## LETTER

## SPECIAL COLLECTION: NANOMINERALS AND MINERAL NANOPARTICLES

# Previously unknown mineral-nanomineral relationships with important environmental consequences: The case of chromium release from dissolving silicate minerals

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## ABSTRACT

Risk assessments that take into account the formation of environmentally dangerous hexavalent chromium in Cr-containing mine tailings, and associated soils and sediments, require an understanding of the occurrence and speciation of Cr in silicate minerals and glasses. Silicates are more soluble and generally more susceptible to weathering than the refractory mineral chromite, the principal ore mineral of Cr. Studies at the nanoscale using a combination of advanced sample preparation via microtoming and focused ion beam techniques, in combination with state-of-the art analytical transmission electron microscopy and electron diffraction, reveal the occurrence of chromite nanoparticles held within clinochlore and lizardite grains in chromitite ore (an igneous cumulate consisting primarily of chromite) from the Black Thor Chromium deposit in Northern Ontario, Canada, and the Mistake Mine, Fresno County, California, U.S.A., respectively. Nanoscale examinations of altered chromitite ore samples from the Black Thor deposit after dissolution experiments in sulfuric acid-bearing solutions of pH 2.5 show that clinochlore alters to amorphous silica depleted in chromite nanoparticles. This observation suggests the release of chromite nanoparticles rather than  $Cr^{3+}$  aqueous species during the weathering of chromite-bearing silicate minerals. This will in turn have an impact on the environmental behavior of  $Cr^{3+}$  and its potential oxidation to  $Cr^{6+}$ . The formation of  $Cr^{4+}_{ad}$  species in this case will require either the initial dissolution of the nanoparticles or the oxidation of Cr3+ species on the surface of the nanoparticles, either process being a rate limiting step in the formation of Cr<sup>6+</sup><sub>aa</sub> species.

**Keywords:** Chromite, nanoparticles, silicates, clinochlore, lizardite, California, Black Thor deposit, Ontario, Nanominerals and Mineral Nanoparticles

## INTRODUCTION

Chromium can be potentially harmful in the environment as a result of industrial waste mismanagement, accidental spills, and the presence of Cr-bearing rocks and minerals in aquifers, soils, mines, and mine waste. In ultramafic rocks, Cr is present as  $Cr^{3+}$  which has relatively low toxicity, is a micronutrient, and is relatively insoluble and immobile at neutral to alkaline pH. In contrast,  $Cr^{6+}$  is an environmental toxin and human carcinogen (Fendorf 1995 and references therein) and is highly mobile at neutral to alkaline pH.  $Cr^{3+}$  can be readily oxidized to  $Cr^{6+}$  by naturally occurring  $Mn^{3+/4+}$ -oxides (e.g., Bartlett and James 1979; Eary and Rai 1987; Fendorf 1995; Weaver and Hochella 2003; Oze et al. 2007). Hexavalent Cr can be reduced by organic carbon, bacteria, sulfides, and  $Fe^{2+}$  (e.g., Kamaludeen et al. 2003 and references therein).

Several studies have focused on the Cr chemistry and mineralogy of serpentinite and serpentine-derived soils as Crbearing silicates of the serpentine and chlorite-groups are more susceptible to weathering than the refractory mineral chromite (e.g., Fandeur et al. 2009; Hseu and Iizuka 2013; Oze et al. 2004 2007; Morrison et al. 2015). Elevated  $Cr^{6+}$  concentrations in soil pore water and groundwater adjacent to serpentinite have been reported (e.g., Izbicki et al. 2008; Wood et al. 2010). In other locations, spectroscopic studies on the speciation of Cr in serpentine-derived soils indicated the absence of detectable amounts of Cr<sup>6+</sup> (Hseu and Iizuka 2013; Fandeur et al. 2009).

Nanoparticles (NPs) are known to adsorb and/or structurally incorporate problematic metallic contaminants, highly actively and efficiently, from their immediate environment and transport them over vast distances via fluvial and alluvial processes (e.g., Hochella et al. 2005; Plathe et al. 2010; Yang et al. 2015). Therefore, it is also likely that, the occurrence of nano-size refractory minerals such as chromite, if present as is the case here (see below), play a significant role in dictating the transport, distribution, bioavailability, and toxicity of Cr in the environment. These phenomena will be particularly critical in areas around mine tailings and chromite-rich soils and sediments down hydrologic gradient. We will show in this study that Cr occurs in the form of chromite NPs that are encapsulated within silicate minerals of the recently discovered large chromite deposits in the McFaulds

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Lake greenstone belt in northern Ontario, Canada (i.e., the Black Label, Black Thor, and Big Daddy chromite deposits; Laarman 2013; Weston and Shinkle 2013) and the well-known Mistake Mine, New Idria district, Fresno, California, U.S.A. The presence of chromite NPs in Cr-rich silicate minerals will have significant implications with respect to our understanding of the mobility of Cr and its retention in soils derived from Cr-rich ultramafic rocks.

The strategy used in this study was to assess samples from the two geographically and geologically distinct chromite ore deposits mentioned above to look at solid state Cr distribution in both the fresh ore, and (in the case of the Black Thor ore) simulated acid-mine drainage (AMD) weathered sample capable of releasing Cr into the surrounding environment. In this study, both the fresh and acid-etched ore came from the same sample obtained by placing a fresh ore block into an acidic solution (pH 2.5) for one month. Sampling the edge of the block proved to be sufficient to observe both altered and unaltered minerals. This resulted in excellent control for sampling for the Cr-state of the ore from fresh to an acid-etched, weathered condition.

The two starting materials were as follows: (1) High-grade chromitite ore samples were obtained from the Black Thor chromite ore deposit, Ontario, Canada, through the Ontario Geological Survey. The deposit occurs within a regional volcanic-intrusive complex known as the "Ring of Fire" ultramafic-mafic intrusive suite (Weston and Shinkle 2013). (2) A chromitite ore sample was also obtained from the Mistake Mine, New Idria district, Fresno, California, U.S.A., through the Natural History Museum of Los Angeles Count (catalog no. NHMCAL: 36235). The chromite is hosted by serpentinites that are part of the Franciscan Complex, a complex of metamorphosed ophiolitic rocks uplifted along the subduction zone of the oceanic crust and the continental crust of the North American Plate (Coleman 1996).

#### METHODS

Details on the dissolution experiments, scanning electron microscopy (SEM), and transmission electron microscopy (TEM) studies as well as additional EDS chemical distribution maps recorded in scanning TEM mode (STEM), the location of the extracted focused ion beam sections and selected area electron diffractions pattern (SAED) of all identified phases are listed in the Supplementary Information section (Figures<sup>1</sup> S1–S10).

#### RESULTS

The chromitite samples from the Black Thor deposit and the Mistake Mine, California, contain the Cr-rich silicates clinochlore,  $Mg_3[Si_4O_{10}(OH)_2][MgAl_{1,33}(OH)_6]$  and lizardite,  $(Mg_{2,3}Al_{0,46})$   $Si_2O_5(OH)_4$ , respectively (Fig. 1). The Cr-content in these silicates is on average 3 and 1 wt% Cr, respectively. The chromites from both deposits are Mg- and Al-rich as follows:  $(Fe_{0.5}Mg_{0.5})$   $(Al_{0.6}Cr_{1.4}O_4)$ , Black Thor; and  $(Mg_{0.7}Fe_{0.3})(Al_{0.4}Cr_{1.6}O_4)$ , Mistake Mine, de facto magnesiochromite.

TEM images and EDS chemical distribution maps for Cr indicate the occurrence of chromite NPs within Cr-rich clinochlore and lizardite from the Black Thor deposit and Mistake Mine, respectively (Figs. 1b–1f; Fig.<sup>1</sup>S3). The NPs can be distinguished in bright-field TEM images from the silicate matrix on the basis of their darker color, being less electron beam transparent (Figs. 1b and 1f), and a characteristic *d*-spacing of their lattice fringes (Figs. 1c and 1f). The presence of chromite NPs in clinochlore can also be recognized in SAED patterns in which diffraction spots indicate the presence of a single crystal of clinochlore, whereas diffuse diffraction rings represent exceptionally small chromite NPs (generally less than 3–5 nm) in random orientations (Fig. 1d). Larger chromite particles also occur in the Cr-rich lizardite matrix (Fig. 1e) in which nano-size fragments of different orientations and sizes are intergrown (Fig. 1S10).

After the dissolution experiment, the chromite from the Black Thor deposit is weakly altered displaying only rarely any etch features or secondary mineral coatings on their surface (Fig.<sup>1</sup> S1). Conversely, areas composed of clinochlore are strongly altered and are often characterized by surface coatings enriched in Si and depleted in Mg and Al. The absence of diffraction spots and rings in SAED patterns taken from silica-rich areas in the FIB section extracted from the surface of the altered chromitite indicate the occurrence of an amorphous silica modification (Fig.<sup>1</sup> S5). TEM images indicate the occurrence of fragments of clinochlore in areas composed of mainly amorphous silica (Figs. 2a-2d). Chemical distribution maps (Figs. 2c-2e), TEM images (Figs. 2a-2b) and SAED pattern (Fig. S5) indicate a higher abundance of chromite NPs (and thus Cr concentrations) in clinochlore fragments and in areas containing clinochlore fragments than in areas exclusively composed of amorphous silica. This can be best recognized in clinochlore inclusions embedded in areas of amorphous silica with only traces of chromite NPs (Fig. 2).

## DISCUSSION

This study presents petrographic associations observed only at the nanoscale with mineral relationships that were not expected and that dictate how these minerals geochemically behave in a weathering environment. This has important environmental consequences in the weathering of chromitite ore, even though the two samples we observed are from two distinct geologic environments (an ultramafic-mafic intrusive complex vs. a metamorphosed ophiolite at a convergent boundary).

Specifically, we demonstrate the occurrence of chromite NPs in silicate minerals associated with chromite. Although the areas investigated in this study with FIB/microtome sectioning/TEM are physically limited, our observations provide an alternative model for the speciation of Cr in minerals of the serpentine and chlorite groups; i.e., the common assumption is that Cr replaces Al in octahedral coordination in silicates such as Cr-rich chlorites (Lapham 1958) although the structural incorporation of Cr into the octahedral sites of boehmite, (orthorhombic  $\gamma$ -AlOOH) has been recently questioned by Chatterjee et al. (2016).

The ratio between chromite NPs and Cr structurally incorporated into the observed silicates is difficult to estimate as the NPs are finely distributed within their matrices. However, it can be noted that the number of chromite NPs observed qualitatively tracks with the total concentration of Cr in the silicates, with a higher number of NPs in the clinochlore (per area in the image) from the Black Thor deposit (Fig. 1b; average 3 wt% Cr) vs. in the lizardite from the Mistake Mine (Fig. 1f; average 1 wt% Cr).

The occurrence of larger chromite aggregates in the chromitite

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FIGURE 1. (a) Backscattered electron image of the chromitite ore from the Black Thor deposit, Northern Ontario Canada, indicating cumulus aggregates of chromite (chr) grains with interstitial clinochlore (chl); (b) TEM image of chromite nanoparticle (black dots) in clinochlore matrix (gray); (c) High-resolution TEM image of a chromite nanoparticle in the clinochlore matrix displaying lattice fringes with d=2.05 Å (highlighted with solid white lines); (d) SAED pattern of the clinochlore matrix with diffraction spots and rings corresponding to a clinochlore single crystal and chromite nanoparticles, respectively; (e) TEM image of the sample from the Mistake Mine in California depicting typical morphological features of lizardite (liz); (f) TEM image of chromite nanoparticles within the lizardite matrix; a fast fourier transform (FFT) image of the lattice fringes in this image is shown in the lower right corner, with the spotted ring corresponding to the (400) lattice spacing of chromite.



FIGURE 2. Images of a treated sample chromitite after the dissolution experiment with a sulfuric acid bearing solution. (a) Image in STEM mode indicating the occurrence of clinochlore fragments within an amorphous silica matrix (encircled); (b) TEM image indicating occurrence and absence of chromite nanoparticles in the clinochlore fragment and silica matrix; (c) chemical distribution maps for Si (red) and Cr (green) indicating the absence of Cr in the silica matrix, respectively; (d–e) chemical distribution maps for G) Si (red). Cr (green) and Mg (blue) and (e) Cr (green) indicating a higher abundance of Cr (green) in clinochlore fragments (violet) than in the silica matrix (red). (Color online.)

sample from California and the occurrence of chromite fragments of different orientations and sizes within these aggregates (Figs.<sup>1</sup> S8 and S10) suggest agglomeration of chromite NPs and the growth of chromite through particle attachment (CPA, De Yoreo et al. 2015). Agglomeration of NPs occurs when Brownian diffusion brings particle surfaces into contact with each other and attractive Van der Waals (vdW) forces are greater than repulsive electrostatic double layer (EDL) forces (theory of Derjaguin and Landau 1941 and Verwey et al. 1948; referred to as DLVO theory) (Hotze et al. 2010; De Yoreo et al. 2015).

Dissolution of the chromitite ore under acidic conditions resulted in the formation of amorphous silica in areas formerly composed of clinochlorite. The formation of the silica has been identified on the surfaces of numerous silicate minerals after dissolution experiments under acidic conditions as well as in acidic mine tailings and soils (e.g., Teng et al. 2001; Hochella et al. 2005; Schindler et al. 2009; Schindler and Hochella 2015, 2016) and has been attributed to either a leaching-proton-exchange process (e.g., Schweda et al. 1997) or an interfacial dissolution-reprecipitation (Hellmann et al. 2012). Either way, the formation of amorphous silica during the non-stoichiometric dissolution of clinochlore under acidic conditions has resulted in the release of chromite NPs. The NPs are no longer visible (Fig. 2), and EDS shows that the Cr concentration in the silica is significantly lower than in the former silicate.

Mobility and retention of NPs are mainly controlled by interactions between other suspended particles and between particles and larger mineral surfaces. Retention of NPs is for example favored when mineral surfaces and NPs have opposing charges (Hotze et al. 2010), which was not the case during the dissolution experiments under acidic conditions (pH = 2.5) where the surfaces of amorphous silica (point of zero charge, pH<sub>pzc</sub> = 4.1, Kosmulski 2009) and chromite (pH<sub>pzc</sub> = 6.5, Souza et al. 2012) had both strong positive charges. This is in accordance with the lower abundance of chromite NPs in the amorphous silica than in clinochlore (Fig. 2).

## IMPLICATIONS

Chromium-bearing silicates such as minerals of the serpentine and chlorite group are more susceptible to weathering than chromite and as such the solubility and dissolution kinetics of these minerals control the amount of Cr released to the environment. Chromium can be structurally incorporated into these minerals, replacing Al<sup>3+</sup> in octahedral coordination. The results of this study show that Cr3+ can also occur as chromite NPs in silicates and that these NPs can be released or dissolved during the non-stoichiometric dissolution of the silicates. The release of chromite NPs rather than Cr<sup>3+</sup> aqueous species during alteration of Cr-bearing silicates would have a large impact on the fate of Cr<sup>3+</sup> in the environment as the behavior of NPs is predominantly governed by the interaction between charged mineral surfaces. For example, common alteration products of Fe-rich minerals of the chlorite and serpentine groups under near neutral pH conditions are Fe-hydroxides (Murakami et al. 1996). These minerals would be able to retain chromite NPs due to their large surface area (Fandeur et al. 2009), especially in the pH range of 7 to 8, where the surfaces of chromite NPs and of some Fe-hydroxides can be negatively and positively charged, respectively (Cornell and Schwertmann 1996).

The release of chromite NPs during the weathering of silicate minerals associated with chromite would also result in (1) lower concentrations of dissolved  $Cr^{3+}$  aqueous species (than expected from the total concentrations of Cr in the silicates) and thus in lower rates of  $Cr^{3+}$ -hydroxide formation if conditions favored that, and (2) lower oxidation rates of  $Cr^{3+}$  to  $Cr^{6+}$  as this would require either the dissolution of the NPs or the oxidation of  $Cr^{3+}$  species on the surfaces of the NPs, processes that would be most likely the rate-limiting steps for the oxidation of  $Cr^{3+}$  to  $Cr^{6+}$  in aqueous solution. Future risk assessments of Cr-contaminated soils, tailings, and mine waste dumps must therefore include an estimate on the abundance of chromite NPs in Cr-rich silicates.

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