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Title: Clumped fluoride-hydroxyl defects in forsterite:

Implications for the upper-mantle

Authors: Céline Crépisson¹, Marc Blanchard¹*, Hélène Bureau¹, Chrystèle Sanloup²,³, Antony C. Withers⁴, Hicham Khodja⁵, Suzy Surblé⁵, C. Raepsaet⁵, Keevin Béneut¹, Clémence Leroy⁴, Paola Giura¹, Etienne Balan¹

¹ Institut de Minéralogie et Physique des Milieux Condensés (IMPMC) UMR CNRS 7590, UMR IRD 206, Université Paris VI, case 115, 4 place Jussieu, 75252 Paris cedex 05, France
² SUPA, Centre for Science at Extreme Conditions and School of Physics and Astronomy, University of Edinburgh, Edinburgh EH9 3JZ, UK
³ Institut des Sciences de la Terre de Paris (ISTEP) UMR CNRS 7193, Université Paris VI, 4 place Jussieu, 75252 Paris, cedex 05 France
⁴ Department of Earth Sciences, Western University, London, ON, Canada
⁵ CEA/DSM/IRAMIS/SIS2M, UMR3299, CEA Saclay, 91191 Gif sur Yvette, France

*Corresponding author. E-mail address: marc.blanchard@impmc.upmc.fr ; Tel.: +33 1 44279822 ; fax: +33 1 44273785

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Abstract

The mechanism and magnitude of fluorine incorporation in H-bearing forsterite were investigated through a combined experimental and theoretical approach. Forsterite samples were synthesized in a piston cylinder press at 2 and 4 GPa, in hydrous conditions, with or without fluorine. High fluorine solubilities of 1715 and 1308 ppm F were measured by particle induced gamma-ray emission (PIGE) in forsterites synthesized at 2 and 4 GPa, respectively. In addition, first-principles calculations based on density functional theory were performed in order to investigate the coupled incorporation mechanisms of fluorine and water in forsterite. Our results demonstrate the close association of fluoride, hydroxyl groups and Si vacancies. Comparison of experimental and theoretical infrared absorption spectra enables assignment of the nine OH stretching bands (3500-3700 cm$^{-1}$) observed in F-rich synthetic forsterite to clumped fluoride-hydroxyl defects in the forsterite crystal structure. Noteworthily, similar bands were previously recorded on some natural olivine with Mg/(Mg+Fe) molar ratio down to 0.86. Fluorine and water cycles are therefore strongly coupled through the nominally anhydrous minerals and the mantle fluorine budget can be entirely accommodated by these mineral phases.
1. Introduction

Very little is known about the deep fluorine cycle. The presence of fluorine in volcanic gases and melt inclusions shows that there is a fluorine flux from the depth to the surface (Schilling et al., 1980; Aoki et al., 1981; Symonds et al., 1988; Bureau et al., 1998; Pyle and Mathers, 2009). Fluxes from surface to depth are also expected at subduction zones, where serpentinites are the most likely carrier of halogen elements including fluorine (John et al., 2011 and references therein). Fluorine released from the dehydration of the oceanic crust can be degassed during through arc volcanism or may be recycled back into the mantle with an efficiency of around 95% (Straub and Layne, 2003; John et al., 2011). Significant fluorine enrichments in mantle rocks are suggested by elevated concentrations (0.2 wt%) in some kimberlite samples (Paul et al., 1975) and by the identification of "fluoride melts" in a metasomatized mantle xenolith from New Zealand (Klemme, 2004). Estimates of the fluorine concentration of the primitive Earth mantle based on CI carbonaceous chondrite range from 15 ppm to 25 ppm (McDonough and Sun, 1995; Palme and Jones, 2003).

Little is also known about fluorine incorporation in the solid Earth. For long it has been proposed that the main fluorine carriers were hydrous and minor accessory minerals such as apatite, micas, amphibole (Smith et al., 1981; Smith, 1981) or clinohumite (Brey et al., 2009). The role of nominally anhydrous and fluorine-free silicates as a deep fluorine reservoir has only recently been proposed (Hervig and Bell, 2005; Bromiley and Kohn, 2007; Beyer et al., 2012; Dalou et al., 2012; Bernini et al., 2013; Mosenfelder and Rossman, 2013a, 2013b; Fabbrizio et al., 2013). In particular, fluorine solubility in olivine, the main constituent of the upper mantle, could reach 900 ppm (Fabbrizio et al., 2013), 1900 ppm (Bernini et al., 2013) or even 4500 ppm (Bromiley and Kohn, 2007). The incorporation mechanism of fluorine in these nominally anhydrous and fluorine-free silicates remains speculative. Crystal-
chemical considerations based on the similar ionic radius of fluoride (0.130 nm) and hydroxyl ions (0.136 nm) suggest that fluoride incorporation could share some similarities with the incorporation of OH groups in these minerals. As a matter of fact, the fluoride for hydroxyl substitution is commonly observed in hydrous minerals (e.g. amphibole, Robert et al., 1999). The substitution has also been reported in some nominally anhydrous silicates, such as garnet (Visser, 1993) and zircon (Caruba et al., 1985; Balan et al., 2013), and in high pressure hydrous phases, such as superhydrous phase B (Hazen et al., 1997).

In the present study, we determine the mechanism and magnitude of fluorine incorporation in H-bearing forsterite (Fo100) by experiment and theory. This work is completed by the further investigation (Particle Induced Gamma-ray Emission measurements PIGE) of the synthetic samples from Withers et al. (2011, 2012), which have a composition similar to mantle olivine (Fo90). The results demonstrate the close association of fluoride, hydroxyl groups and Si vacancies. Implications for fluorine storage in the upper mantle are discussed.

2. Materials and Methods

2.1. Syntheses and preparation

Forsterite samples were synthesized in a piston cylinder press at the University of Edinburgh, CSEC. The starting material was prepared by homogenizing a mixture of high purity (reagent grade) reactants (MgO, SiO₂, Mg(OH)₂, and NaF). The Mg/Si molar ratio of the starting material was 1.75, initial H₂O and F contents were ~5 and ~1.1 wt%, respectively. Experiments were performed at 2 GPa (run #PC36F) and 4 GPa (run #PC38 and #PC38F), at ~ 1250°C, in a 3 cm long cell-assembly composed of a graphite heater surrounded by pyrex and talc sleeves. Two welded 5 mm height and 3 mm outer diameter Au₇₅Pd₂₅ capsules, with and without NaF, were simultaneously inserted in the press on top of each other and separated
by an alumina disk. MgO powder dried overnight at 1000°C was packed around the Au$_{75}$Pd$_{25}$ capsules. Experiments were run for 3 days and terminated by turning-off the heater before a slow decompression. Temperature was monitored with a Type C thermocouple (W$_{74}$Re$_{26}$-W$_{95}$Re$_{5}$) whose extremity was in contact with the top capsule.

2.2. Electron microscopy and microprobe analysis

Forsterite grains were mounted on a metallic disk with carbon tape and coated with a 10 nm thick carbon film. The texture of the samples was examined with a Zeiss Ultra 55 field emission scanning electron microscope (SEM) with a working distance of 3±0.2 mm and an acceleration voltage of 15 keV. Major elements were analyzed using a Cameca SX FIVE electron microprobe at CAMPARIS facility (UPMC). Acceleration voltage was set to 15 keV, current to 10 nA and beam was focused to 10 µm diameter at the surface of the sample. Counting times were 10 s both on the sample and the background.

2.3. Vibrational spectroscopy

IR measurements were done on a Bruker IFS 66v/S Fourier transform infrared spectrometer working in vacuum and aligned in transmission geometry. The sample was disposed in a homemade sample chamber at the focal point of two cassegrainian reflectors. The analysis area of the sample was selected with slits located between the sample and the detector. Every spectrum was acquired in the frequency region 550-9000 cm$^{-1}$ with resolution of 4 cm$^{-1}$ and accumulation of 64 scans using the mid infrared instrumental configuration i.e. Globar, KBr and MCT as respectively source, beam splitter and detector. The background was measured after each sample measurement without changing analysis conditions. Absorbance was obtained after subtraction of the baseline using the OPUS/IR software. Seven to ten grains (50 to 150 µm thick) from each capsule were analyzed. After normalization to
thickness and subtraction of epoxy signal and baseline, average unpolarized absorption spectra were calculated and fit using Lorentzian line-shapes (Table 1). In the wavenumber range of interest, the epoxy signal corresponds to a strong band at ~3440 cm\(^{-1}\), where there is no olivine OH band, plus a weak and broad band centred at ~3550 cm\(^{-1}\). The epoxy correction was made for two of the seven spectra of the PC38 synthesis and three of the ten spectra of the PC36F synthesis.

Raman spectroscopic measurements were performed with a Renishaw InVia Raman spectrometer coupled with an optical microscope, using a 514.5 nm laser excitation radiation (1 mW power) and a 1200 mm\(^{-1}\) grating. Spectra were acquired from 100 to 2200 cm\(^{-1}\), and averaged from 4 accumulations, each with a counting time of 60 s.

2.4. Ion beam analysis

Particle induced gamma ray emission (PIGE), Rutherford backscattering spectrometry (RBS) and Elastic Recoil Detection Analysis (ERDA) were performed at the nuclear microprobe of CEA Saclay SIS2M / LEEL (Khodja et al., 2001). In both cases particle induced X-ray emission (PIXE) and Rutherford Backscattering RBS measurements were simultaneously associated to PIGE and ERDA. PIGE is based on the detection of γ-ray emission during a nuclear reaction triggered by a high-energy proton beam and makes it possible to determine accurately fluorine concentrations. The γ-ray emission at 109.9 keV of the \(^{19}\text{F}(p,p'\gamma)\) nuclear reaction induced by a 3 MeV proton beam was detected with an HP-Ge detector (Mosbah et al., 1991). ERDA is based on the detection of protons at a low ‘grazing’ angle of 15° from a \(^4\text{He}\) incident beam of 3 MeV after they have been ejected from the samples through elastic collisions, this method has been recently used to quantify water in nominally anhydrous minerals; details about the procedure are described in (Raepsaet et al., 2008; Bureau et al., 2009, Withers et al., 2012). PIXE and RBS measurements allow chemical
characterization of the sample with respect to major and trace elements. RBS measurements were performed using an annular detector positioned at 170° with respect to the incident beam direction, in order to monitor the electrostatic charge delivered to the sample. Further details can be found in Bureau et al. (2009) and Habrioux et al. (2012). All analyses were performed by scanning a 4 µm × 4 µm microbeam on 100 µm × 100 µm areas.

In PIGE analysis, the fluorine concentration is determined by comparison with a reference sample using the following relation:

$$\frac{[X_{\text{sample}}]}{[X_{\text{std}}]} = \frac{S_{\text{sample}}}{S_{\text{std}}} \times \frac{A_{\text{sample}}}{A_{\text{std}}} \times \frac{N_{\omega_{\text{std}}}}{N_{\omega_{\text{sample}}}}$$  (1)

where $A$ is the peak area associated with $\gamma$-ray transitions observed for element $X$, $S$ the stopping power of the sample and $N_{\omega}$ the number of protons by solid angle unit delivered to the selected area. The reference sample was a pantellerite glass with F content of 4200 ppm and a similar stopping power to the forsterite samples (KE12; Métrich and Rutherford, 1991). Calibration was verified on a Macusanite glass with known F content of 1.33 wt% determined by EMPA (Pichavant et al., 1987). Data were processed with the RISMIN software (Daudin et al., 2003) following the procedure described in Bureau et al. (2009). Statistical errors are around 10%. A fit of the RBS spectra with the SIMNRA software (Mayer, 1997), using the olivine matrix composition, enables determination the $N_{\omega}$ parameters.

2.5. Theoretical methods

Calculations were performed using the theoretical approach and convergence parameters described in Balan et al. (2011). Calculations are based on the density functional theory, with the generalized gradient approximation (GGA) to the exchange-correlation functional as proposed by Perdew, Burke and Ernzerhof (Perdew et al., 1996) and periodic boundary conditions. Structure relaxations were done on a 2×1×2 supercell (112 atoms) of
forsterite using the PWscf code of the Quantum Espresso package (Giannozzi et al., 2009; http://www.quantum-espresso.org). The theoretical relaxed primitive orthorhombic-cell parameters of anhydrous forsterite are $a = 4.78$ Å, $b = 10.28$ Å, and $c = 6.01$ Å ($Pbnm$ space group). The optimized cell parameters of pure forsterite were kept constant during the relaxation of the F- and OH-bearing supercells. During this step, no symmetry constraint was applied to the atomic positions and the forces on atoms were minimized to less than $10^{-4}$ Ry/a.u. Following the same procedure as Balan et al. (2011) high-frequency OH stretching modes and corresponding IR absorption spectra were calculated from the dynamical matrix, Born effective charge tensors and dielectric tensor, obtained using the density functional perturbation theory (Baroni et al., 2001) as implemented in the Phonon code of the Quantum Espresso package (Giannozzi et al., 2009; http://www.quantum-espresso.org).

3. Results

3.1. Chemical composition and infrared spectra of synthetic forsterite samples

The investigated samples consist of transparent and large (> 100 µm) forsterite crystals with no observed inclusions (Fig. 1). Coexisting enstatite crystals are also identified at the top of the capsule by Raman measurements. Major element analyses indicate that the forsterite grains are close to the stoichiometric composition (Table 1). Beside OH defects, the major measured impurity is fluorine. Concentrations of 1715 ppm F and 1308 ppm F are measured by PIGE at 2 GPa (PC36F) and 4 GPa (PC38F) respectively (Table 1). Minor Na concentrations are also detected by PIGE, i.e. 131 ppm Na and 81 ppm Na at 2 GPa (PC36F) and 4 GPa (PC38F) respectively. These Na concentrations do not exceed the typical range of few hundreds of ppm expected for Na in olivine (e.g. Borisov et al., 2008).

The OH-stretching infrared absorption spectrum of the F-free forsterite sample (#PC38, Fig. 2) is similar to that reported in previous studies (Lemaire et al., 2004; Bali et al.,
It is dominated by the "hydrogarnet-type" defect \((4H)_{Si}\) responsible for a cluster of narrow absorption bands in the 3620-3550 cm\(^{-1}\) region (Balan et al., 2011; Umemoto et al., 2011). Two broader bands observed in the same region have been recently attributed to interstitial OH-groups (Ingrin et al., 2013; Balan et al., 2013). Weak and broad bands ascribed to protonated Mg vacancies (e.g. Balan et al., 2011) are observed at 3160 and 3220 cm\(^{-1}\); whereas weak and narrow bands observed between 3300 and 3400 cm\(^{-1}\) could be related to trace amounts of trivalent impurities (Berry et al., 2007).

The OH-stretching spectra of F-bearing forsterite samples (#PC36F and #PC38F) are similar to each other. They are also similar to the unpolarized spectrum of a F-rich forsterite (900 ppm F) synthesized at 1200°C and 2 GPa by Fabbrizio et al. (2013). Although still dominated by a group of intense bands observed in the 3700-3500 cm\(^{-1}\) region, they differ significantly from that of the F-free sample (#PC38). These bands are not observed in F-free forsterite and are rarely seen in natural samples. The most intense band is located at 3570 cm\(^{-1}\) with a shoulder at 3555 cm\(^{-1}\). Seven other intense bands occur at 3674, 3641, 3624, 3597, 3591, 3535 and 3514 cm\(^{-1}\) (Table 1). The 3612 cm\(^{-1}\) band related to \((4H)_{Si}\) is still visible but very weak (Fig. 2). Small variations in the relative intensities of the F-related bands in #PC36F and #PC38F cannot be considered as significant owing to the limited number of measurements used to build average IR spectra (Fig. 3). Some bands observed here are close in frequency to OH bands reported previously in natural olivine samples and assigned to hydrous nano-inclusions, such as talc, serpentine or Ti-clinohumite (Beran and Libowitzky, 2006). Among the defects related to Ti-clinohumite, Ti planar defects give rise to a band at 3410 cm\(^{-1}\) and the Ti point defect leads to two bands at 3572 and 3525 cm\(^{-1}\) (Berry et al., 2005; Balan et al., 2011). In the present spectra (Fig. 2), no bands are observed at 3525 and 3410 cm\(^{-1}\), which would suggest the absence of Ti related defects. Furthermore, no significant presence of Ti has been detected by PIXE during the ERDA analysis of the samples. Talc
corresponds to an OH band at 3677 cm\(^{-1}\) and bands in the range 3640-3700 cm\(^{-1}\) can be referred to serpentine (Beran and Libowitzky, 2006). The two highest bands, observed here at 3674 and 3641 cm\(^{-1}\), could then be assigned to talc or serpentine nano-inclusions that would be nearly homogeneously distributed in all crystals sampled from the two experiments but as it will be shown further by the calculations, intrinsic hydroxylated point defects in the crystal structure of F-bearing forsterite can already explain these bands. The significant modifications observed between the F-free and the F-bearing forsterite samples thus suggest that fluoride ions are associated with these OH-bearing defects.

Interestingly, the infrared spectrum of the F-rich synthetic forsterite samples is nearly identical to that of a natural F-rich forsterite sample from Pamir, Tadzikistan (Libowitzky and Beran, 1995), here referred to as the Pamir olivine (Fig. 4a). All the bands observed in the synthetic samples are present in the Pamir sample, except a shoulder at 3555 cm\(^{-1}\) and the small band at 3612 cm\(^{-1}\) related to \((4H)^{\text{Si}}\) defects. Noteworthily, other natural olivine samples described in the literature display absorption bands at wavenumbers close to those reported here for the F-bearing synthetic samples (Sykes et al., 1994; Kishina et al., 2001; Bell et al., 2004, Matsyuk and Langer, 2004; Koch-Müller et al., 2006, Mosenfelder et al., 2011). For these natural samples, Mosenfelder et al. (2011) specifically attributed the three bands above 3600 cm\(^{-1}\) to coupled substitutions of hydroxyl and fluoride.

Water contents are 448±90 ppm H\(_2\)O in F-free forsterite; 267±53 ppm H\(_2\)O and 887±124 ppm H\(_2\)O in F-rich forsterite samples synthesized at 4 and 2 GPa, respectively (Table 1). As a comparison, Lemaire et al. (2004) measured water contents ranging from 42 to 750 ppm in synthetic forsterite synthesized at 2 GPa and Withers et al. (2012) measured a water content of 462 ppm in olivine (Mg# = 90) synthesized at 3 GPa.

Water and fluorine contents are higher for forsterite synthesized at 2 GPa (#PC36F) than at 4 GPa (#PC38F), however a tendency can hardly be defined with only two pressures
investigated. Further experiments are needed to define the effect of pressure on simultaneous water and fluorine incorporation in olivine.

3.2. Theoretical results

Infrared spectroscopic measurements suggest that fluoride ions are associated with OH defects in the structure of F-bearing forsterite. Starting from the models of \((4H)^x_{\text{Si}}\) and \((2H)^x_{\text{Mg}}\) defects previously determined by Balan et al. (2011), the structure of fluorinated defects was obtained by substituting fluoride ions for OH groups. The most stable configurations of clumped fluoride and OH defects associated with Si vacancies are displayed in Fig. 4.a. For a single F-for-OH substitution, three configurations of almost equal stability (within 2 kJ.mol\(^{-1}\)) are obtained, denoted as \(F_{(O1)}\), \(F_{(O2)}\) and \(F_{(O3)}\) for F in O1, O2 and O3 site, respectively. Other orientations of OH groups lead to significantly less stable configurations (by ~30 kJ.mol\(^{-1}\)). When two F for OH substitutions are considered, two configurations are found equally stable (within 1.1 kJ.mol\(^{-1}\)): \(2F_{(O2, O1)}\) and \(2F_{(O2, O3)}\) corresponding respectively to OH/F substitutions in O2 and O1 sites and to OH/F substitutions in O2 and O3 sites. For three F for OH substitutions, the most stable configuration is obtained for F ions located in O2 and in the two symmetrical O3 sites (\(3F_{(O3, O3, O2)}\)) with the OH group pointing toward the center of the vacancy.

For a Mg vacancy, the most stable configuration corresponds to a F substituting for the OH at the O2 site. The position of the remaining OH group in O2 site is almost unchanged. Its OH bond length is slightly shortened (1.000 Å versus 1.003 Å in the fully protonated Mg vacancy).

Assuming that both Mg and Si vacancies are present in the crystal, the preferential fluorination of Mg or Si vacant sites can be described by three reactions, depending on the number of fluoride ions considered:
The total energy difference associated with these three reactions is always significantly negative (-101 kJ.mol$^{-1}$, -98 kJ.mol$^{-1}$ and -81 kJ.mol$^{-1}$ for reactions 2, 3 and 4, respectively).

This suggests that OH/F substitutions are more likely to occur in Si protonated vacancies than in Mg protonated vacancies. Similarly, the relative stability of mixed OH/F defects associated to Si vacancies can be compared to that of equivalent proportions of pure F and OH defects:

\[
\frac{3}{4}[V_{Si}4(OH)_{o}]^{r} + \frac{1}{4}[V_{Si}4F_{o}]^{r} \leftrightarrow [V_{Si}3(OH)_{o}F_{o}]^{r} \tag{5}
\]

\[
\frac{1}{2}[V_{Si}4(OH)_{o}]^{r} + \frac{1}{2}[V_{Si}4F_{o}]^{r} \leftrightarrow [V_{Si}2(OH)_{o}2F_{o}]^{r} \tag{6}
\]

\[
\frac{1}{4}[V_{Si}4(OH)_{o}]^{r} + \frac{3}{4}[V_{Si}4F_{o}]^{r} \leftrightarrow [V_{Si}3(OH)_{o}3F_{o}]^{r} \tag{7}
\]

The total energy difference associated with these reactions is always significantly negative (-19 kJ.mol$^{-1}$, -34 kJ.mol$^{-1}$ and -32 kJ.mol$^{-1}$ for reactions 5, 6 and 7 respectively). Although the absolute abundance of mixed OH/F defects also depends on the fluorine activity and H$_2$O fugacity, the greater stability of mixed OH/F defects with respect to the defect end-members is consistent with the weakness of the signal ascribed to fully protonated Si vacancies.

The theoretical IR absorption spectrum corresponding to the OH-stretching modes of the seven most stable geometries of the OH- and F-bearing defects associated with Mg and Si vacancies have been computed.

In the case of an Mg vacancy, one OH/F substitution induces an upshift of the band from 3222 to 3291 cm$^{-1}$, compared to the fully protonated Mg vacancy, while the polarization
along the z axis remains unchanged (Balan et al., 2011). This frequency shift is directly related to the geometrical changes related to the presence of the fluoride ion.

The IR spectra generated for the six most stable configurations of OH/F defects in Si site are displayed in Fig. 4. For one OH/F substitution, each of the three configurations \( \text{F(O1)}, \text{F(O2)}, \text{and F(O3)} \) lead to three distinct bands (Fig. 4b). The most intense is strongly polarized along x (respectively at 3611, 3583 and 3556 cm\(^{-1}\)) whereas the two other bands at lower frequency are polarized along y and z (respectively around 3577, 3534 and 3505 cm\(^{-1}\), with a splitting smaller than 10 cm\(^{-1}\)). Most of the bands result from a coupling between 2 or 3 OH-groups of the defects. OH-bands related to two OH/F substitutions fall very close to contributions from the 1 OH/F substitution (Fig. 3c). Configuration \( 2\text{F(O2, O1)} \) leads to one band strongly polarized along x (3581 cm\(^{-1}\)) and one band polarized along z (3554 cm\(^{-1}\)) whereas configuration \( 2\text{F(O2, O3)} \) leads to one band strongly polarized along x (3610 cm\(^{-1}\)) and one band polarized equally along x and z (3546 cm\(^{-1}\)). All absorption bands arise from coupled vibrational motions of the two OH groups. Finally, \( 3\text{F(O3, O3, O2)} \) is associated with one band at higher frequency (3674 cm\(^{-1}\)) strongly polarized along x (Fig. 4d).

Lack of polarized measurement prevents a thorough comparison of the theoretical IR spectra with the spectrum of the present synthetic F-rich forsterite. However, the 3700-3500 cm\(^{-1}\) region of the average IR spectrum of the synthetic samples is very similar to the spectrum of the Pamir olivine (Fig. 5a). We thus consider that the theoretical IR spectra can be compared with the polarized spectrum of the Pamir olivine reported on Fig. 5 (Libowitzky and Beran, 1995). We point out that different polarizations for what seems to be single experimental bands on the Pamir olivine spectrum are often explained by theoretically calculated contributions from different defects.

The band at 3674 cm\(^{-1}\) (band a, Fig. 5) is well explained by the configuration \( 3\text{F(O3, O3, O2)} \) which shows a band strongly polarized along x at the same wavenumber. Although the two
theoretical frequencies almost coincide at 3610-3611 cm\(^{-1}\), the bands at 3641 and 3624 cm\(^{-1}\) (bands b and b') are best explained by the 2F\(_{(O2, O3)}\) and F\(_{(O1)}\) defects, with the main polarization along x and a much smaller polarization along z. The weakly split bands around 3597 and 3591 cm\(^{-1}\) (bands c and c') can be attributed to contributions from the configurations 2F\(_{(O1, O2)}\) and F\(_{(O2)}\) for the x component and from configuration F\(_{(O1)}\) for the weaker z and y components. The intense band at 3570 cm\(^{-1}\) (band d) is well reproduced by configuration F\(_{(O3)}\) giving a band at 3556 cm\(^{-1}\) with a polarization along x, whereas the contribution along z could arise from the 2F\(_{(O1, O2)}\) defect. Finally, the band e at 3535 cm\(^{-1}\) could be related to the configuration F\(_{(O3)}\). However, the polarization properties, relative intensity and band positions of the experimental band e are not well reproduced and this assignment is still very speculative. Furthermore, there are differences between the spectrum of the synthetic forsterite and that of the Pamir olivine with the presence of a band at 3515 cm\(^{-1}\) (Fig. 5): the band e could actually turn into a doublet reproduced by the configuration F\(_{(O3)}\) though polarization remains unexplained.

3.3. *Integrated molar absorption coefficients of OH defects in forsterite*

For each configuration we calculate the integrated molar absorption coefficients for F- and OH-bearing defects. Results are given in Figs. 4b, 4c and 4d for F-bearing Si vacancies and we find a value of 135668 cm\(^{-2}\) per mol H\(_2\)O/L for an Mg defect with one OH/F substitution. These new data are aligned along a similar trend to OH defects in F-free forsterite (Fig. 6), i.e. on a line rather parallel to the general trend from Balan et al. (2008) but with a downward offset of ~50000 cm\(^{-2}\) per mol H\(_2\)O/L. This shift is consistent with previous findings on F-free forsterite by Balan et al. (2011). Therefore, the molar absorption coefficient derived from F-free olivine by Withers et al. (2012) can also be applied to F-rich olivine. Noteworthily, the 3F\(_{(O3, O3, O2)}\) defect is not on the same trend as the other defects. Despite a
higher frequency, its molar absorption coefficient is close to values obtained for the other
defects. This suggests that all Si defects in forsterite can be approximately treated with a
single molar absorption coefficient.

4. Discussion

4.1. Clumped OH/F defects in natural olivine samples

The synthetic forsterite samples display a characteristic IR spectrum in the OH
stretching-region that we confidently assign to the presence of clumped fluoride-hydroxyl
defects. The reported spectra are similar to those of Fabbrizio et al. (2013) and can be
observed also in various natural F-rich olivines from crustal environments (Sykes et al., 1994;
Libowitzky and Beran, 1995) and in a kimberlite megacryst from Monastery mine with up to
47 ppm F (Bell et al., 2004; Mosenfelder et al., 2011, 2013a) and in some olivines from the
Udachnaya kimberlite outcrop (Kishina et al., 2001; Matsyuk and Langer, 2004; Koch-Müller
et al., 2006). The latter have not yet been analyzed for fluorine, but the presence of melt
inclusions with F-rich phases suggests that some F may have been incorporated in the host
olivines (Golovin et al., 2007). This indicates that the results of our experiments as well as
those from Fabbrizio et al. (2013) are directly applicable to natural conditions. In particular,
the Mg/(Mg+Fe) molar ratio, which varies from 86.00 to 99.65 in the natural samples
discussed above (Sykes et al., 1994; Libowitzky and Beran, 1995) does not modify the IR OH
stretching bands. This is also true for the kimberlitic olivines investigated by Bell et al. (2004)
and Mosenfelder et al. (2011). The positions of their OH stretching bands are similar, within 5
cm⁻¹, to those reported in the present study for Fe-free samples. Consequently the
incorporation mechanisms and solubility of fluorine are not expected to be significantly
changed in presence of iron. The present results also indicate that some OH IR bands
observed by Sykes et al. (1994) in a boron and fluorine rich olivine (0.33-0.55 wt% F and
0.35-0.40 wt% B) could be explained without having to consider F-B coupled substitutions. Additionally FTIR measurements prove to be useful for the identification of F-rich olivine including Fe-rich olivine. It must be noted however that this IR signature is not observed for synthetic hydrous olivines from Withers et al. (2011) and (2012) that contain up to 47 ppm F, despite not having been intentionally doped with fluorine (Table 1). On the other hand, the natural olivines ROM177 and GRR1012-2 containing 37 and 47 ppm F, respectively (Mosenfelder et al., 2013a), display the IR signature assigned here to clumped fluoride-hydroxyl defects. This apparent discrepancy might be explained by the ratio of the fluorine to water contents. With less than 220 ppm H$_2$O, the fluorine-related defects are still distinguishable by FTIR measurements, whereas in the synthetic olivines containing more water, the F-free hydroxyl defects are dominant.

Similar studies on other NAMs may reveal comparable processes, even though first studies on F-bearing pyroxenes (Mosenfelder and Rossman, 2013a and 2013b) have not revealed so far characteristic IR OH stretching bands that could be linked to incorporation of fluorine.

4.2. Cooperative solubility of water and fluorine in olivine

In nominally anhydrous silicates, the hydrolytic weakening mechanism consists in an increase of cationic vacancy concentration in presence of water due to a thermodynamic stabilization of the vacancies by the presence of the charge compensating protons (Brodholt and Refson, 2000). Our experimental and theoretical results indicate that mixed OH/F defects display an even greater stability. Therefore, the presence of fluoride ions should contribute to the hydrolytic weakening mechanism. An important consequence of this cooperative effect is that the dependence of water solubility in forsterite on water fugacity should be modified in presence of fluoride. Indeed, this dependence is a function of the number of OH groups
associated with a specific OH defect (Keppler and Bolfan-Casanova, 2006). In high pressure F-free forsterite samples, water incorporation is dominated by (4H)$_{Si}^x$ defects and to a lesser proportion by interstitial defects (Ingrin et al., 2013). Thus, the replacement of (4H)$_{Si}^x$ defects by clumped F/OH defects should lead to a decrease in the fugacity exponent between the F-free and F-bearing system.

4.3. Geodynamical implications

We show that fluorine solubility can reach more than 1300 ppm in hydrous olivine at pressures and temperatures relevant of the upper-mantle. Such high solubility implies that significant amounts of fluorine can be stored in nominally anhydrous minerals in the upper mantle. Our experiments establish the link between F and OH assimilation in mineral networks. This must be particularly true in subduction settings where water is recycled back to the mantle. Indeed, if F and OH are associated, F could be recycled in the mantle together with water through F incorporation in NAMs in a peridotitic layer at the bottom of the down-going oceanic lithosphere. In that case fluorine could be transferred in deeper regions of subduction zones "en route" to the transition zone and lower mantle during deep hydrous mineral transformations in subducted slabs such as in the phase B (Hazen et al., 1997). Furthermore, recent results have demonstrated the capability of the transition zone to contain fluorine through an experimental study of the solubility of fluorine in ringwoodite and wadsleyite (Roberge et al., 2013).

To conclude, water and fluorine geodynamical cycles are likely to be strongly connected in the mantle owing to the existence of clumped defects in nominally anhydrous minerals.

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Synthesis conditions, chemical compositions of forsterite samples and positions of the observed absorption bands with their corresponding area and width. Infrared band position, area and full width at half height (FWHH) from fit of the spectra by Lorentzian functions (see text). Detailed polarized IR measurements of samples A710 and M475 are to be found in Withers et al. (2011).
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1 determined by EPMA, 2 by PIGE and 3 by ERDA. For IR absorption bands, *indicates F-related bands and † possible bands related to OH-interstitials.
Figure 1:
Secondary electron image of forsterite grains (synthesis #PC36F)

Figure 2:
Unpolarized average infrared spectra of synthetic forsterite (black dots) and fit of spectra by a sum of Lorentzian functions in grey line (see parameters of the Lorentzian functions in Table 1). Lowermost spectrum absorbance is multiplied by a factor of 4 and spectra are offset vertically for clarity.

Figure 3:
Unpolarized infrared spectra of forsterite grains derived from the synthesis #PC38F.

Figure 4:
a) Structural models of the most stable configurations for OH/F substitutions in Si vacancy investigated by first-principles calculations. In 3F_{(O3, O3, O2)}, the OH group points toward the center of the vacancy. In 2F_{(O2, O1)} the two OH groups point toward the F in O1 site. In 2F_{(O2, O3)} the OH in O1 site points toward the F in O3 site while the OH in O3 site points toward O in O1 site. In F_{(O1)} all OH groups are pointing toward the F atom. In F_{(O2)} the OH in O1 site points toward the F while the two OH in O3 sites point toward the O1 atom. In F_{(O3)} the OH in O1 site points toward the F while the other two OH point toward O in O1 site. Theoretical polarized IR spectra for one (b), two (c) and three (d) OH/F substitutions. Red vertical lines correspond to the position of the OH bands in the fully protonated Si vacancy (Balan et al., 2011). The integrated molar absorption coefficient of the corresponding OH defects is indicated in the figure’s legend (in cm$^{-2}$/mol H$_2$O/L).
Figure 5:

a) Polarized IR spectra of Pamir olivine from Libowitzky and Beran (1995) compared to the unpolarized average IR spectrum of run #PC38F. a.b. indicates the absence of the broad shoulder observed in F-rich forsterite samples at ~3555 cm$^{-1}$.

b) Calculated polarized spectra considering one, two and three OH/F substitutions in the Si vacancy. Proportions of the various configurations were taken into account following Umemoto et al. (2011) though variation in energy is too weak to induce major changes of relative intensities.

Figure 6:

Theoretical integrated molar absorption coefficients for the different F- and OH-bearing defects. The solid line corresponds to the theoretical correlation obtained by Balan et al. (2010), dashed line is a guide for the eyes.
Fig. 1.
Fig. 2.
Fig. 3.

#PC38F
4 GPa / 1308 ppm F
Fig. 4.
Fig. 5. 

Theoretical spectra
Fig. 6.

- Si no F
- Si 1 substitution
- Si 2 substitutions
- Si 3 substitutions
- Mg no F
- Mg F

General trend (Balan et al., 2008)

Kint (cm$^2$ per mol H$_2$/L)

wavenumber (cm$^{-1}$)