TECHNICAL ARTICLE

Predicting Geochemical Behaviour of Waste Rock with Low Acid Generating Potential Using Laboratory Kinetic Tests

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Abstract Prediction of contaminated neutral drainage using laboratory kinetic tests designed for acid mine drainage prediction is challenging because of the low metal concentrations generated by low sulfide oxidation rates. Fresh and weathered samples from the Tio mine waste rock piles were submitted to humidity cell tests. The waste rocks were demonstrated to be non-acid generating in the long term, as interpreted by conservative oxidation-neutralization curves. The results demonstrate that even though the main neutralizing minerals react differently after 25 years of natural weathering (with regard to Ca, Mg, Al, and Si release), the response of the fresh waste rocks during humidity cell leaching was very similar to those of the weathered waste rocks, when considering all the elements related to silicate dissolution, including those implicated in secondary phase precipitation. However, Ni generation was

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greater in the weathered waste rocks even though sulfide oxidation rates were similar, as Ni sorption properties reach saturation. Although the Ni concentrations from the leachates of humidity cell tests remain below regulated values, they are bound to increase with continued weathering if no preventive or control measures are undertaken at the site.

Keywords Contaminated neutral drainage · Humidity cell tests - Nickel - Tio mine waste rocks - Weathering

Introduction

The remediation costs of mine waste that generates acid mine drainage (AMD) are about 10 times that of non-acid-generating wastes (Aubertin et al. [2002](#page-16-0); Bussière [2007](#page-16-0)); therefore, accurate AMD prediction is crucial for an integrated mine waste management and proper remediation of mine sites. However, many toxic metals, such as Ni, Zn, Co, As, and Sb, are soluble at near-neutral pH, and can potentially contaminate mine effluents, even without acidic conditions; this phenomenon is called contaminated neutral drainage (CND) or simply neutral drainage (Bussière [2007](#page-16-0); Heikkinen et al. [2009](#page-17-0); Nicholson [2004;](#page-17-0) Pettit et al. [1999](#page-18-0)). CND occurs when sufficient neutralization is available in the mine wastes and/or when sulfide oxidation is sufficiently weak (Heikkinen et al. [2009](#page-17-0)). Prediction techniques developed for AMD generation prediction (Blowes et al. [2003](#page-16-0); Lawrence and Scheske [1997;](#page-17-0) Lawrence and Wang [1997](#page-17-0); MEND [1991;](#page-17-0) Paktunc [1999b](#page-17-0); White et al. [1999\)](#page-19-0) might not be suitable for CND generating sites (Nicholson [2004](#page-17-0)). Li ([2000\)](#page-17-0) demonstrated that for low-sulfide, lowneutralization potential mine wastes, the proportion of dissolved carbonates that effectively neutralize acid

decreases when sulfide oxidation drops below a certain rate, because the bicarbonate ions are dissolved and flushed out of the material without acid neutralization.

Many cases can be found in the literature where Ni CND is generated from tailings with significant carbonate neutralization and various sulfide oxidation levels (Heikkinen and Räisänen [2008](#page-17-0); Heikkinen et al. [2009\)](#page-17-0). It was also demonstrated (Li [2000\)](#page-17-0) that the silicate contribution to neutralization becomes increasingly important as sulfide oxidation rates drop. Many studies are found in the literature where the unoxidized portion of AMD generating tailings show CND-like features such as near-neutral pH and high dissolved nickel levels (e.g. Gunsinger et al. [2006](#page-16-0); Heikkinen and Räisänen [2008](#page-17-0); Heikkinen et al. [2009](#page-17-0); Holmström et al. [2001](#page-17-0); Johnson et al. [2000](#page-17-0); McGregor et al. [1998\)](#page-17-0), where Ni levels are controlled mainly by sorption and/or coprecipitation with iron oxyhydroxides. However, no studies were found where Ni CND resulted from low sulfide oxidation with significant silicate neutralization. The present study focuses on water quality prediction of Ni CND generating mine waste rocks by means of kinetic tests, at a site where acid neutralization is mainly provided by silicate minerals. The tests were performed on waste rocks sampled from the Tio mine, a hematite-ilmenite deposit near Havre-Saint-Pierre, Québec, Canada (Fig. [1](#page-2-0)), exploited since the early 1950's through an open pit operation by Rio Tinto, Iron and Titanium Inc. The gangue material of the Tio ore is mainly composed of a calcic plagioclase mineral close to labradorite composition (approximate formulae $Na_{0.4}Ca_{0.6}Al_{1.6}Si_{2.4}O₈; see Pepin 2009; Planete$ $Na_{0.4}Ca_{0.6}Al_{1.6}Si_{2.4}O₈; see Pepin 2009; Planete$ $Na_{0.4}Ca_{0.6}Al_{1.6}Si_{2.4}O₈; see Pepin 2009; Planete$ et al. [2010a](#page-18-0)). The gangue also contains pyroxene, chlorite, mica and biotite. Effluents from the waste rock piles are near-neutral and sporadically show Ni concentrations slightly higher than those allowed by applicable local regulations (QIT [2005](#page-18-0)). Preliminary studies performed on a concentrated sulfides fraction of the Tio mine waste rocks, prepared from froth flotation, showed that Ni is generated mainly from Ni-bearing pyrite $(F \in S_2)$ and millerite (NiS) that seem to be associated with ilmenite in the Tio ore deposit (Bussiere et al. [2005](#page-16-0)). Other sulfide oxidation products include Co and Zn. A Tio mine waste rock sample previously studied was shown to be non-acid-generating (Bussiere et al. [2005\)](#page-16-0). The work also demonstrated that the waste has important metal retention potential, occurring most probably via surface sorption (Bussiere et al. [2005](#page-16-0); Pepin et al. [2008;](#page-17-0) Plante et al. [2008;](#page-18-0) Plante [2010](#page-18-0)).

Literature Background

The two main gangue minerals found in the Tio mine waste rocks are plagioclases and pyroxenes (Plante et al. [2010a](#page-18-0)), and the waste rocks can contain up to approximately

70–80% residual ilmenite ore. The dissolution mechanisms of these minerals and the formation of layered double hydroxides, which are believed to play an important role in the metal uptake of silicates, are reviewed below.

Plagioclase Reactivity

The plagioclases are a solid solution series within the feldspar mineralogical family. The series range from albite to anorthite end-members with respective compositions of $NaAlSi₃O₈$ and $CaAl₂Si₂O₈$, where sodium and calcium atoms can substitute for each other in the mineral's crystal lattice structure. Plagioclase dissolution have been long known to be incongruent, with Al, Na and Ca being preferably dissolved at the surface (e.g. Blum and Stillings [1995](#page-16-0); Carroll and Knauss [2005](#page-16-0); Casey et al. [1989;](#page-16-0) Inskeep et al. [1991](#page-17-0); Muir et al. [1989,](#page-17-0) [1990a](#page-17-0), [b](#page-17-0); Schweda et al. [1997](#page-18-0)). This dissolution generates an Al–Na-Ca-poor thin layer (which is consequently enriched in Si) at the plagioclase surface (Blum and Stillings [1995](#page-16-0)). The formation of silica enriched residual layers on feldspars in acidic solution is a multi-step process. First, the exchange of $Na⁺$ and Ca^{2+} for H⁺ (Blum and Stillings [1995](#page-16-0); Muir and Nesbitt [1997](#page-17-0); Muir et al. [1990a](#page-17-0), [b](#page-17-0); Scheweda et al. [1997\)](#page-18-0) results in an increase in solution pH (Blum and Stillings [1995](#page-16-0)). Second, Al^{3+} is preferentially released into solution due to the breaking of Si–O–Al bonds preferentially over Si-0-Si bonds (Muir and Nesbitt [1992](#page-17-0), [1997](#page-17-0); Muir et al. [1990a,](#page-17-0) [b;](#page-17-0) Xiao and Lasaga [1994](#page-19-0)), which is also an acidconsuming step (Muir et al. [1990a](#page-17-0), [b\)](#page-17-0). The second reaction step implies that Al is preferentially released from the plagioclase into solution, after which Si is released (Muir and Nesbitt [1997](#page-17-0)). A third step is also suggested for feldspar dissolution, which implies repolymerization of Si–O bonds, releasing acid into solution (Muir et al. [1990a,](#page-17-0) [b](#page-17-0); Schweda et al. [1997\)](#page-18-0). Since labradorite (the plagioclase in the Tio waste rocks, see Table [3](#page-6-0)) is net-neutralizing (Jambor et al. [2007\)](#page-17-0), the overall reaction must be acidconsuming; therefore, the third reaction step must release less acid than was consumed in the first two reaction steps.

According to Blake and Walter ([1999\)](#page-16-0), the thickness and composition of leached layers on feldspars depends on solution composition and pH. Dilute acidic conditions produce relatively thick Si-rich layers, up to several thousands of \AA (e.g. Blum and Stillings [1995;](#page-16-0) Casey et al. [1989](#page-16-0); Hellmann et al. [2003](#page-17-0); Muir and Nesbitt [1992](#page-17-0), [1997](#page-17-0)), whereas dissolution at near-neutral pH produces only thin leached layers of tens of \AA between pH 5 and 8 (Blake and Walter [1999](#page-16-0); Blum and Stillings [1995](#page-16-0); Muir and Nesbitt [1997](#page-17-0)). The circumneutral environments prevailing in the Tio mine waste rock kinetic tests should promote the formation of such thin leached layers (tens of \dot{A}) on the plagioclase feldspar surfaces.

Fig. 1 Geographic location of the Tio mine site

The Al released from labradorite dissolution could precipitate as various possible secondary phases, such as kaolinite ($Al_2Si_2O_5(OH)_4$) or $Al(OH)_3$, as demonstrated by geochemical simulations with VMinteq and JCHESS (Figs. [7](#page-11-0), [8](#page-12-0) respectively). The labradorite reaction with acid leading to kaolinite precipitation can be generalized and simplified as Eq. 1, while the same reaction leading to aluminum hydroxide is generalized as Eq. 2:

$$
Ca_{0.6}Na_{0.4}Al_{1.6}Si_{2.4}O_8 + 1.6H^+ + 2.4H_2O
$$

\n
$$
\rightarrow 0.8Al_2Si_2O_5(OH)_4 + 0.8H_4SiO_4 + 0.4Na^+
$$

\n
$$
+ 0.6Ca^{2+}
$$
\n(1)

$$
Ca_{0.6}Na_{0.4}Al_{1.6}Si_{2.4}O_8 + 1.6H^+ + 6.4H_2O
$$

\n
$$
\rightarrow 1.6Al(OH)_3 + 2.4H_4SiO_4 + 0.4Na^+ + 0.6Ca^{2+}
$$
 (2)

Pyroxene Reactivity

The pyroxenes found in the Tio mine waste rocks are represented mainly by enstatite (6.1–24.9 wt%) in Table [3.](#page-6-0) Enstatite neutralizes acid through Eq. 3 (Oelkers and Schott [2001](#page-17-0); Stefánsson [2001\)](#page-18-0):

$$
MgSiO_3 + 2H^+ + H_2O \rightarrow Mg^{2+} + H_4SiO_4 \tag{3}
$$

Enstatite dissolution proceeds via magnesium releasing exchange reactions between aqueous H^+ and Mg^{2+} in the enstatite structure, followed by the relatively slow detachment of silica from partially liberated tetrahedral chains (Oelkers and Schott [2001](#page-17-0); Oelkers et al. [2009](#page-17-0); Zakaznova-Herzog et al. [2008\)](#page-19-0). Enstatite is believed to exchange about 3.1 hydrogen atoms for each Mg at the reacting sites, following Eq. 4:

$$
MgSiO3 + 3.1H+ \to H3.1SiO31.1 + Mg2+
$$
 (4)

Thus, pyroxenes preferentially release Mg over Si in early dissolution stages, much like the plagioclases preferentially release Ca and Na over Si initially. The preferential release of Mg from pyroxene is expected to lead to the formation of Mg-depleted/Si- and H-enriched surfaces. From XPS (X-ray photoelectron spectroscopy) results of enstatite leached at pH 6, model calculations indicate an Mg-depleted weathered surface layer of only a few atoms thick, while continuous leaching near pH 1 and at elevated temperatures (up to 60° C) leads to formation of a weathered layer of pure silica (Schott et al. [1981](#page-18-0)).

Ilmenite Reactivity

Ilmenite (FeTiO₃) minerals may contain many substitutes within its crystal structure, most notably Mg and Mn (e.g. Lener [1997](#page-17-0), and references therein). Ilmenite dissolution is known to be associated with Mg and Mn release (e.g. Grey et al. [2005](#page-16-0); Hodgkinson et al. [2008;](#page-17-0) Nair et al. [2009](#page-17-0); Schroeder et al. [2002](#page-18-0)); during ilmenite dissolution, Mg and Mn removal is believed to follow Fe release (Lener [1997](#page-17-0)). In the case of naturally weathered ilmenite, Grey and Reid [\(1975](#page-16-0)) proposed a two-stage alteration mechanism, which is currently the generally accepted model (Janßen et al. [2008\)](#page-17-0). In the first stage, ilmenite undergoes weathering through oxidation and removal of Fe to form a transitional phase consisting of an apparently continuous series of compositions from ilmenite to pseudorutile (ideally $Fe₂Ti₃O₉$ (Eq. 5):

$$
\text{FeTiO}_3 + 2\text{H}^+ + \frac{1}{2}\text{O}_2 \rightarrow \text{Fe}_2\text{Ti}_3\text{O}_9 + \text{Fe}^{2+} + \text{H}_2\text{O} \qquad (5)
$$

The Fe is assumed to diffuse out through the unaltered oxygen lattice. In the second stage, pseudorutile undergoes incongruent dissolution (Eq. 6), resulting in the formation of rutile (TiO₂) and/or leucoxene (Frost et al. 1983 ; Nair et al. [2009\)](#page-17-0), which is believed to be the ultimate alteration product of ilmenite (Lener [1997](#page-17-0); Nair et al. [2009](#page-17-0); Schroeder et al. [2002\)](#page-18-0):

$$
Fe_2Ti_3O_9 + 4H^+ \rightarrow 3TiO_2 + 2Fe^{2+} + 2H_2O + \frac{1}{2}O_2
$$
 (6)

It was also proposed (White and Peterson [1996](#page-18-0); White et al. [1994](#page-18-0)) that ilmenite could dissolve in anoxic conditions at pH 1–7 following Eq. 7:

$$
\text{FeTiO}_3 + 2\text{H}^+ \rightarrow \text{TiO}_2 + \text{Fe}^{2+} + \text{H}_2\text{O} \tag{7}
$$

Ilmenite dissolution is a neutral process (it neither consumes nor produces acid) at near-neutral/oxic conditions such as those encountered in the present study, because the oxidation and hydrolysis of each ferrous ion released by ilmenite dissolution will produce $2H^+$ (Eqs. 8) and 9), which are equivalent to the H^+ consumed by each ilmenite or pseudorutile while dissolving:

$$
Fe^{2+} + \frac{1}{4}O_2 + H^+ \rightarrow Fe^{3+} + \frac{1}{2}H_2O
$$
 (8)

$$
\text{Fe}^{3+} + 3\text{H}_2\text{O} \to \text{Fe(OH)}_3 + 3\text{H}^+ \tag{9}
$$

Layered Double Hydroxides (LDH)

Metal uptake by mineral surfaces may be attributed to a number of different processes. Sparks (2001, and references therein) indicates that sorption of metals such as Ni and Co on soils results in the formation of metal hydroxide precipitate phases. In the case of Al-bearing soil mineral sorbents, the precipitates are metal-Al hydroxides called layered double hydroxides (LDH) (e.g. d'Espinose de la Caillerie et al. [1995](#page-16-0); Eick et al. [2001;](#page-16-0) Roberts et al. [1999](#page-18-0); Scheidegger et al. [1997](#page-18-0), [1998;](#page-18-0) Scheinost and Sparks [2000](#page-18-0); Scheinost et al. [1999](#page-18-0); Sparks [2001;](#page-18-0) Yamaguchi et al. [2002](#page-19-0)).

These superficial precipitates occur at metal releases far below theoretical monolayer coverage, in a pH-range well below that where the precipitation of metal hydroxides would be expected based on thermodynamic solubility, and at time scales as fast as 15 min (Scheidegger et al. [1998](#page-18-0); Sparks [2001](#page-18-0)). It was established that such LDH phases preferentially form in the presence of Al-containing sorbents like kaolinite, gibbsite, and alumina above pH 7.0 (Scheinost and Sparks [2000\)](#page-18-0). Ni–Al LDH was established to form on gibbsite of low-surface area over pH 7.5, while on high-surface area gibbsite, Ni was sorbed as an innersphere complex (Yamaguchi et al. [2002](#page-19-0)). It was also shown to form on a soil clay fraction (composed of aluminum hydroxy interlayered vermiculite, kaolinite, mica with minor amounts of gibbsite and quartz) at pH 7.5 (Roberts et al. [1999\)](#page-18-0), less so at pH 6.8, but not at pH 6.0. Moreover, Ni sorption rates significantly increased with pH. It was also demonstrated that LDH were thermodynamically and/ or kinetically favored rather than the formation of hydroxides when an Al-releasing sorbent is available at near-neutral pH (Scheinost et al. [1999;](#page-18-0) Peltier et al. [2006](#page-17-0)).

It was also demonstrated that Ni–Al LDH become increasingly stable as they age and that over time, these precipitates will transform into more stable Ni-phyllosilicates (Ford et al. [1999](#page-16-0); Peltier et al. [2006;](#page-17-0) Roberts et al. [1999;](#page-18-0) Scheckel and Sparks [2001;](#page-18-0) Scheckel et al. [2000](#page-18-0)).

Materials and Methods

Characterization Methods

The specific gravity (Gs) of the waste rock samples was determined with a Micromeritics helium pycnometer. The specific surface area (S.S.A.) was determined by using a Micromeritics surface area analyzer using the B.E.T method (Brunauer et al. [1938](#page-16-0)). The grain size distribution was determined by sieving for the fractions between 10 cm and $0.355 \mu m$ and by a laser diffraction grain size analyzer for the $\langle 0.355 \text{ um fraction using a Malvern} \rangle$ Instruments Mastersizer S. The Tio mine waste rock chemical analysis was performed using acid digestion $(HNO₃-Br₂-HF-HCl)$ followed by ICP-AES analysis for over 20 elements. Silica is partially evaporated during the digestion procedure and therefore is not reported in this study. Sulfide sulfur was determined by subtracting the sulfate sulfur (determined by a 40% HCl extraction; method adapted from Sobek et al. [1978](#page-18-0)) from the total sulfur (ICP-AES analysis). The waste rock samples mineralogical characterization was performed with a Bruker A.X.S. D8 advance x-ray diffraction (XRD) instrument equipped with a copper anticathode. Mineralogical quantification was performed with Rietveld [\(1993](#page-18-0)) fitting of the XRD data with TOPAS software, with a detection limit and precision of approximately 0.1–0.5 wt%. Acid–base accounting (ABA) was determined following the protocol prescribed by Lawrence and Wang [\(1997](#page-17-0)). The acid-generation potential (AP) was calculated assuming that the sulfide sulfur content was exclusively expressed as pyrite and that all pyrite was available for oxidation, and converted to calcite equivalents (kg $CaCO₃/t$) by multiplying by a factor of 31.25. The neutralization potential (NP) was determined by HCl additions followed by back-titration of the excess acid to a pH 8.3 endpoint. The net neutralization potential (NNP) is the difference between the AP and NP $(NNP = NP-AP)$, and the NP/AP ratio was also determined for interpretation purposes. Scanning electron microscope (SEM) observations of backscattered electrons (BSE) were made on a Hitachi S-3500 N microscope equipped with an x-ray energy dispersive spectrometer (EDS) analyzed with INCA software, at 20 kV and $100-130 \mu\text{A}$. Oxygen was stoichiometrically determined by EDS analyses.

Kinetic Prediction Procedure

The geochemistry of the Tio mine waste rock was evaluated using humidity cell tests (ASTM D5744-96 [2001](#page-16-0)), which consist of weekly drying-wetting cycles ending with flushing of the studied material (1 kg) with deionized water (1 L), and analysis for various geochemical parameters such as pH and metals concentrations in the leachates (using ICP-AES). These cell tests were performed for 76 cycles (539 days) on the \lt 6.3 mm fraction of the mine wastes (believed to be the most reactive, e.g. Price and Kwong [1997\)](#page-18-0) without grinding, since that would generate unwanted fresh surfaces in the weathered waste rock samples.

Geochemical Simulations

Geochemical speciation models are widely used to describe solid/water chemistry in mine drainage waters (e.g. Alpers and Nordstrom [1999;](#page-16-0) Blowes et al. [2003;](#page-16-0) Bussiere et al. [2004](#page-16-0); Villeneuve et al. [2003\)](#page-18-0). Such models calculate the ion activities and speciation over a wide range of conditions (such as temperature, pressure, concentration, pH, and Eh), and are used to calculate the saturation indexes of a wide variety of minerals and of the distribution of the elements between the aqueous, gaseous, and solid phases. Two different geochemical speciation models were used in this study: Visual Minteq (Felmy et al. [1984\)](#page-16-0), which is a windows version of the MINTEQA2 ver.4.0 database (USEPA [1999](#page-18-0)), was employed for saturation index calculations, and; JCHESS (van der Lee and De Windt [2002\)](#page-18-0), a graphical front-end of the CHESS model built from the EQ3/6 database (Wolery [1992](#page-19-0)), was employed for Eh–pH diagrams.

Samples Tested

Since the Tio mine has been in operation since the early 1950's, the waste rock piles have been submitted to natural weathering phenomena (up to over 5 decades at the time of sampling). Moreover, the waste rock composition varies significantly, as the target cut-off for ore processing is 76% hemo-ilmenite. Therefore, the hemo-ilmenite content of the waste rocks range between 0 and 76%. Also, the mine exploits two different ore bodies (Tio and Northwest) that have slightly different compositions (Plante et al. [2010a\)](#page-18-0). The waste rock samples were carefully selected to represent the waste rock pile heterogeneity in terms of composition, origin, and age. Table [1](#page-5-0) lists the general characteristics of the samples selected for the present study, as well as the ilmenite levels target during sampling $(C1\leq$ $C2 < C3$ for fresh and $C4 < C5 < C6$ for weathered

unitless)

Table 1 Physical characteristics of the waste rock samples studied (D_x in μ m, refers to the size for which x% of the sample is under this grain size)

samples). All samples were screened to $<$ 10 cm at the mine site.

Characterization Results

The physical characteristics of the Tio mine waste rock samples are summarized in Table 1. The weathered waste rock samples are generally finer than the fresh waste rock samples, as indicated by the D_{10} (C1–C3: 163–320 μ m, C4–C6: 73–174 μ m), D₅₀ (C1–C3: 1,285–2,000 μ m, C4–C6: 1,200–1,305 μ m), D₉₀ (C1–C3: 4,310–5,050 μ m, C4–C6: 4,380–4,860 μ m) and % <80 μ m (C1–C3: 4.7– 7.1%, C4–C6: 6.7–10.4%) values. The chemical element analyses most relevant for the purpose of this study are shown in Table 2. Fe (19.4–33.7 wt%) and Ti (9.84–17.6 wt%), associated mainly with ilmenite and hematite, follow the levels targeted in the sample preparation ($C1 < C2 < C3$ and $C4 < C5 < C6$). Since Al, Ca, and Mg are associated with gangue minerals, their levels follow the inverse of Fe and Ti and vary from 3.54 to 8.68 wt% for Al, from 1.45 to 3.65 wt% for Ca and from 1.12 to 3.14 wt% for Mg in the waste rock samples. Mg levels are generally higher in weathered samples (2.34–3.14 wt%) than in fresh ones (1.12–1.75 wt%). Ni (280–430 ppm), Co (240–450 ppm), Cu (80–170 ppm), Mn (52–90 ppm), and Sb (270–470 ppm) levels are generally correlated with Fe and Ti. S_{sulfide} values are significantly higher in the fresh waste rocks than in the weathered ones, most probably because of weathering (sulfide oxidation). S_{sulfide} values range between 0.335 and 0.466 wt% in the fresh waste rock samples and from 0.134 to 0.167 wt% in the weathered ones. It was previously demonstrated (Pepin [2009](#page-17-0); Plante et al. [2010a\)](#page-18-0) that Ni in the Tio mine waste rocks was mainly mineralogically associated with the ilmenite and sulfides.

The ABA of the Tio mine waste rock samples are also shown in Table 2. The AP values of the fresh waste rock samples (10.5–14.6 kg $CaCO₃/t$) are higher than those of the weathered ones $(4.2-5.2 \text{ kg } CaCO₃/t)$. The NP values range from 6.1 to 8.6 kg $CaCO₃/t$. Consequently, all the NNP values fall inside the uncertainty zone of the test $(\pm 20 \text{ kg } CaCO₃/t)$ as defined by SRK [\(1989](#page-18-0)). NP/AP ratios

Table 2 Chemical and ABA characterization of the studied samples (elemental values in wt%, AP, NP, NNP in kg $CaCO₃/t$, NP/AP

range between 0.6 and 1.7 and classify as likely (when $\lt 1$) or possibly (between 1 and 2) ARD generating according to the Price et al. [\(1997](#page-18-0)) criterion. However, the NP values are quite low and the uncertainty of the method $(\pm 2 \text{ kg})$ CaCO₃/t, based on a 39 kg CaCO₃/t reference sample; Plante [2005\)](#page-18-0) is quite high for the low NP values, leading to relative errors from 23 to 33%. Thus, ABA interpretation from such low values is difficult.

The XRD quantification results are presented in Table [3.](#page-6-0) Pyrite is the main sulfide mineral detected (1.1–3.0 wt%), while chalcopyrite was detected at trace levels (0.1–0.6 wt%); no nickel sulfide was detected by XRD in the samples studied. The ore minerals ilmenite (23.0–45.2 wt%) and hematite (7.9–20.1 wt%), referred to as hemo-ilmenite since

they occur as exsolution lamellae in the Tio mine deposit, confirm the variation following the low, intermediate, and high ilmenite levels targeted in sample preparation $(C1\lt C2)$ \langle C3 and C4 \langle C5 \langle C6). The plagioclase mineral labradorite (main gangue mineral in the Tio deposit) levels are inversely proportional to the hemo-ilmenite content and ranges between 16.0 and 44.6 wt%. Other important gangue minerals in the Tio mine waste rock samples include the pyroxenes enstatite and pigeonite, with levels up to 24.7 and 3.7 wt% respectively. The micas biotite and muscovite $(\leq 3.1 \text{ wt\%})$, chlorites $(\leq 3.3 \text{ wt\%})$, K-felsdpar orthoclase (up to 6.8 wt%), rutile $(\langle 2.0 \text{ wt\%})$, and spinel (up to 6.5 wt%) complete the Tio mine waste rock samples mineralogy.

Results and Interpretation

Humidity Cells

As shown in Table 3, no carbonate mineral was detected in the Tio mine waste rocks. Consequently, neutralization of the acid resulting from sulfide oxidation is provided mainly by silicate minerals, particularly by the calcic plagioclase mineral (identified as labradorite in the XRD analyses) and the pyroxenes (enstatite and pigeonite), and possibly also from the minor gangue minerals phlogopite mica (biotite) and chlorite, based on the classification of silicate dissolution in static ABA testing (Jambor et al. [2007](#page-17-0)). The dissolution products considered in the present study that come from the neutralizing processes are Ca, Al, Mg, and Si. According to the classification of the relative mineral reactivities at pH 5 of Kwong [\(1993](#page-17-0)), pyroxenes react faster than plagioclases. However, the Jambor et al. ([2002,](#page-17-0) [2007](#page-17-0)) NP comparisons on pure minerals showed that the plagioclase mineral labradorite neutralizes more acid than pyroxenes.

The humidity cell tests results are shown in Figs. [2](#page-7-0) and [3](#page-7-0). The pH of all the cell leachates remained near-neutral over 529 testing days. The weathered waste rocks generally generate leachates with lower pH values (6.5–8) than the fresh waste rocks (7.5–9.0) during the first 150 days, after which no significant differences were observed (pH between 6.5 and 8.5). The electric conductivities of the leachates stabilized rapidly between 10 and $100 \mu S/cm$, and remained so until the last leaching cycle. Leachate Ca concentrations stabilized after approximately 50 days; the weathered waste rocks cells generated lower Ca levels $(1-2 \text{ mg/L})$ than the fresh ones $(2-10 \text{ mg/L})$. The Al concentrations from the fresh waste rock cells diminished from the beginning of the test until approximately 400 days, after which they increased until the end of the test. Al concentrations generated from the weathered samples remained very close to or below the analytical detection limit (0.01 mg/L) throughout the tests. The weathered waste rock cells generated higher Mg and Si levels than fresh ones (Mg: up to 0.13 mg/L for C1–C3 and up to 0.61 mg/L for C4–C6; Si: \lt 5 mg/L for C1–C3 and up to 22 mg/L for C4–C6). No significant differences between S (sulfates) values were noticed between the fresh and weathered waste rocks (0.3–1.1 mg/L). The Ni values were generally less than the detection limit (0.004 mg/L) for the fresh waste rock leachates and the C5 sample. The Ni loadings of the C4 and C6 weathered waste rock leachates stabilized between 0.005 and 0.010 mg/L after the first 130 days. Co and Zn levels were generally close to

Mineral	Formulae	C ₁	C ₂	C ₃	C ₄	C ₅	C ₆
Labradorite	(Ca, Na) $(Si, Al)4O8$	44.6	19.1	16.0	32.2	31.3	20.3
Ilménite	FeTiO ₃	23.0	37.0	45.2	23.2	26.9	41.2
Hématite	Fe ₂ O ₃	8.1	9.3	20.1	7.9	8.6	16.5
Pyrite	FeS ₂	1.3	3.0	1.6	1.1	1.3	1.7
Chalcopyrite	CuFeS ₂	0.5	0.5	0.2	0.4	0.6	0.1
Enstatite	$Mg_2Si_2O_6$	6.1	12.9	8.5	24.7	20.4	13.9
Pigeonite	(Mg, Fe^{2+}, Ca) $(Mg, Fe^{2+})Si_2O_6$	3.7	3.2	0.5	2.1	2.8	2.2
Biotite	$K(Mg, Fe^{2+})_3$ (Al, $Fe^{3+})Si_3O_{10}(OH, F)_2$	1.6	1.7	0.4	1.0	1.1	1.0
Muscovite	KAl ₂ (Si ₃ Al) O_{10} (OH,F) $_2$	0.1	1.1	0.5	1.2	0.7	0.0
Chlorite	$(Mg,Fe,Al)_{6}(Si,Al)_{4}O_{10}(OH)_{8}$	2.8	2.9	0.9	3.3	3.1	0.0
Orthoclase	KAlSi ₃ O ₈	6.8	0.6	4.5	1.2	0.5	0.9
Rutile (anatase)	TiO ₂	1.0	1.5	0.6	1.0	1.9	0.9
Spinel	MgAl ₂ O ₄	0.2	6.5	0.7	0.1	0.2	0.7
Total		99.8	99.2	99.4	99.4	99.1	99.4

Table 3 Mineralogical characterization of the kinetic and weathering cells materials by XRD and Rietveld quantification

not shown)

Fig. 2 Evolution of pH, conductivity, Ca, Al, Mg, and Si concentrations in humidity cell tests on Lac Tio waste rocks (note the log scale for conductivity, Ca, and Mg; dashed lines represent the ICP-AES detection limit)

the detection limit and are not relevant for the purpose of this study. Fe levels were below the detection limit most of the time in all cells; therefore, they are not shown.

Mineralogical Evolution During Tio Mine Waste Rock Weathering

The depletion curves for Ca (indicative of plagioclase dissolution) and S (as sulfates, indicative of sulfide oxidation) in humidity cells are shown in Fig. [4.](#page-8-0) The depletion curves show a first stage characterized by a rapid elemental decrease, followed by stabilization in a second stage, generally after 10–50 days. Such behavior was also observed by others (Benzaazoua et al. [2004;](#page-16-0) Cruz et al. [2001;](#page-16-0) Scharer et al. [1991,](#page-18-0) [1994](#page-18-0)). The rapid elemental depletions observed in the first depletion stage and the subsequent slowdowns are explained by at least three mechanisms: (1) dissolution of oxidation products and readily soluble phases, (2) fine particle disappearance due to complete dissolution and (3) surface passivation due to secondary minerals precipitation (Benzaazoua et al. [2004](#page-16-0); Cruz et al. [2001;](#page-16-0) Scharer et al. [1991](#page-18-0); Villeneuve et al. [2003](#page-18-0)). The Ca depletion slopes of the fresh samples are steeper than those of the weathered ones, while the S depletion slopes of the weathered samples are more similar to those of fresh ones. Ni depletions were negligible and are not reported in Fig. [4.](#page-8-0)

The evolution of the cumulative normalized loadings (mg/kg) in the flushed waters shown in Fig. [5](#page-8-0) enable the determination of the elemental release rates (presented in

Table [4](#page-9-0)), by linear regression within the stabilized portion of the curves. It is important to point out the difference between the weathering rate and the release rate. According to Sapsford et al. [\(2009](#page-18-0)), the weathering rate is defined as the rate (mass per unit time, often normalized to unit mass or unit area) at which a primary mineral is transformed into secondary products, whether soluble or insoluble, congruently or incongruently, whereas the release rate is the rate at which an element or species is driven away from a unit mass of rock per unit time. In kinetic tests such as those used in the present study, the release rate is stoichiometrically related to the weathering rate of a given primary mineral only when the considered reaction products are entirely flushed (not retained in the material).

The applicability of the linear regressions used for the determination of the release rates are evaluated by the determination coefficients (R^2) shown in Table [4.](#page-9-0) Most R^2

values in Table [4](#page-9-0) are close to unity ($R^2 > 0.97$), enabling interpretation and comparison of these release rates, especially for Ca, Mg, Si, and S. However, some R^2 values are farther from unity (\mathbb{R}^2 as low as 0.37, Table [4](#page-9-0)), especially for elements with concentrations very close to the analytical detection limits in the leachates, like Al and Ni. The S release rates are similar for all samples studied and vary between 0.040 and 0.071 mg/kg/d (0.120 and 0.213 mg/kg/d as sulfates) in humidity cell tests. The S release rates obtained from the fresh and weathered waste rocks are of the same order. Assuming that all the sulfate ended up in the leachates, as no S-bearing oxidation products are suspected to form under the conditions of the kinetic tests (Online Supplementary Figures 1 and 2c), the S release rates are indicative of the sulfide weathering rates. The very low sulfate levels in the post-testing solids $(\leq 30 \text{ ppm},$ not reported in this study; see Plante [2010\)](#page-18-0) support this

Fig. 5 Cumulative and normalized humidity cell loadings for oxidation/neutralization-related elements

Table 4 Elemental release rates and associated determination coefficients (R^2) from humidity and weathering cells tests (>200 days)

	Release rates (mg/kg/d)											
	Ca	R^2	S	R^2	Si	R^2	Mg	R^2	Al	R^2	Ni	R^2
C ₁	0.429	0.998	0.047	0.993	0.159	0.992	0.012	0.996	5.8E-03	0.974	4.7E-06	0.368
C ₂	0.245	0.995	0.040	0.988	0.119	0.991	0.008	0.991	1.8E-03	0.865	1.5E-04	0.860
C ₃	0.495	0.998	0.071	0.993	0.212	0.997	0.016	0.989	4.8E-03	0.972	$1.2E-04$	0.948
C ₄	0.108	0.999	0.052	0.995	0.267	0.969	0.031	0.998	$7.2E-04$	0.903	1.8E-04	0.792
C ₅	0.120	0.994	0.056	0.992	0.246	0.967	0.023	0.994	6.8E-04	0.851	$6.0E-0.5$	0.772
C ₆	0.088	0.997	0.050	0.994	0.319	0.976	0.021	0.998	1.1E-03	0.893	$4.5E-04$	0.990

assumption. Consequently, sulfide oxidation appears to occur at similar rates in the fresh and weathered waste rock samples when submitted to humidity cells.

The Ca release rates were significantly higher in the fresh waste rocks samples (0.245–0.495 mg/kg/d) than in the weathered ones (0.088–0.120 mg/kg/d), as expected from the previously explained plagioclase reaction mechanism (preferential Ca release form plagioclase in early weathering stage). Since no secondary Ca minerals should have precipitated, according to the thermodynamic simulations (Online Supplementary Figures 1 and 2), the Ca leached from the plagioclase is expected to have remained in solution during the kinetic tests. Consequently, the Ca release rates in the fresh waste rock samples are expected to be stoichiometrically related to the plagioclase weathering rate.

In the fresh waste rock samples, the Al release rates decreased at approximately 250 days, while the Si release rates increased after 400–450 days. The kinetic tests gave higher Si release rates in weathered samples, which suggest that weathered plagioclase surfaces in the Tio mine waste rock samples are already Ca and Al depleted, in accordance with the first two plagioclase reaction steps previously mentioned. As demonstrated in Online Supplementary Figure 1, there is only a slight difference in the Saturation Index values of the Si-bearing secondary minerals (kaolinite, imagolite, halloysite) between fresh and weathered waste rock samples. Therefore, secondary mineral precipitation in fresh waste rocks doesn't seem to be the main reason explaining the higher Si release rates in weathered samples. The higher Si release rates and lower Ca release rates in weathered waste rocks are consistent with the development of a Ca-poor, Si-rich layer over the weathered plagioclase surface; once the Ca is leached from the surface layer of the plagioclase, more Si is released into solution, as expected from the plagioclase reaction steps previously discussed. The third plagioclase dissolution step (Muir et al. [1990a](#page-17-0), [b;](#page-17-0) Schweda et al. [1997](#page-18-0)) is believed to release some of the acid adsorbed in the first two steps. The generally lower pH values obtained in the leachates obtained from weathered samples in humidity cell tests support this mechanism.

The Mg release rates, related mostly to pyroxene dissolution but also probably to ilmenite in lesser amounts since ilmenite releases Mg during its dissolution (e.g. Grey et al. [2005](#page-16-0); Hodgkinson et al. [2008](#page-17-0); Nair et al. [2009;](#page-17-0) Schroeder et al. [2002](#page-18-0)), were higher in weathered waste rocks (0.021–0.031 mg/kg/d) than in fresh ones (0.008–0.016 mg/kg/d), suggesting that pyroxenes and/or ilmenite seem to dissolve faster in weathered waste rocks. However, it is impossible from these results to specifically distinguish the behaviors of ilmenite and pyroxene. SEM images of altered ilmenite grains from the weathered sample C6 are shown in Fig. [6](#page-10-0). Ilmenite grains in Fig. [6](#page-10-0)a, b, and d show obvious weathering of ilmenite preferentially over hematite. The EDS analyses of the weathered ilmenite shows almost complete Fe depletion, suggesting the remaining phase is something between pseudo-rutile (ideally, $Fe₂Ti₃O₉$) and leucoxene (believed to be similar to rutile, $TiO₂$). On every altered ilmenite grain observed from the present study, there are clear boundaries between the altered and primary ilmenite, which has also been observed in other studies (Mücke and Bhadra Chaudhuri [1991](#page-17-0); Nair et al. [2006,](#page-17-0) [2009](#page-17-0)). Each ilmenite grain in Fig. [6](#page-10-0) bears alteration products on the surface, even the apparently unaltered ilmenite grain shown in Fig. [6](#page-10-0)c; EDS analyses suggest that these secondary phases are mainly composed of Si, Al, and Mg. Moreover, thermodynamic simulations with JCHESS (Online Supplementary Figure 2b) suggests Mg-montmorillonite ((Na,Ca)_{0.33}(Al,Mg)₂Si₄O₁₀(OH)₂·nH₂O) precipitation in the typical conditions encountered in the humidity cell tests in this study. More insight is needed to understand the nature of this secondary phase. This mineral is not included in the VMinteq database used in the thermodynamic simulations, which explains its absence in the possible secondary phases considered in Online Supplementary Figure 1. Finally, the Ni concentrations in the drainage waters are often below the analytical detection limit. Therefore, the rates had poor R^2 values (0.37–0.86) for most of the waste rock samples; only the C3 and C6 samples had high R^2 values (>0.95). It is therefore hazardous to link the Ni release rates to any mineralogical weathering rate in the present study.

Fig. 6 SEM (BSE mode) images of altered hemo-ilmenite grains (Alt: Si, Al, Mg secondary phase; H: hematite; Ilm: ilmenite; PR: pseudorutilelike phase)

The differences in plagioclase weathering levels between samples are illustrated by plotting the cumulative normalized loadings of Ca vs. Si, and Al vs. Si, in Fig. [7a](#page-11-0) and b, respectively. It can be seen that the slopes of the Ca and Al vs. Si loadings from the fresh waste rocks tend to decrease over time. However, the slopes corresponding to the weathered waste rocks are significantly lower than those of fresh waste rocks. These tendencies show that weathering of the plagioclases within the fresh waste rocks in humidity cells have not reached the weathering level of the weathered waste rocks samples, even though there is a shift of the slopes from the fresh samples towards those of the weathered samples in Fig. [7a](#page-11-0) and b.

In the near-neutral conditions encountered in this study, Al could precipitate as various possible secondary phases like Al(OH)₃, kaolinite (Al₂Si₂O₅(OH)₄), or Mg-montmorillonite, as demonstrated by geochemical simulations with VMinteq and JCHESS (Online Supplementary Figures 1a, 1b and 2b). Consequently, Al precipitation as secondary minerals (such as kaolinite, Eq. [1\)](#page-2-0) in the conditions of the kinetic tests could explain the low Al levels. No Al secondary minerals were detected by XRD, meaning that either these secondary minerals were present at less than the XRD detection limit (approximately 0.5 wt%) and/or that they are poorly crystallized and therefore undetectable by XRD. Poorly crystalline secondary minerals were established to form on weathered feldspar surfaces (Carroll and Knauss [2005](#page-16-0); Casey et al. [1989;](#page-16-0) Hellmann et al. [2003](#page-17-0); Nugent et al. [1998;](#page-17-0) Oberlin and Couty [1970](#page-17-0); Zhang and Lüttge [2009,](#page-19-0) and references therein). The SEM images of weathered hemo-ilmenite grains (Fig. 6) also suggest that poorly crystallized secondary minerals containing Al, Mg, and Si were present. Another secondary mineral, which most probably precipitates in the Tio mine waste rocks, is illite $((K,H_3O)(A1,Mg,Fe)_2(Si,A1)_4O_{10}[(OH)_2,(H_2O)]),$ although it was not detected in thermodynamic simulations (because K was not analyzed in the leachates and therefore not considered), nor by XRD. However, K is released by Tio waste rocks and illite minerals most probably precipitate, as demonstrated in another study on Tio waste rocks (Plante [2010\)](#page-18-0). Moreover, K-bearing secondary minerals were observed in polished sections of waste rock samples from the present study (Plante et al. [2010b](#page-18-0)).

The higher Si release rates in weathered waste rocks may be partly attributable to the dissolution of poorly crystallized amorphous Si-bearing secondary minerals that

Fig. 7 Comparison of Ca–Si, Al–Si, and S-Ni cumulative values in humidity cells

could have formed on the surfaces either in the field or during kinetic testing. To illustrate the dissolution of such secondary phases, Eqs. 10 to 12 depict kaolinite dissolution. According to Cama et al. ([2002\)](#page-16-0), kaolinite dissolution in acid is expressed by Eq. 10. In near-neutral conditions, each Al^{3+} ion produced is hydrolyzed to $Al(OH)_3$, generating $3H^+$ ions (Eq. 11). The combination of kaolinite dissolution and $AI(OH)$ ₃ precipitation can be expressed as a neutral process (Eq. 12), as it neither produces nor consumes acid at near-neutral conditions (with Al^{3+} hydrolysis).

$$
Al_2Si_2O_5(OH)_4 + 6H^+ \rightarrow 2Al^{3+} + 2H_4SiO_4 + H_2O
$$
\n(10)

 $Al^{3+} + 3H_2O \rightarrow Al(OH)_3 + 3H^+$ (11)

$$
Al_2Si_2O_5(OH)_4 + 5H_2O \rightarrow 2Al(OH)_3 + 2H_4SiO_4 \qquad (12)
$$

Therefore, the dissolution of Si-bearing secondary minerals like kaolinite could partially explain the higher Si release rates in weathered waste rock samples, in addition to incongruent dissolution of weathered plagioclase surfaces releasing more Si into solution than fresh ones and the increase of pyroxene reactivity. Plagioclase samples $(Ab_{90}An_{10}$, which is 90% albite and 10% anorthite, containing low Ca levels) naturally weathered in slightly acidic soils (pH 4.9–5.6) showed a largely amorphous Al enriched and Na depleted coating, and the depleted feldspar surfaces beneath the coatings were Al and Na depleted (Nugent et al. [1998\)](#page-17-0). It was also

demonstrated (Nugent et al. [1998\)](#page-17-0) that these coatings developed within the first few months of field weathering. Finally, the coatings were obvious by AFM (atomic force microscopy) and detectable by SIMS (secondary ion mass spectrometry) and XPS (x-ray photoelectron spectroscopy) techniques, but largely undetected by SEM. Thus, it is possible that the plagioclases in the weathered Tio mine waste rocks (and after weathering of fresh waste rocks in the kinetic tests) were coated by an amorphous Al-bearing coating undetected by SEM and XRD in the present study.

Long Term Assessment of AMD Generation

The main neutralizing minerals in the Tio mine waste rocks are plagioclase (labradorite; 16–45 wt% in Table [3\)](#page-6-0) and pyroxenes (enstatite/pigeonite; 9–27 wt% total in Table [3](#page-6-0)). The pyroxenes are known to dissolve faster than labradorite by about an order of magnitude (Brantley et al. [2003](#page-16-0); Kwong [1993](#page-17-0); Wilson [2004](#page-19-0)). However, labradorite is believed to be twice as effective in acid neutralization as pyroxenes when compared in static tests (Paktunc [1999a](#page-17-0); Jambor et al. [2002,](#page-17-0) [2007\)](#page-17-0). Hence, the reaction of pyroxenes to acid production could be increased by plagioclase depletion. The higher Mg release rates in weathered waste rock samples than those observed for fresh ones suggest that the pyroxene contribution to acid neutralization increases in the weathered waste rock (considering that Mg release from ilmenite is the same or lower in weathered samples). Ilmenite release is known to be related to Mg and

Mn release (e.g. Schroeder et al. [2002](#page-18-0); Grey et al. [2005](#page-16-0); Hodgkinson et al. [2008](#page-17-0); Nair et al. [2009\)](#page-17-0); the low Mn values (not reported in this study) obtained in the leachates from all waste rock samples do not enable an efficient discrimination of pyroxene and ilmenite dissolution (the main Mg sources in the waste rocks).

The long term assessment of AMD generation potential of mine wastes can be done using oxidation-neutralization curves (Villeneuve et al. [2003;](#page-18-0) Benzaazoua et al. [2004](#page-16-0); Villeneuve [2004\)](#page-18-0). These curves compare the evolution of the products of sulfide oxidation (dissolved S as sulfates in the present work) with those of acid neutralization (such as Ca and Mg from plagioclases and pyroxenes in the present study). The long term assessment of AMD generation is performed by extrapolating the oxidation-neutralization curves and by plotting the initial solid sample values for comparison; if the points plot on the oxidation side of the extrapolation, the material is likely AMD generating in the long term. This interpretation is based on the interdependent hypotheses that the ratio between the oxidation and neutralizing products stays linear, and that mineralogy will not affect the linearity of the long-term relation (e.g. depletion of a mineral, precipitation of a new secondary phase). It was demonstrated that these hypotheses were respected for the long term weathering of high neutralization potential mine tailings, for which the neutralization was mainly provided by carbonates (Benzaazoua et al. [2004\)](#page-16-0) and by silicates (Hakkou et al. [2008](#page-16-0)).

In the Tio mine waste rocks, the main neutralization product not suspected to be affected by secondary mineral precipitation is Ca; the Ca cumulative normalized loadings against S in humidity cell tests are shown in Fig. 8a. The curves show two phases separated by an inflection point: a lower slope value in the first days of testing followed by a steeper slope for the rest of the duration. Such an inflection point towards the neutralization products was observed in studies involving carbonate neutralization (Villeneuve et al. [2003;](#page-18-0) Villeneuve [2004](#page-18-0)), and is caused by (1) surface passivation of the sulfides and/or (2) an exaggeration of the leaching of neutralization products (Villeneuve et al. [2003](#page-18-0); Villeneuve [2004](#page-18-0)). Mineral surface passivation is the decrease in reactivity following secondary mineral precipitation; such passivation was measured on weathered pyrite by cyclic voltammetry (Cruz et al. [2001;](#page-16-0) Villeneuve et al. [2003](#page-18-0); Villeneuve [2004\)](#page-18-0). The surface passivation of the sulfides could explain the curve shapes obtained for the cumulative normalized S loadings shown in Fig. [5e](#page-8-0). Preferential Ca release is believed to take place on the fresh waste rocks samples but not in the weathered ones, because of the plagioclase dissolution behavior presented in the literature background. In fact, the slope difference before and after the inflection point is more pronounced in the fresh than in the weathered waste rock samples, suggesting that the inflection point is mainly due to preferential Ca leaching from the plagioclases in early weathering stages. The slope and \mathbb{R}^2 values of the linear regressions of the cumulative Ca vs. S curves are presented in Table [5](#page-13-0); the slopes of the fresh waste rock samples $(6.4–10.0)$ are significantly higher than those of the weathered ones (1.65–2.06), with satisfactory $R²$ values ranging between 0.991 and 0.999. These results are a consequence of the non-stoichiometric Ca release in early plagioclase weathering stages.

The extrapolation of the oxidation-neutralization curves and the points of the initial Ca and S values are plotted in Fig. [9](#page-13-0). The C1 and C2 initial Ca values plot very close to the extrapolated curves, while for C3, they plot as acid-generating (Fig. [9](#page-13-0)a); therefore the long-term AMD generation potential of the fresh waste rocks is uncertain to acid-generating. On the other hand, the weathered waste rocks Ca and S initial values (Fig. [9b](#page-13-0)) plot far from the extrapolations on the neutralization side, demonstrating that the waste rocks should not generate AMD in the long term. The preferential Ca release in fresh waste rocks, because of the plagioclase dissolution mechanism, explains these results. However, the interpretation hypotheses of these oxidation-neutralization curves are not respected for

Fig. 8 Oxidation-neutralization curves for humidity cells considering Ca (a) and Ca + Mg + Al + Si (b) vs. S

Table 5 Characteristics of oxidation-neutralization curves for humidity cells $(>70$ days)

Waste rock	cumul. norm Ca/S		cumul. norm $Ca + Mg + Al + Si/S$			
sample	Slope	R^2	Slope	R^2		
C1	10.0	0.991	13.8	0.993		
C ₂	6.4	0.994	9.55	0.994		
C ₃	7.0	0.999	10.0	0.998		
C ₄	1.94	0.996	7.34	0.990		
C ₅	2.06	0.998	6.51	0.990		
C ₆	1.65	0.995	7.56	0.975		

curves extrapolation for fresh (a) and weathered (b) waste rocks

the fresh waste rocks in the present study, as plagioclase dissolution behavior changes with time (preferential Ca and Al dissolution in the early weathering stages). Nevertheless, one can assume that the weathered waste rocks response to kinetic tests are more representative of the long term behavior needed to assess the long-term AMD generation potential. Therefore, only the weathered waste rock results should be considered in the long term assessment of AMD generation of the Tio mine waste rocks, which suggests that the waste rocks are not acid-generating. Using the oxidation-neutralization curves involving only Ca as the neutralizing products takes into account only the plagioclase contribution to overall neutralization, and possibly the contribution of other minor Ca-bearing minerals. Not taking the pyroxene contribution in the overall neutralization is therefore conservative as pyroxenes are the second most important neutralizing minerals in the Tio mine waste rocks (see Table [3\)](#page-6-0).

In addition to Ca released by the plagioclases in response to sulfide oxidation, the Mg, Al, and Si releases are also related to silicate dissolution. Therefore, all these elements could be considered as neutralization products in the oxidation-neutralization curve; the cumulative normalized loadings of $Ca + Mg + Al + Si$ vs. S are shown on Fig. [8](#page-12-0)b, while the slopes and \mathbb{R}^2 values are presented on Table 5. The comparison of the initial $Ca + M$ $g + Al + Si$ values of the solid samples is not relevant, because Mg, Al, and Si are associated with other nonneutralizing minerals. However, it is interesting to note that the slopes of the fresh and weathered waste rocks $Ca + Mg + Al + Si$ vs. S loadings are much more similar (between 7 and 14) than the slopes of only the Ca vs. S loadings (between 1 and 10). The similarities are explained by the fact that plagioclase in weathered waste rocks release more Si and Mg than the fresh ones, which release more Ca. Thus, the higher Mg and Si releases compensate for the lower Ca releases in weathered waste rocks and consequently, the slopes considering $Ca + Mg + Si + Al$ are similar. Other studies obtained oxidation-neutralization curve slopes from fresh and weathered tailings that were similar to each other, where the neutralization was provided mainly by carbonates (Benzaazoua et al. [2004](#page-16-0); Villeneuve [2004](#page-18-0); Villeneuve et al. [2003](#page-18-0), [2009](#page-18-0)).

These results have important implications for long-term extrapolations of humidity cell test results. First, it means that even though the main neutralizing minerals react differently after 25 years of natural weathering (regarding Ca, Mg, Al, and Si release), the response of the fresh waste rocks to humidity cell tests are very similar to those of the weathered waste rocks when considering all the elements related to silicate dissolution, including those implicated in secondary phase precipitation. Therefore, the hypotheses of the oxidation-neutralization curves are respected when silicate neutralization products (Ca, Mg, Al, and Si) are considered. This in turn implies that the precipitation of secondary Mg, Al, and Si minerals for the fresh and

weathered waste rocks do not affect the linearity of the oxidation-neutralization relation in the long term (at least for 25 years of weathering). Therefore, long-term extrapolation of humidity cell results can be done for the Tio mine waste rocks (and possibly for other waste rocks where neutralization is provided by plagioclase and pyroxenes), as was done by Benzaazoua et al. [\(2004](#page-16-0)) for mine tailings in which neutralization mainly came from carbonates. Even though the kinetic tests on the Tio mine waste rocks do not predict the onset of CND conditions as they occur in the field, the results show that long-term extrapolation of the kinetic tests results are acceptable when the hypotheses of the oxidation-neutralization curves are respected for 25 years of weathering.

Discussion

Since sulfides oxidize at similar rates in fresh and weathered Tio mine waste rocks (Table [4\)](#page-9-0), the Ni release rates of fresh and weathered waste rocks should be of the same order. However, the Ni release rates of the weathered waste rocks are about an order of magnitude greater than those of the fresh waste rocks (Table [4](#page-9-0)) in humidity cell tests. The exact difference between fresh and weathered Ni release rates is difficult to establish because the Ni levels in leachates are regularly close to or below the detection limit, and therefore linear regressions of the cumulative Ni releases vs. time renders poor determination coefficients $(R²)$ in most of the humidity cell tests and for the fresh samples in weathering cells (Table [4\)](#page-9-0). The relation between S and Ni releases are illustrated in Fig. [7](#page-11-0)d by comparing the S vs. Ni cumulative normalized loadings. The slopes of the cumulative normalized loadings of S vs. Ni of the fresh waste rocks tend to decrease and therefore to get closer to those of the weathered waste rocks with prolonged kinetic testing. Assuming that Ni is released from sulfide oxidation, two main hypotheses have to be considered in order to explain the Ni release rates differences between the fresh and altered waste rocks: (1) precipitation as secondary Ni phases and (2) Ni sorption within the waste rock materials.

Geochemical modeling using Vminteq (Online Supplementary Figure 1) show that no secondary Ni hydroxide $(Ni(OH)_{2(am)}$ and $Ni(OH)_{2(c)})$ is suspected to precipitate in the conditions of the kinetic tests. Geochemical simulations in JCHESS (Online Supplementary Figure 2) indicate that the secondary Ni silicate $Ni₂SiO₄$ species is suspected to precipitate at $pH > 8$. The pH values measured in the humidity cell test leachates from the fresh waste rocks tended to decrease from 8.5 at the beginning to below 8 after 250 days, while in the case of the weathered waste rocks, the pH was rarely over 8 (Fig. [2](#page-7-0)). Moreover, the formation of secondary Ni minerals generally tend to be kinetically slow (Gunsinger et al. [2006,](#page-16-0) from Alpers et al. [1994](#page-16-0) and Xue et al. [2001](#page-19-0)) and discrete minerals may be absent, even though they are thermodynamically stable. Therefore, secondary Ni minerals are most likely not responsible for the absence of Ni in the leachates of the kinetic tests in the present study.

Ni sorption is a phenomenon known to occur on Tio mine waste rock surfaces (Plante et al. [2010a\)](#page-18-0). Many of the minerals found in the Tio mine waste rocks (ilmenite, plagioclase, chlorite, mica, spinel) show Ni sorption capacities (Plante et al. [2010a\)](#page-18-0); plagioclase and ilmenite were demonstrated to be the main Ni sorption supports in the Tio mine waste rocks. Ni sorption is probably occurring on plagioclase surfaces via layered double hydroxides (LDH), as discussed earlier in the present paper. XPS measurements showed that the sorbed Ni was similar to $Ni(OH)₂$ on all these minerals (Biesinger et al. [2009;](#page-16-0) Plante et al. [2010a](#page-18-0)).

In addition to Ni sorption on primary minerals of the Tio mine waste rocks, some of the secondary minerals believed to form in the waste rocks are known to have Ni sorption capabilities. For example, kaolinite metal sorption is a well-documented phenomenon (e.g. Dadhich et al. [2003](#page-16-0); Gu and Evans [2008](#page-16-0); Gupta and Bhattacharyya [2008](#page-16-0); Nachtegaal and Sparks [2003;](#page-17-0) Yavuz et al. [2003\)](#page-19-0). Co (whose behavior is similar to Ni) uptake by kaolinite occurs in two stages (Thompson et al. [2000](#page-18-0)) during sorption experiments: a rapid initial uptake followed by slow uptake that continued even after thousands of hours, illustrating that equilibrium was difficult to achieve in the experiments. Consequently, these authors point out that equilibrium models probably offer only approximate simulation of kaolinite metal sorption and similar laboratory sorption systems, particularly when experiments are of much shorter duration. Ni uptake by kaolinite without reaching equilibrium within up to 7 months was also observed (Nachtegaal and Sparks [2003\)](#page-17-0), revealing the formation of a Ni–Al LDH at the kaolinite surface resistant to mild desorption agents (0.02 M NaCl at pH6, 0.1 M NaNO₃ at pH 6, and $HNO₃$ at pH 4). Illite, a secondary mineral also believed to be present in the weathered Tio mine waste rocks, is also known as a Ni-sorbent material (e.g. Bradbury and Baeyens [2009](#page-16-0); Cama et al. [2005](#page-16-0); Echeverría et al. [2003](#page-16-0); Gu and Evans [2007\)](#page-16-0); Ni is retained on illite surfaces by precipitation of Ni–Al LDH (Elzinga and Sparks [2001\)](#page-16-0), adsorbed as an outer-sphere complex (Elzinga and Sparks [2001](#page-16-0)) or inner-sphere complex (Ech-everría et al. [2003\)](#page-16-0). Therefore, illite must be considered as a potential Ni sink in the present kinetic test work. Ni sorption on weathered waste rock samples were shown not to reach equilibrium within 72 h, while equilibrium was attained on the fresh waste rocks; these behaviors may be explained by the hypothesis that secondary minerals play a significant role in Ni sorption of the weathered Tio mine waste rocks.

Iron secondary minerals likely to precipitate in CND conditions (such as goethite, ferrihydrite or amorphous Fe(OH)3, lepidocrocite, magnetite, and hematite; see Online Supplementary Figure 1) are known to play an important role in controlling the metal release in mine waters by sorption or co-precipitation (Alpers et al. [1994](#page-16-0); Heikkinen and Räisänen [2008;](#page-17-0) Jambor and Blowes [1998](#page-17-0)). Also, both circumneutral pH and the presence of sulfates in drainage waters have been found to increase metal uptake by secondary iron oxyhydroxides phases (Heikkinen and Räisänen [2008](#page-17-0); Swedlund and Webster [2001;](#page-18-0) Swedlund et al. [2003,](#page-18-0) [2009](#page-18-0)). Sequential extractions on naturally weathered Tio mine waste rocks and post-kinetic test samples show that a significant portion of Ni is associated with reducible phases, mainly composed of iron oxyhydroxides (Plante et al. [2010a](#page-18-0)) SEM observations on polished sections of the Tio mine waste rocks show abundant iron oxyhydroxides phases, particularly around weathered iron sulfide particles (Pepin [2009](#page-17-0); Plante [2010\)](#page-18-0).

The fact that the weathered samples release more Ni vs S than the fresh ones (Fig. [7](#page-11-0)d) suggest that their Ni uptake has reached a level close to its maximum uptake capacity, leaving more Ni in solution than in fresh waste rocks, and/ or that secondary Si and Al mineral precipitation and their subsequent Ni uptake is less significant in weathered waste rocks than fresh waste rocks, as suggested by geochemical simulations (Online Supplementary Figures 1, 2). Both these hypotheses are consistent with the general observation that weathered waste rocks release more Ni than fresh ones.

Some of the Tio mine waste rocks piles generate Nicontaminated neutral drainage in the field that may exceed the regulated levels. However, the humidity cell leachates show Ni concentrations way below the regulated levels, even for weathered waste rock samples. Consequently, humidity cell tests on the Tio mine waste rocks do not adequately predict the Ni levels that are observed on field test pads and in actual waste rock piles at the mine site (Bussière et al. 2008 ; Pepin 2009 ; Plante 2010 ;). These results highlight the difficulty of extrapolating humidity cell results to the field scale for CND-generating mine waste. Only taking humidity cell tests into account in the drainage quality prediction of the Tio mine waste rocks, without separate sorption studies and field observations and investigations, would have generated a false safe assessment for drainage quality at the Tio mine site. However, humidity cell tests do evaluate the waste rock geochemical behavior, which is critical in understanding the development of CND conditions at the field scale.

Conclusion

The main conclusions that can be drawn from this study are:

- The acid-neutralization capacities of the Tio mine waste rocks are provided mostly by plagioclase and pyroxene minerals in the waste rocks;
- Secondary minerals are ubiquitous in the weathered waste rocks and control the mobility of Al, Mg, and Si in drainage waters;
- Even though the main neutralizing minerals react differently after 25 years of natural weathering (regarding Ca, Mg, Al, and Si release), the response of the fresh waste rocks during humidity cell leaching are very similar to those of the weathered waste rocks when considering all the elements related to silicate dissolution, including those implicated in secondary phase precipitation;
- The long-term extrapolation of kinetic cell results to the long term can be done for the Tio mine waste rocks (and possibly for other waste rocks whose neutralization is provided by the neutralizing silicates plagioclase and pyroxenes), as was done by Benzaazoua et al. [\(2004](#page-16-0)) for mine tailings in which neutralization mainly came from carbonates;
- The Tio mine waste rocks would not be not acid generating in the long term, as determined by humidity cells on fresh and weathered waste rocks interpreted by conservative oxidation-neutralization curves;
- Even though sulfide oxidation occurs at similar rates in fresh and weathered waste rocks, Ni generation is greater in weathered waste rocks, as Ni sorption capacities of the weathered waste rocks are approaching saturation.

These results highlight the fact that metal mobility in near-neutral drainage is often driven by sorption phenomena that are not straightforward. Although humidity cells implicitly account for sorption phenomena in the overall geochemistry of the system, the present study demonstrates that separate sorption studies are needed to adequately interpret the humidity cell results for the Tio mine waste rocks. The present study also highlights the limits of humidity cell tests in predicting field-scale metal loadings: the Ni levels remain well below the observed levels on the field. More insight is needed in order to extrapolate lab results to the field scale.

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