed after it was incorporated into the Bory granulite.

Grain shape fabrics are more strongly developed immediately adjacent to garnet porphyroclasts, indicating that matrix strain was focussed in these areas, probably to maintain constant bulk strain rate around the relatively rigid garnets. The kelyphitic pargasite trails define a delta geometry (Figs 2c, 4c), indicating that the rigid garnet porphyroclasts rotated, consistent with non-coaxial deformation during Stage 2 deformation. Although not well developed, shear sense indicators are present as denoted by larger grain size in the strain shadow areas, presence of fine-grained material along the garnet’s margins, and local imbrication of garnets. Shear sense indicators are best developed with a consistent sense of shear in the Y–Z plane. This type of pattern is expected for pure-shear dominated transpressional deformation (e.g., Fossen and Tikoff 1993). Numerical modelling of olivine LPO development in transpression deformation also shows results that are consistent with the observed patterns (Tommasi et al. 1998). Consequently, we interpret that the Sklené peridotite was deformed during “transpressional” deformation (widening/thinning shear of Tikoff and Fossen 1993) that produced an oblate strain ellipsoid. We also interpret the observed pyroxene LPO patterns as reflecting this transpressional deformation. Given the relatively poor development of shear sense indicators, we cannot unambiguously rule out the possibility of a flattening-
dominated non-coaxial shear, which would also produce an oblate strain ellipsoid (lengthening-widening shear of Tikoff and Fossen 1993).

We emphasize that the enclave nature of the Sklené outcrop does not allow us to explicitly link these fabrics to spatial orientations with respect to the development of the Variscan orogen. Consequently, the “transpressional” fabric does not necessarily require a transient component of displacement for the mountain belt, although the fabrics are consistent with transpressional deformation inferred for the orogen by Schulmann et al (2005).

7. Conclusions

The patent and cryptic features of the Sklené peridotite, as described above, reveal a prolonged evolution from its inception in the mantle to its incorporation in the Bory granulite, as summarized in the following sequence of events:

- early partial fusion, melt extraction, and depletion of incompatible elements at moderate pressures in the spinel stability field,
- stabilization of garnet under high-pressure conditions in a subsolidus regime at $\sim1230 \, ^{\circ}C$ and 50 kbar, slightly within the stability field of majoritic garnet,
- metasomatic addition of Rh, Th, U, Pb, and Sr by transient, primitive basaltic melts,
- dynamic recrystallization, possibly with a gradual transition to grain size sensitive creep, at $\sim915 \, ^{\circ}C$ and 24 kbar, accompanied by stabilization of pargasitic amphibole and development of [010] maxima and [100] and [001] girdles in olivine in a flattening-dominated, transpressive deformation regime, prior to incorporation in granulite, but likely reflecting the early stages of convergence between Brunia and Moldanubia,
- local, static recrystallization and growth of spinel within kelyphite at $\sim900 \, ^{\circ}C$ and 17–21 kbar, probably after incorporation in granulite and isofacial with granulite peak metamorphism,
- and finally, development of a talc–amphibole reaction rim between peridotite and granulite, due to the instability of olivine with quartz and plagioclase in the presence of a hydrous fluid phase.

The Sklené peridotite yields a Sm–Nd mineral isochron cooling age of $338 \pm 13 \, Ma$, which is comparable to that for other Variscan garnet peridotites, pyroxenites, and eclogites in the Gföhl Terrane. With respect to mineral paragenesis, PT conditions, major-element composition, and Nd and Sr isotopic compositions, the Sklené peridotite is most similar to Type III subcontinental lithosphere. However, the pattern of trace-element metasomatism resembles that in Type II–Mg peridotite in the nearby Horní Bory quarry. Thus, we suggest that the Sklené peridotite represents a fragment of Type III subcontinental lithosphere, which experienced only incipient effects of Type II metasomatism, in contrast to the more pervasive metasomatism recorded by Type II Mg and Fe peridotites in the Horní Bory quarry.

Garnet peridotites in the Gföhl Terrane include a diversity of mantle lithologies and histories, and each peridotite occurrence must be investigated in detail to reveal the full panorama of mantle evolution in the Variscan orogenic belt.

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Moldanubian garnet peridotite
Appendix – analytical methods

A.1. Electron probe microanalysis

Minerals were analyzed by wavelength-dispersion spectrometry (WDS) with a Cameca SX50 instrument at the University of Wisconsin. Operating conditions were: 15 kV accelerating voltage, 20 nA beam current (Faraday cup), and beam diameter of 1 µm. Combinations of natural minerals and synthetic materials were used as standards for each of the mineral species, e.g. natural olivine for Mg, Fe, and Si and Ni metal for Ni in unknown olivine, synthetic spinel for Mg and Al and natural chromite for Fe and Cr in unknown spinel, and comparably appropriate combinations for pyroxenes and amphibole. Data reduction was performed by Probe for Windows software, utilizing the $\phi(\rho z)$ matrix correction of Armstrong (1988).

A.2. Electron back-scattered diffraction

EBSD data were collected from oriented, polished thin sections that had been subjected to further chemical-mechanical polishing for ~20 minutes with colloidal silica and very thinly carbon-coated to prevent specimen charging. Data were acquired on a Hitachi S3400N SEM at the University of Wisconsin–Madison and processed using HKL CHANNEL5 software. A general description of this method is provided by Prior et al. (1999). Crystallographic orientation data were collected using a step size of 15 µm, <1/3 of the smallest grain size of the matrix material. Patterns were collected relatively slowly, to maximise quality, and indexed on the basis of ≥ 6 Kikuchi bands so that only limited (<1%) amount of mis-indexing of crystal orientation could have occurred. Resultant data collection rates were 1 point every ~0.4 s. Data were processed (see Bestmann and Prior 2003) to produce microstructural maps, and orientation data sets based on one point per grain, where grains are separated by boundaries with misorientations ≥10°. Grains containing at least 3 analysis points (i.e. ≥ 40 µm in diameter) were selected for plotting on LPO figures. This sampling scheme produces data directly comparable with data collected by universal stage techniques (e.g. Webber et al. 2008). LPO fabric plots were generated using the program PFCCh5.app courtesy of D. Mainprice, and are presented on lower hemisphere equal area stereographic projections standard X–Y–Z fabric reference frame as described in the caption of Fig. 6. Fabric strengths were quantified using the M-index of Skemer et al. (2005), which is a measure of the deviation of the distribution of uncorrelated misorientations from the expected random distribution in a crystal aggregate. Possible M-indices range from 0 for a completely random distribution, to 1 for a single crystal. These calculations were performed using the matlab script Mindex_GUI.m (courtesy of P. Skemer).

A.3. Wet chemical analysis

Major and minor elements in whole-rock samples were determined by traditional wet-chemistry methods at the Faculty of Science, Charles University. Replicate analyses of reference material, e.g., PCC–1, yield an average precision of ±5% (1σ).

A.4. Trace element ICP-MS analyses

Whole-rock ICP-MS trace-element analyses were obtained using the methods of Strnad et al. (2005) at the Faculty of Science, Charles University. Clinopyroxene and garnet separates were prepared using a combination of heavy liquid and magnetic separations, which resulted in > 95 % pure separates. Separates were then leached in hot acids (~60 ºC) to remove possible grain-boundary contamination, using the following leaching steps: garnet – 6N HCl for 1 day, 7N HNO₃ for 1 day and 11N HF for 10 min; clinopyroxene – 6N HCl for 1 hour and 6N HNO₃ for 1 hour. Trace-elements in clinopyroxene and garnet were measured by ICP-MS at Charles University, using the same procedures as mentioned above (Strnad et al. 2005).
A.5. Rb-Sr and Sm-Nd isotopes

For isotope analyses, whole-rocks, pyroxenes and garnets were spiked with a mixed $^{84}\text{Sr}/^{87}\text{Rb}$ and $^{149}\text{Sm}/^{150}\text{Nd}$ tracer solution. All samples were digested in 52% HF for seven days at 180 °C in a Teflon bomb surrounded by a steel jacket. Before digestion, garnets were leached in 6 M HCl (1 day), 7 M HNO$_3$ (1 day) and 11 M HF (10 minutes) at 60 °C whereas pyroxenes were leached in 6 M HCl (1 hour) and 6 M HNO$_3$ (1 hour) at the same temperature. The whole-rock powders were not leached prior to digestion. Digested samples were dried and redissolved in 6N HCL, dried again and redissolved in 2.5N HCL. Rb, Sr and light rare-earth element separation was accomplished on quartz columns by conventional ion exchange chromatography with a 5 ml resin bed of Bio Rad AG 50W–X12, 200–400 mesh. Nd was separated from Sm and other rare-earth elements on quartz columns using 1.7 ml Teflon powder coated with HDEHP, di(2-ethylhexyl) orthophosphoric acid, as cation exchange medium. All isotopic measurements were made by Thermal Ionization Mass Spectrometry, on a Finnigan MAT 262 mass spectrometer. Sr was loaded with a Ta–HF activator on pre-conditioned W filaments and was measured in single-filament mode. Rb was loaded with ultra-pure H$_2$O on pre-conditioned Re-filaments and measurements were performed in a Re double filament configuration. Sm and Nd were loaded as phosphates and measured in a double Re-filament configuration mode. The $^{87}\text{Sr}/^{86}\text{Sr}$ ratios were normalized to $^{86}\text{Sr}/^{88}\text{Sr} = 0.1194$, the $^{143}\text{Nd}/^{144}\text{Nd}$ ratios to $^{140}\text{Nd}/^{144}\text{Nd} = 0.7219$, and Sm isotopic ratios to $^{147}\text{Sm}/^{152}\text{Sm} = 0.56081$. Analyses of two different loads of La Jolla standard gave $^{143}\text{Nd}/^{144}\text{Nd}$ ratios of 0.511842 ± 0.000008 and 0.511833 ± 0.000008 (errors are 2$\sigma$) while two analyses of the NBS 987 Sr standard yielded a $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of 0.710259 ± 0.000010 and 0.710246 ± 0.000009 (2$\sigma$). Total procedural blanks (chemistry and loading) were 125 pg for Sr, 35 pg for Nd and 4 pg for Sm. Single stage model-ages (T$_{DM}$) were calculated with depleted present-day parameters $^{143}\text{Nd}/^{144}\text{Nd} = 0.513151$ and $^{147}\text{Sm}/^{144}\text{Nd} = 0.219$ (Liew and Hofmann 1988). $^{143}\text{Nd}/^{144}\text{Nd}$ ratios are quoted in the εNd notation of DePaolo and Wasserburg (1976) as deviations from a chondritic reference (CHUR) with present-day $^{143}\text{Nd}/^{144}\text{Nd} = 0.512638$ (Jacobsen and Wasserburg 1980). Least-square regression of the Sm–Nd isotopic data with assessment of fit using the mean square of the weighted deviates (MSWD), were calculated after Ludwig (2003).

A.6. Analyses of highly siderophile elements (HSE)

HSE concentrations and osmium isotopic compositions were obtained at the University of Maryland. Whole-rock powder (~ 2 g) was mixed with Re–Os and Ir–Ru–Pt–Pd spikes and dissolved in Carius tubes using reverse concentrated aqua-regia at 280 °C for 2–3 days (Shirey and Walker 1995). After digestion, osmium was separated using solvent extraction by CCl$_4$ and back reduction to HBr (Cohen and Waters 1996), and the final fraction was purified by microdistillation (Birck et al. 1997). The remaining HSE were separated by anion exchange chromatography.

Os concentration and isotopic composition were analyzed by Negative Thermal Ionization Mass Spectrometry (N–TIMS) on a VG Sector 54 instrument, using an electron multiplier in a peak jumping mode. The measured Os isotopic ratios were corrected for mass fractionation using $^{192}\text{Os}/^{190}\text{Os} = 3.08271$ (Shirey and Walker 1998). External precision was monitored using an in-house UMCP standard solution and was better than ± 0.3% (2σ). Iridium, Ru, Pt, Pd and Re were analyzed using a Nu Plasma multi-collector ICP-MS with two–three electron multipliers in a static mode. The isotopic fractionation of corresponding elements was corrected using a linear law and in-house Ir, Ru, Pd, Pt and Re standard UMCP solutions (0.2–0.3 ppb) that were run with samples. HSE measurements are estimated to be accurate and precise to ± 5%.