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This is the author's manuscript

Original Citation:

Availability:
This version is available http://hdl.handle.net/2318/117842 since 2015-12-28T11:58:47Z

Published version:
DOI:10.1016/j.ibiod.2012.07.018

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(Article begins on next page)
Lichen deterioration of asbestos and asbestiform minerals of serpentine rocks in Western Alps

Sergio E. Favero-Longo, Francesco Turci, Bice Fubini, Daniele Castelli, Rosanna Piervittori

Abstract

Naturally Occurring Asbestos (NOA) on serpentine outcrops are exposed to abiotic and biotic environmental forces which may affect their toxicity-relevant properties. Lichen colonization and deterioration were examined on monomineralic and polymineralic veins, containing asbestos (chrysotile, tremolite) and/or asbestiform minerals (antigorite, balanergoite, carlosturanite, diopside). The hyphal penetration of Candelariella vitellina within the different veins ranged from few hundreds of micrometers to several millimetres. The various NOA mineral species contacted by lichen hyphae were differently modified in their chemical composition. NOA were incubated in the laboratory with oxalic acid to mimic the dissolution process driven by lichen metabolites bearing acidic and chelating functions, typically pulvinic acid in the case of C. vitellina. The incubated fibres were chemically modified to the same extent of those colonized by C. vitellina in the field and were used to examine the effects of lichen deterioration on the toxicity-relevant release of Reactive Oxygen Species (ROS). Incongruent dissolution patterns observed in the cases of chrysotile-, chrysotile+ balanergoite- and tremolite bearing veins were associated to a significant (even if not complete) inactivation of the surface. Conversely, the veins bearing minerals subjected to congruent dissolution (antigorite, diopside) maintained their reactivity. Lichen colonization may thus be associated to a bioattenuation process on some kinds of fibrous veins, but such effect cannot be generalized to all colonized NOA.

1. Introduction

The six commercially relevant asbestos minerals have been strictly regulated (USA, China) or banned (27 EU countries and 39 more) under worldwide workplace health and safety and public health regulations (International Ban Asbestos Secretariat, 2011). Nevertheless, some concern is growing about the widespread presence of Naturally Occurring Asbestos (NOA; US-EPA, 2011) – asbestos minerals found in place in their natural state, not commercially mined or used – which can be released into the air by natural processes (e.g., rock and soil weathering) or human activities (e.g., agriculture, construction), thus becoming a threat to human health (Culley et al., 2010). Shared scientific and procedural basis for assessing NOA occurrence in a given area and evaluating the associated risk has been invoked in order to develop effective public policies and minimize fibre hazard (Lee et al., 2008).

The natural occurrence of asbestos minerals is mostly associated to serpentine rocks, where the fibres occur within different generations of metamorphic veins (O’Hanley, 1996; Virta, 2005). The largest deposits of asbestos were punctually mined in the serpentine belt of several ophiolitic complexes around the world (eastern Canada, Russian Urals, California, north-eastern Greece, Cyprus and northwest Italy) (Ross and Nolan, 2003). Beside these wide, but rare commercial deposits, small occurrences of fibre-bearing surfaces are extremely common throughout all the serpentine outcrops and may act as sources of fibre dispersion in natural environments (Lee et al., 2008).

The invoked characterization of the potential risk associated to NOA in the case of the widespread, but small sources of NOA, should take into account the following points:

(a)
the six regulated asbestos minerals (namely, chrysotile, tremolite, actinolite, anthophyllite, riebeckite-crocidolite and grunerite-amosite) only partially account for the whole set of fibrous minerals comprised in NOA. In fact many studies have reported on asbestiform minerals not commercially exploited, but sharing a fibrous habit with asbestos, whose pathogenicity is mostly unknown (Baris et al., 1987; Comba et al., 2003; McDonald et al., 2004; Turci et al., 2009; Pugnaloni et al., 2010);

(b)

NOA are more than likely to appear in polymineralic veins, including other fibrous minerals (Groppo and Compagnoni, 2007a);

(c)

opposite to airborne fibres generated by human activities, naturally released fibres from serpentinite outcrops have been often affected by abiotic and biotic environmental forces for a long time, which may have modified their toxicity-related physico-chemical features (Favero-Longo et al., 2005a, 2009a; Daghino et al., 2008).

Serpentinites of the Italian Western Alps contain two asbestos minerals: (a) chrysotile, the serpentine asbestos most widely mined worldwide, always showing fibrous habit, and (b) tremolite, an amphibole which only occasionally occurs with fibrous habit (Groppo and Compagnoni, 2007a). Widespread asbestiform minerals are: (c) antigorite, having the same composition of chrysotile, but different crystalline structure, which in most cases displays a lamellar habit, but occasionally a pseudo-fibrous, columnar habit (Groppo and Compagnoni, 2007b); (d) carlosturanite, a serpentine-type phase (Compagnoni et al., 1985), mostly displaying a fibrous habit; (e) balangeroite, a Fe-rich mineral of the gageite group, mostly found with fibrous habit in association with chrysotile (Compagnoni et al., 1983); (f) diopside, a pyroxene which usually displays prismatic habit, but also occurs as a fibre in chrysotile + carlosturanite bearing veins (Ferraris, 1995; Groppo and Compagnoni, 2007a).

The toxicity of asbestos and asbestiform minerals is currently related to several physico-chemical features acting together such as fibrous habit, high biopersistence and redox-reactive iron ions at the surface, generating Reactive Oxygen Species (ROS) (Kane, 1996; Fubini and Fenoglio, 2007; Ballirano et al., 2008).

In mountain areas, physical environmental forces including temperature changes, freezing and thawing, and washing away by rain, were shown in laboratory experiments to induce a partial dissolution process of chrysotile, tremolite and balangeroite and, consequently, to affect their surface reactivity (Favero-Longo et al., 2009a). On the other hand, serpentinite rocks of mountain areas, including western Alps, are widely colonized by lichen-forming fungi (Favero-Longo et al., 2004), well known agents of physico-chemical deterioration on natural and artificial mineral substrata (Warscheid and Braams, 2000; St. Clair and Seaward, 2004; Gadd et al., 2012). Lichens physically support disaggregation processes through their expanding (wetting) and contracting (drying) thalli adhered to the rock surfaces and the penetration of their hyphae along mineral planes of weakness and intergranular voids (Adamo and Violante, 2000; de los Ríos and Ascaso, 2005). The release of molecules with acidic and chelating functions, leaching and/or complexing metal ions, mostly accounts for lichen-driven mineral dissolution and/or neoformation (Adamo and Violante, 2000). Primary metabolites, particularly the well-known chelator oxalic acid, and/or a wide range of slightly soluble secondary metabolites can be involved in these biogeochemical processes depending on species (Gadd, 1999; Chen et al., 2000).
Lichen communities were described on the asbestos-rich serpentinites of the chrysotile mine of Balangero (Torino, Italy) where the fibre rough texture improves bioreceptivity by increasing water retention and, possibly, propagule deposition (Favero-Longo et al., 2006). Chrysotile fibres contacted by the hyphal penetration component (sensu Favero-Longo et al., 2005b) of several lichen species, including Candelariella vitellina (L.) Müll. Arg., Xanthoparmelia pulla (Ach.) O. Blanco, A. Crespo, Elix, D. Hawksw. & Lumbsch, Xanthoparmelia tinctina (Maheu & A. Gillet) Hale, Lecanora rupicola (L.) Zahlbr. and Acarospora cervina A. Massal., were affected in their chemical composition (Favero-Longo et al., 2009a, b). The octahedral Mg-rich sheets, which compose the serpentine structure in association with tetrahedral silicate sheets, were selectively leached. The finding of oxalates at the interface between the mineral substrate and most of the investigated lichen species suggested oxalic acid as the most common, although not exclusive, leaching agent (Favero-Longo et al., 2007, 2009a, b). The removal of poorly coordinated iron ions, replacing some magnesium in chrysotile and related to its surface reactivity, upon the incubation with lichen chelants (including oxalic acid and lichen secondary metabolites, as pulvinic and norstictic acid), was shown in laboratory following a biomimetic approach (Turci et al., 2007). The induced chemical modification was associated to a partial decrease in the surface reactivity of the fibres, suggesting a bioattenuation role for lichens on asbestos-bearing rocks. The work also indicated that the specific chemical feature of each chrysotile source and, in general, of each fibrous mineral should be independently considered with regard to inactivation.

This paper aims to compare the lichen-driven modification of chemical composition and surface reactivity of chrysotile, occurring in monomineralic or polymineralic veins (chrysotile, chrysotile-balangeroite, chrysotile-diopside-carlosturiane), with that of other asbestos (tremolite) and asbestiform minerals (antigorite).

Four types of fibrous veins uncolonized or penetrated by lichen hyphae in the field were examined by means of light polarizing microscopy and scanning electron microscopy (SEM); elemental compositions of the fibres were obtained with energy dispersive X-ray spectroscopy (EDS). Observations and analyses were focused on veins covered by thalli of C. vitellina, a lichen species extremely common on serpentinites (Favero-Longo et al., 2004), which was ubiquitous on all the four selected vein types.

The asbestos and asbestiform fibres below the thalli, occurring in low amount and difficultly separable from other organic and inorganic materials, were not suitable to carry out the tests on the toxicity relevant surface reactivity. Fibres which were similarly modified in their chemical composition to those found below C. vitellina in the field were thus produced in the laboratory by incubating fresh fibres with oxalic acid. Although oxalic acid may not be secreted by C. vitellina (Favero-Longo et al., 2007, 2009b, this work), it is commercially available and displays a high solubility (14.3 g/100 ml at 25 °C) which makes it mostly suitable to generally mimic in vitro, in reasonable times, the deterioration driven by lichen metabolites with acidic and chelating functions (Turci et al., 2007).

The chemical modification of fibres was experimentally characterized in terms of: i) ion release in the supernatant by atomic emission spectroscopy (ICP-AES) and ii) mode of mineral dissolution (congruent vs. incongruent) by the SEM-EDS analysis of the cation to silicon ratio in solid residuals. The potency to release ROS in the presence of hydrogen peroxide (Fenton-like activity, Fubini et al., 1995) was finally evaluated for unaltered and leached fibres using the spin trapping technique coupled with electron paramagnetic resonance spectroscopy (EPR).

2. Material and methods
2.1. Analysis of lichen deterioration of fibrous minerals in the field

The physical interaction and the chemical deterioration effect of lichens on asbestos and asbestiform minerals were investigated on four types of fibrous veins (A-D) exposed at the surface of serpentinite rocks in Western Alps (Table 1):

Table 1.

Asbestos and asbestiform minerals of serpentinite rocks in Western Alps: ideal and calculated formulas of the investigated samples.

<table>
<thead>
<tr>
<th>Minerals</th>
<th>Abbr.</th>
<th>Ideal formula</th>
<th>Sample (Vein-min.)</th>
<th>Calculated formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serpentinite Chrysotile Ch</td>
<td>Mg₃Si₂O₅(OH)₆</td>
<td>A-Ctl</td>
<td>Mg₃Si₂O₅(OH)₆</td>
<td>Mg₃Si₂O₅(OH)₆</td>
</tr>
<tr>
<td>Antigorite Atg</td>
<td>Mg₃Si₂O₅(OH)₆</td>
<td>B-Ctl</td>
<td>Mg₃Si₂O₅(OH)₆</td>
<td>Mg₃Si₂O₅(OH)₆</td>
</tr>
<tr>
<td>Carlosturanite Car</td>
<td>Mg₃Si₂O₅(OH)₆</td>
<td>C-Mgt</td>
<td>Mg₃Si₂O₅(OH)₆</td>
<td>Mg₃Si₂O₅(OH)₆</td>
</tr>
<tr>
<td>Tremolite Trem</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
<td>D-Trm</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
</tr>
<tr>
<td>Diopside Dio</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
<td>E-Dio</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
</tr>
<tr>
<td>Ralstonite Rag</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
<td>F-Rag</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
<td>CaMg₃Si₂O₅(OH)₆</td>
</tr>
</tbody>
</table>

Vein A, consisting of chrysotile (A-Ctl), from natural, non-mined serpentinite outcrops in the abandoned Balangero mine (Lanzo Valley).

Vein B, consisting of chrysotile (B-Ctl), carlosturanite (B-Cst) and fibrous diopside (B-Dio), from the serpentinite outcrops of Ciampansio (Varaita Valley, Monviso ophiolitic massif). On the basis of the complete dissolution of the vein sample, the relative amount of chrysotile, carlosturanite and diopside was 47:48:5 (see Supplementary Material 1, also including X-ray diffraction analysis).

Vein C, consisting of fibrous antigorite (C-Atg), from the serpentinites of Mompantero (Susa Valley). The serpentine fibres were recognized as antigorite on the basis of Raman analyses (Groppo et al., 2006; see Supplementary Material 2).

Vein D, consisting of fibrous tremolite (D-Trm), from the serpentinite lenses embedded within calcschists of Jovencaux (Susa Valley).

Serpentine samples crossed by the veins were taken both from within the outcrops, where the fibres were not exposed to weathering and biodeterioration forces, and from rock surfaces colonized by C. vitellina. Petrographic cross sections for each vein type were prepared according to Piervittori et al. (1991), examined under transmitted light, using a polarizing microscope Olympus BX4, and, after coating with carbon, by scanning electron microscopy (SEM) in both the secondary electron and back-scattered electron mode, using a Stereoscan 410 Leica electron microscope equipped with a link ISIS EDS apparatus. EDS analyses were performed on the different mineral fibres, either uncolonized or contacted by lichens. The sections were analysed unpolished in order to preserve the possible local surface modification, but preventing the production of fine quantitative analyses. Furthermore, the lichen-derived loss of the original mineral stoichiometries prevented the recalculation of the mineral formula from the EDS analyses. Accordingly, EDS data are reported as atomic weight ratios between main octahedral cations and silicon (Mg/Si for all the minerals, and Ca/Si for diopside and tremolite) in order to highlight the effects of incongruent leaching processes (Turci et al., 2007).
EDS results were statistically analysed by means of ANOVA with post-hoc Student’s t-test or Tukey’s test. SYSTAT release (version 10; SYSTAT, Evanston, IL, USA) was used to perform the tests (p < 0.05).

2.2. Mimicking lichen deterioration in the lab

According to Turci et al. (2007), the deterioration by lichen metabolites with acidic and chelating functions was mimicked by incubating the mineral fibres with the commercially available (Merck Pro-analysis) oxalic acid 0.5 mM.

Veins B, C, D which were examined in field conditions as control were used as source of fibres for incubations. In the case of chrysotile from the Balangero mine, control fibres of the Ctrl-vein A were not sufficient to be processed and used for incubations, so that two different chrysotile veins deriving from the same mining area were used (Table 1):

Vein E: a relatively pure sample of short fibre chrysotile prepared for commercialization (E-Ctl).

Vein F: a mixed field sample (from the same outcrops analysed in the field) of long-fibre chrysotile (F-Ctl) intergrown with brown, Fe-rich asbestiform balanergoite (F-Blg).

Separation and incubation of asbestos fibres. Small bundles of fibres from each vein type were separated from serpentine rocks with Teflon pincers and gently crushed in a hand agate mortar in order to further isolate the fibres and favour their good dispersion in the following incubation step. Each sample was suspended in 0.5 mM oxalic acid, at 25 °C, in the dark, using a solid/liquid ratio of 1 mg mL⁻¹ (200 mg of fibres within 200 mL of solution). Sodium formaldehyde (1%) was added as antiseptic agent. After 35 days of incubation, the fibres were filtered and dried.

Analysis of the solution filtrates. Inductively coupled plasma atomic emission spectrometry (ICP-AES) analyses were performed on the incubating solution in order to evaluate the leaching of calcium, magnesium and silicon from the fibres. Due to the low solubility of iron hydroxides at pH 4.5 (oxalic acid 0.5 mM), the free iron in solution was not searched. The analyses were performed with an IRIS II Advantage/1000 radial plasma spectrometer by Thermo-Jarrel Ash Corp. The optical system is sealed with inert gas, with no moving parts, high-resolution (ER/S) capable. The Echelle grating and dispersion prism monochromator range is extended between 165 and 800 nm, with an optical resolution of 0.007 nm (at 200 nm). The photo device is a CID (charge injection device) camera frozen to −50 °C. As the result of ICP-AES describes an average process concerning the leaching of a representative amount of starting material (200 mg), a single analysis is reported for each supernatant.

Elemental analysis of the solid residuals of the fibres. Energy dispersive X-ray spectroscopy (EDS) coupled with scanning electron microscopy (SEM) was performed on the unpolished, carbon-coated, incubated fibres using a Stereoscan 410 Leica electron microscope equipped with a link ISIS EDS apparatus. The samples were analysed unpolished in order to preserve the possible local surface modification, but preventing the production of fine quantitative analyses. Furthermore, the leaching-derived loss of the original mineral stoichiometries prevented the recalculation of the mineral formula from the EDS analyses.
Accordingly, EDS data are reported as atomic weight ratios between main octahedral cations and silicon (Mg/Si for all the minerals, and Ca/Si for diopside and tremolite, and Fe/Si for balangeroite) to evaluate the congruence or incongruence of the leaching process (Turci et al., 2007).

Evaluation of the surface-induced ROS production. The spin trapping agent 5,5’-dimethyl-pyrroline-N-oxide (DMPO), which gives a relatively stable [DMPO-OH] adduct, was used to detect the formation of the OH radical in aqueous suspension of the fibros samples contacted with H₂O₂ (Fenton activity). Following a well established technique described in previous studies (Fubini et al., 1995), the nature and quantity of the stabilized radical was measured by means of Electron Paramagnetic Resonance (EPR) spectroscopy. Fibres were suspended (22 mg mL⁻¹) in H₂O₂ (0.250 ml, 0.5 M in H₂O), DMPO (0.250 ml, 0.05 M) and phosphate buffer (0.500 ml, 1 M, pH 7.4). The radical formation was evaluated by recording at 10’, 30’, 60’ the EPR spectrum of the [DMPO-OH] adduct. All spectra were recorded on a Miniscope MS 100 (Magnettech, Berlin, Germany) EPR spectrometer. The instrument settings were as follows: microwave power, 10 mW; modulation, 1000 mG; scan range, 120 G; centre of field, approx. 3345 G. The number of radicals released is proportional to the intensity of the EPR signal. The signals were double integrated and numeric values were reported, as nanomoles of radicals normalized per milligramme of fibres, in order to quantitatively represent the production of free radicals by mineral fibres. Blanks were performed in the absence of any fibre. All the experiments were repeated at least twice.

Results of EDS and EPR spectroscopies were statistically analysed by means of ANOVA with post-hoc Student’s t-test or Tukey’s test using SYSTAT 10.

3. Results

3.1. Lichen penetration and deterioration of fibrous veins in the field

All the fibrous vein types examined in the study were penetrated by the lichen C. vitellina. The maximum depth of its well-developed hyphal penetration component ranged from few hundreds of microns to several millimetres depending on the fibre orientation, the grain-size, and the occurrence of fractures. Fig. 1 summarises the wide range of different colonization outcomes found in the field. Chrysotile fibres of vein A were contacted by lichen hyphae down to a depth greater than 2 mm from the rock surface (Fig. 1a); the polymineralic Ctl-Cst-Dio vein B was generally less penetrated (down to av. 200 µm) by Candelariella hyphae (Fig. 1b and c); the rigid and brittle fibrous antigorite of vein C was completely surrounded by lichen hyphae and the related extracellular polymeric matrices (Fig. 1d and e); the brittle and powdery tremolite vein D poorly supported the colonization and the outcrop surface was rarely contacted by Candelariella thalli. However, below the established thalli a dense hyphal net surrounded the needle-like tremolite fibres (Fig. 1f and g). An intergranular rather than intragranular penetration was generally observed, with hyphae occupying and/or generating discontinuities around fibre bundles. A hyphal-driven disaggregation of the colonized veins was observed in the order: vein D > veins A, C > vein B. The grain size of fibrous bundles seems to be the key factor in the degree of disaggregation, with the smaller bundles (tremolite vein D) being more disaggregated than bigger ones.
The medullar layer of *Candelariella* thalli did not contain microcrystalline deposits displaying high birefringence between crossed polars which may indicate the possible occurrence of oxalates, ubiquitously detected on serpentinite rocks in the thalli of several other lichen species (see Supplementary Material 3). Accordingly, no crystals having the typical habit of oxalates were observed throughout the SEM observations. On the other hand, yellowish crystals were observed at the cortex layer of the overall examined thalli and identified on the basis of Raman and UV–Vis spectroscopies as pulvinic acid and its derivatives (pulvinic dilactone and calycin), which typically characterize *C. vitellina* and the whole *Candelariella* genus (see Supplementary Material 3).

The effect of lichen colonization induced chemical modification of the fibres to a very different extent. Chrysotile (A-Ctl) from Balangero outcrops was significantly modified, showing a lower Mg/Si ratio with respect to the uncolonized fibre (−24.9%; Fig. 2a); in the complex Ctl-Cst-Dio vein B (Fig. 2d) a lower Mg-depletion was detected in chrysotile (B-Ctl, Mg/Si: −5.6%) with respect to carlosturanite (B-Cst, Mg/Si: −10.0%), while diopside (B-Dio) was unmodified in both its Mg/Si and Ca/Si ratios; antigorite (C-Atg) contacted by lichen hyphae showed a similar Mg/Si to that of uncolonized one (Fig. 2e); tremolite fibres (D-
Trm) were also not significantly modified by lichens in both Mg/Si and Ca/Si ratios with respect to the control (Fig. 2f).

Fig. 2.
Chemical deterioration (variation in the cation/Si wt% ratio) of asbestos and asbestiform minerals from different vein types. a, chrysotile vein A (Ctl); b, pure-chrysotile sample E (Ctl); c, chrysotile (Ctl) - balangeroite (Bg) vein F; d, chrysotile (Ctl)-carlosturanite (Cst)-diopside (Dio) vein B; e, fibrous antigorite (Atg) vein C; f, fibrous tremolite (Trm) vein D. Mg/Si (light grey bars), Fe/Si (dark grey bars) and Ca/Si (empty bars) measured by SEM-EDS on the fibrous minerals exposed at the surface of serpentinite rocks, uncolonized (Control) and colonized by the lichen Candelariella vitellina (C.v.) in the field, and after incubation in the laboratory with oxalic acid (Ox. 0.5 mM). Data are reported as average of independent
measures (at least four) ± standard error. For each cation/Si ratio of each mineral of each vein type, columns which do not share at least one letter are statistically different (p < 0.05)

### 3.2. Fibre leaching by oxalic acid

In the laboratory, the incubation of the fibrous minerals for 35 days with oxalic acid 0.5 mM determined leaching outcomes mostly similar to those detected in the field below the thalli of *C. vitellina*. In Fig. 2, the “Ox. 0.5 mM” columns report the cations/Si ratios for the laboratory-leached minerals and in Fig. 3 the concentration of the main structural ions found in the supernatant of the leached veins (B–E) is reported.

![Fig. 3.
Concentration of magnesium (○), calcium (○) and silicon (■) in the supernatant of fibre suspensions. Chrysotile (Ctl) vein E, chrysotile-balangeroite (Ctl-Blg) vein F, chrysotile-carlostatic-dioptase (Ctl-Cst-Dio) vein B, antigorite (Atg) vein C and tremolite (Trm) vein D after incubation with oxalic acid 0.5 mM for 35 days in the dark.

Chrysotile “lab” samples from Balangero, both the pure (E-Ctl) and the balangeroite-mixed one (F-Ctl), showed a significant decrease of the Mg/Si ratio (~24.4% and ~26.9%, with respect to the respective controls; Fig. 2b–c), similarly to what detected below lichens in the field. For balangeroite (F-Blg), the Mg/Si ratio was not modified, while the Fe/Si ratio significantly increased (+67.6%), suggesting a preferential removal of Si (and Mg) with respect to Fe (Fig. 2c). Both chrysotile (B-Ctl) and carlostatic (B-Cst) in the polymineralic vein B displayed a Mg/Si decrease (~20.7% and ~29.8% with respect to the controls), indicating a selective solubilisation of the brucitic layer, in this case significantly higher than that detected below *C. vitellina* (Fig. 2d). Unmodified cations/Si ratios in diopside (B-Dio), antigorite (C-Atg) and tremolite (D-Trm) were observed with respect to the respective controls (Fig. 2d–f).

Upon the incubation of pure chrysotile (E-Ctl) a higher concentration of Mg was detected in the supernatant (13.5 mg L⁻¹) with respect to silicon (1.1 mg L⁻¹) (Fig. 3–Vein E), yielding a Mg/Si ratio of approx. 12 consistent with the incongruent dissolution suggested by EDS measurements. By contrast, a similar release of Mg (9.2 mg L⁻¹) and Si (10.3 mg L⁻¹) was observed in the case of the mixed vein F, depending on the parallel leaching contributions of chrysotile and balangeroite (Fig. 3–Vein F).

In the case of the polymineralic vein B, the amount of Mg in the supernatant (10.5 mg L⁻¹), which relied on the independent leaching contributions of each mineral, was higher than that of Si (6.0 mg L⁻¹; Fig. 3–Vein B). Poor Ca occurrence in the supernatant (1.6 mg L⁻¹) indicated the slight solubilisation of diopside (B-Dio), which followed the congruent dissolution indicated by EDS.
Antigorite (Fig. 3-Vein C) also released higher amounts of Mg (8.7 mg L⁻¹) with respect to Si (5.9 mg L⁻¹), but their ratio in the supernatant (Mg/Si = 1.5) was strongly lower than that detected in the case of pure chrysotile. This datum is consistent with the absence of variation in the Mg/Si ratio in the solid residual indicated by EDS.

Tremolite (Fig. 3-Vein D) leaching determined a Mg amount (1.7 mg L⁻¹) in the supernatant much lower than what detected for all the other fibrous samples. Ca was poorly released (2.3 mg L⁻¹), similarly to what observed for diopside in the polymineralic vein B. Both Mg and Ca were however higher than Si (0.8 mg L⁻¹) yielding a relatively high cations/Si ratio in the supernatant [(Mg + Ca)/Si = 4.7]. This indicates a leaching mechanism incongruent in nature, though only few surface atomic layers were affected, as indicated by the low overall dissolution values and the unaltered bulk EDS analysis.

3.3. Effect of fibre deterioration on ROS release

All fibrous samples, including chrysotile-bearing veins, asbestos tremolite and asbestiform antigorite, showed Fenton reactivity, i.e. they released •OH radicals in the presence of hydrogen peroxide (Fig. 4). Different absolute values of radical release were detected for the different veins: these variations depend on several factors (including surface area) that complicate the comparison of the specific fibre reactivity and a direct correlation with their toxic potential, the analysis of which goes well beyond the aim of this paper which is limited to the analysis of the deterioration effect on each fibrous mineral or assemblage.

The different fibrous samples were differently affected in their surface reactivity after they were chemically modified in the lab in a similar way to what was observed below lichen thalli in the field. Pure chrysotile (E-Ctl), the mixed chrysotile-balangeroite sample (F-Ctl and F-Blg) and fibrous tremolite (D-Trm) showed a significant decrease of radical release upon the incubation with oxalic acid 0.5 mM. By contrast, no significant modification of reactivity was detected in the case of the polymineralic vein B and antigorite vein C.

4. Discussion

The hyphal penetration of lichenized- and non-lichenized fungi within a rock and the following metabolite release imply modifications of the chemical and crystallographic features of the contacted minerals (Adamo and Violante,
deteriorated. The biogeochemical action, recorded in the field and mimicked in the laboratory, affects differently the chemical composition of the various fibrous minerals, thus yielding vein-dependent modifications of the surface reactivity.

The incubation with oxalic acid - a well-known chelator able to leach divalent cations from the octahedral mineral layers (Thomassin et al., 1977; Turci et al., 2007) - yields asbestos and asbestiform fibres similarly modified in the chemical composition to those observed in the field below the lichen thalli. Only in the case of chrysotile and carlosturanite in vein B, the chemical modification obtained in the laboratory is higher than that detected below C. vitellina, possibly reflecting the poor hyphal penetration observed within this vein type. However, the absence of oxalate deposits at the Candelariella-substrate interface - already reported on solid culture media (Favero-Longo et al., 2007), on other rock substrates (Jorge Villar et al., 2005) and on asbestos cement roofs (Favero-Longo et al., 2009b) - rules out oxalic acid as the major agent of the Candelariella-driven dissolution processes. On the other hand, the detected mineral deterioration may be related to the release of secondary metabolites such as pulvinic acid, the precursor of the crystalline deposits observed in the cortex layer and characterizing the Candelariella genus (Culberson, 1979). Pulvinic acid, as oxalic acid, displays acidic and chelating functions that make it active in complexing metals, including Mg$^{2+}$, Fe$^{2+}$ and Fe$^{3+}$, in the alkaline range (Hauck et al., 2009) which often characterizes water and soils deriving from serpentinized-ultramafic rocks (Alexander et al., 2007), including asbestos-rich serpentinite sediments (Schereier et al., 1987). Accordingly, the fibres produced upon incubation for 35 days with oxalic acid not only are similarly modified in their chemical composition to those colonized by C. vitellina, but are likely modified through a similar dissolution pathway, thus being suitable for testing the potency to release ROS of lichen-deteriorated fibres. It is worth noting that the very low solubility of Candelariella secondary metabolites (Elix and Stocker-Wörgötter, 2008) prevents to obtain pulvinic acid at a suitable concentration to mimic, in times compatible with laboratory experiments, the mineral dissolution observed in the field after a years- or decades-long lichen colonization. Accordingly, previous experiments showed that leaching effects upon 35 days incubation with saturated pulvinic acid (0.003 mM) are limited to the uppermost atomic layers of the minerals (Turci et al., 2007).

4.1. Chrysotile in pure and mixed veins

Chrysotile, as previously reported with regard to abiotic and biotic leaching systems (Thomassin et al., 1977; Morgan, 1997; Turci et al., 2008; Oze and Solt, 2010), exhibits an incongruent dissolution in all the four studied veins (A, B, E, F). In fact, Mg is leached out more easily than Si, as shown by the decrease of the Mg/Si ratio in the lichen colonized (field) and oxalic acid treated (laboratory) fibres (see Fig. 2a, b, c, d: Ctl columns). Pioneer work by Wilson et al. (1981) reported the complete Mg depletion of chrysotile colonized by Lecanoraatra, yielding amorphous silica as deterioration product. Incongruent ion depletion was also previously reported for other phyllolites, as biotite and muscovite, contacted by lichen thalli (Ascaso and Wierczhos, 1996; Prieto et al., 1997; Wierczhos and Ascaso, 1998).

The incongruent dissolution determines a high Mg/Si ratio in the supernatant of the pure chrysotile from vein E. This depends on the oxalic acid-driven selective uptake of Mg$^{2+}$ and likely Fe$^{2+/3+}$, its minor substitute in the octahedral brucite-like layer. Iron is indeed chelated by several acidic fungal metabolites, including oxalic acid (Gadd, 1999), and its occurrence in chrysotile accounts for the detrimental surface chemical reactivity (Gazzano et al., 2007). Accordingly, the ROS production is significantly reduced after the leaching of vein E in the laboratory (Fig. 4), suggesting that a similar bioattenuation process, likely driven by the Mg- and Fe-chelating activity of pulvinic acid (Hauck et al., 2007, 2009), may occur in the field in the case of the lichen-modified vein A.

The submicroscopic association of chrysotile with balangerite or with carlosturanite and diopside may modulate the impact of lichen deterioration on the chemical reactivity in veins F and B, respectively. In the case of the polymineralic
vein F, the decrease of ROS production by the leached fibres with respect to the control (−85%) is even higher than that detected for the pure chrysotile in vein E (−67%). Balangeroite shows an incongruent dissolution in the opposite direction with respect to chrysotile, with silicon being preferentially removed with respect to the divalent structural cations (mainly Fe) (Favero-Longo et al., 2009a), as indicated by the increase of the cations/Si ratio detected by EDS after incubation and the high amount of silicon found in the supernatant. The incongruent dissolutions of both chrysotile and balangeroite leading to a modified surface chemistry may account for the reactivity loss (Turci et al., 2007).

In polymineralic vein B, chrysotile and carlosturanite, which also displays a serpentine-like structure (Ferraris, 1995), show an incongruent dissolution because of the selective Mg-depletion of the brucitic-like layer, which is even higher in the laboratory than in the field. However, no significant changes are shown in terms of ROS production after leaching. The Ca occurrence in the incubation supernatant and the absence of variation in the Mg/Si and Ca/Si in the solid residuals with respect to control account for a congruent dissolution of diopside, likely related to a persistent radical reactivity (see Section 4.3). However, as their close intergrowth prevents the availability of pure samples for the three phases (Groppo and Compagnoni, 2007a), no certainty is possible on both the dissolution mode and extent of carlosturanite and diopside which could account for the definition of the final responsible for the unvaried surface reactivity following the leaching process.

4.2. Other fibrous minerals

Antigorite shares the same chemical composition with chrysotile, but maintains its surface reactivity unmodified upon leaching. The different crystal structure of the two serpentine polymorphs accounts for two different dissolution modes, with antigorite being congruently dissolved both in the field below lichens and in the laboratory. The congruent dissolution pattern of antigorite is unveiled by EDS analyses and by the evaluation of Mg and Si amounts in the supernatant which are closer than in the case of chrysotile. This dissolution mode, with a higher Si release, suggests a higher renewal of the exposure of the brucitic-like layers at the surface, including the Fe substituting for Mg, finally accounting for the maintenance of the surface reactivity upon leaching.

Chrysotile displays a cylindrical structure with a continuous tetrahedral silica layer exposing a leachable brucitic octahedral layer on the outwards of the fibre, whereas antigorite shows an alternating-wave structure in which the silica tetrahedra are tilted periodically below and above the brucitic layer (Wicks and O’Hanley, 1988). The rare fibrous habit depends on peculiar mechanisms of vein formation (Groppo and Compagnoni, 2007b).

When the brucitic layer is removed, the two polymorphs expose a different structured silica layer, with chrysotile exposing a protonable O (Si–O → Si–OH), previously shared with Mg, and antigorite exposing a bridge O involved in a siloxane bridge that is forced to break upon protonation at acidic pH (Si–O–Si → Si–OH + Si–O–). Such structural features may account for the observed solubility pathways: incongruent for chrysotile and congruent for antigorite).

According to its well-known biopersistence, which accounts for its long durability in human fluids and high toxicity (McDonald and McDonald, 1997), tremolite displays a very low ion release in the supernatant, which justifies the slight variation in the solid residuals and in the field samples below lichen thalli. However, dissolution of cation (Mg, Ca) rich layers, where Fe substitutes for Mg, is significantly higher than that of Si, accounting for a poor, but incongruent dissolution, which justifies the significant decrease in surface reactivity. It is worth noting that another amphibole asbestos, crocidolite which does not occur in Western Alps, but was widely used in asbestos-cement together with chrysotile, also showed a trend towards a poor selective depletion of Mg and Fe with respect to Si below thalli of Candelariella (Favero-Longo et al., 2009b). On the other hand, the non-fibrous amphibole ferrohastingsite contacted by lichen hyphae was shown not to change its chemical and structural identity (Barker and Banfield, 1996).

4.3. Congruent vs. incongruent dissolution and surface reactivity
The overview of the deterioration processes of asbestos and asbestiform minerals below lichens in the field and in oxalic acid solution in the laboratory highlights that the fibre surface reactivity is modulated by the dissolution mode, rather than by the dissolution extent. The more the dissolution is incongruent, i.e. with the selective cation depletion where reactive iron appears as occasional substitute, the more the surface is modified with respect to the control and becomes less reactive. A congruent dissolution – even in the case of antigorite which shows a significant leaching of both Mg and Si – is not associated with a surface modification, the surface being only renewed and thus reactivity maintained. It is worth noting that pure chrysotile from Central Alps (Val Malenco), showing a higher but more congruent leaching of Mg and Si with respect to pure chryso tile from Balangero, showed a significantly lower reduction of the surface reactivity (Turci et al., 2007).

5. Conclusions

The evaluation of the health risk posed by NOA and asbestiform minerals has to take into account surface modifications induced by weathering (Favero-Longo et al., 2009a) and biodeterioration forces on natural outcrops. Present data confirm that the reactivity of fresh fibres cannot be used to assess the hazard of NOA. Lichen deterioration driven by metabolites with acidic and chelating functions, mimicked in the laboratory using the commercially available oxalic acid, modifies chemistry and surface reactivity of fibre-bearing veins to a different extent. The veins exclusively bearing minerals subjected to incongruent dissolution (Veins E, F, D) show a significant (even if not complete) inactivation of the surface, while the veins bearing minerals subjected to congruent dissolution (Veins B, C) maintain their reactivity. Lichens incongruently leach chrysotile both in monomineralic and polyminer alic veins, but in the latter ones the surface reactivity is maintained or diminished according to the associated minerals and their dissolution mode. Although a bioattenuation role of lichens is envisaged for some veins bearing chrysotile and, surprisingly, tremolite, other veins result unaffacted, suggesting that this positive attenuating effect of lichens cannot be generalized for all colonized NOA and has to be evaluated case by case.

Acknowledgements

This research has been carried out with the financial support of Regione Piemonte, Environmental section, in the context of a multidisciplinary project “Asbestos hazard in Western Alps”. The authors are grateful to two anonymous reviewers for manuscript improvement.

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