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From highly polluted Zn-rich acid mine drainage to non-metallic waters: Implementation of a multi-step alkaline passive treatment system to remediate metal pollution

Francisco Macías ^{a,*}, Manuel A. Caraballo ^{a,b}, Tobias S. Rötting ^c, Rafael Pérez-López ^{a,d}, José Miguel Nieto ^a, Carlos Ayora ^d

- ^a Geology Department, University of Huelva, Campus "El Carmen", E-21071 Huelva, Spain
- ^b Department of Geosciences, Virginia Tech, Blacksburg, VA 24061, USA
- ^c Department of Geotechnical Engineering and Geosciences, Technical University of Catalonia UPC—Barcelona Tech, Jordi Girona 1-3, D2-006, E-08034 Barcelona, Spain
- ^d Institute of Environmental Assessment and Water Research, CSIC, Jordi Girona 18, E-08034 Barcelona, Spain

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ABSTRACT

Complete metal removal from highly-polluted acid mine drainage was attained by the use of a pilot multistep passive remediation system. The remediation strategy employed can conceptually be subdivided into a first section where the complete trivalent metal removal was achieved by the employment of a previously tested limestone-based passive remediation technology followed by the use of a novel reactive substrate (caustic magnesia powder dispersed in a wood shavings matrix) obtaining a total divalent metal precipitation. This MgO-step was capable to abate high concentrations of Zn together with Mn, Cd, Co and Ni below the recommended limits for drinking waters. A reactive transport model anticipates that 1 $\rm m^3$ of MgO-DAS (1 $\rm m$ thick×1 $\rm m^2$ section) would be able to treat a flow of 0.5 L/min of a highly acidic water (total acidity of 788 $\rm mg/L$ CaCO₃) for more than 3 years.

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1. Introduction

Acid mine drainage (AMD) is a major cause of water pollution arising from poor management of mine activities, and it can be considered as the principal source of inorganic water pollution in mining districts (Younger et al., 2002). AMD polluted waters typically show low pH values, high acidity and high dissolved metal contents. The Iberian Pyrite Belt (IPB) can be considered one of the largest massive sulfide districts in the world. This region although mined since 3000 BC (Nocete et al., 2005), experimented its most extensive and intense mining period during the 19th and 20th centuries with more than 100 active mines (Sáez et al., 1999). These abandoned and uncontrolled mines have left an enormous amount of pollution sources discharging AMD into the surrounding basins (Ofias et al., 2004; Sarmiento et al., 2009a; 2011), particularly in the Odiel basin, where 427 km of its rivers are affected by AMD (Sarmiento et al., 2009b).

Iron and Al are two of the main contaminants present in the highly metal-polluted AMD typically found in the IPB, discharging to the Atlantic Ocean an average load of 7900 t/year Fe and 5800 t/year Al (Olías et al., 2006). Iron and Al present strong buffering systems in

these waters (Sanchez España and Trevor, 2007), and both systems play a key role in the natural attenuation of AMD (Bigham and Nordstrom, 2000), and therefore, they must be taken into account in remediation attempts. Another important AMD constituent in the IPB is Zn which has an estimated discharge to the Atlantic Ocean of 3500 t/year, an almost 15% of global gross flux of continental Zn to the oceans (Olías et al., 2006; Sarmiento et al., 2009b). Also other divalent metals such as Mn, Cu, Cd, Co, Ni and Pb are present in these AMDs, all of them remain in solution until pH values of about 8.5 (Cortina et al., 2003), being this an important factor to be considered in remediation strategies.

Given this severe and widespread metal pollution, the IPB is regarded as a world-class example of AMD polluted region and as the perfect "field-laboratory" to develop and test remediation strategies applicable to a wide range of metal and metalloid pollutants, as well as other AMD polluted sites around the world where similar acid systems are present and efforts are made for its remediation (e.g. Strosnider and Nairn, 2010; Caraballo et al., 2010; Behum et al., 2011).

Furthermore, the European Water Framework Directive 2000/60/EC exhorts the countries belonging to the European Union to achieve good ecological and chemical quality of all the European water bodies by 2015. However, in the context of the Odiel basin, due to its specific characteristics related to an ancient and uncontrolled AMD pollution, the regional authorities propose 2021 and 2027 as new deadlines for

^{*} Corresponding author. Tel.: +34 95 921 9834; fax: +34 95 921 9810. E-mail address: francisco.macias@dgeo.uhu.es (F. Macías).

compliance with the Directive, depending on the severity of AMD pollution (BOIA, 2011).

AMD pollution sources in the Odiel basin, and in many other old mining districts in Europe and in the world, typically correspond to orphan sites. In addition, the lifetime of this pollution generation has been estimated to range between tens to hundreds and hundreds to thousands of years for coal or sulfide mining districts, respectively (Younger, 1997). Taking these considerations into account, passive treatment systems are proposed as the most feasible and sustainable remediation option from both economical and environmental points of view (PIRAMID-Consortium, 2003; Johnson and Hallberg, 2005).

Conventional passive treatment systems (e.g., anoxic limestone drains, reducing and alkalinity producing systems or limestone sand reactors) are based on limestone dissolution or a combination of limestone dissolution and sulfate reducing bacteria activity (Hedin et al., 1994; Jage et al., 2001; Watten et al., 2005). Their application, although successful in waters with low to intermediate metal concentrations (e.g., AMDs in coal mining districts), is highly discouraged in highly polluted AMDs because their high Fe and Al concentrations promote serious problems of clogging and loss of reactivity (Rötting, 2007). Moreover, the pH values reached by limestone dissolution are insufficient to precipitate divalent metals (Cortina et al., 2003) and the high Zn and Cu concentrations of these waters may be lethal for sulfate reducing bacteria (Cabrera et al., 2006).

During the last ten years, the investigations on remediation of highly metal-polluted AMD in the IPB have been focused on the use of a novel reactive substrate known as Dispersed Alkaline Substrate (DAS). The concept of DAS relies on a mixture of a fine-grained alkaline reagent (e.g., limestone sand or MgO powder) and an inert coarse matrix (e.g., wood shavings). Limestone-DAS is suitable for trivalent metal removal like Fe or Al (Caraballo et al., 2009a; 2011a, b), whereas MgO-DAS efficiently removes divalent metals like Zn, Mn, Cd, Co, and Ni (Rötting et al., 2006; Caraballo et al., 2009b). In addition, a Natural Fe-Oxidizing Lagoon (NFOL) was designed and tested in the field to oxidize and remove some of the abundant Fe prior to the limestone-DAS treatment (Macías et al., 2012).

The aim of the present study is to obtain for the first time at the IPB a complete metal removal from a highly divalent and trivalent metal polluted AMD. To this end the hydrochemistry and mineralogy of the system will be discussed, paying special attention in the process involved in divalent metal removal. The most innovative aspect of the work is the implementation on a pilot scale of an MgO-based passive treatment focused on retention of divalent metals as the last step of a more complex system capable of treating highly polluted AMD. Moreover, the MgO-based treatment of divalent metal pollution in surface waters would not only be a local case study applied to acid mine water, but also circum-neutral mine water discharges with high divalent metal concentrations (e.g. Zn) can occur in mining areas where ore sulfides are hosted in carbonate rocks, for instance, Mississippi Valley-type lead–zinc deposits.

2. Materials and methods

2.1. Field remediation site

The passive pilot plant constructed at Monte Romero abandoned mine complex is located in the northern part of the IPB (SW Iberian Peninsula) (Fig. 1A). The first step is a natural pretreatment (NFOL), constructed near the mine shaft where AMD groundwater emerges. The outflow of the NFOL is connected with the DAS passive treatment system by a pipe (Fig. 1B). A schematic view of the DAS system is shown in Fig. 1C. The AMD is gravity forced through a downflow reactor of 3 m³ in volume (T1) filled with limestone-DAS (80% (v/v) pine wood shavings and 20% (v/v) limestone sand) connected in series to two decantation ponds (D1, D2) of 6 m³ in volume each with aeration cascades. AMD then flows through an identical limestone-DAS reactor

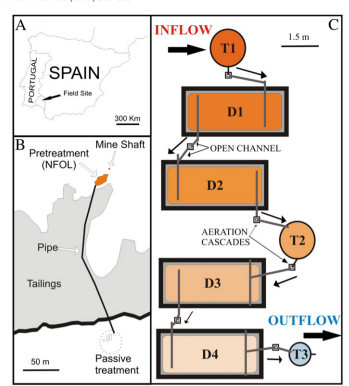


Fig. 1. General (A) and detailed (B) field site location, and schematic view of the DAS passive treatment system (C).

(T2) and two further aeration cascades and decantation ponds (D3, D4). Finally, AMD flows through a reactive tank of 1 m³ in volume (T3) filled with MgO-DAS reactive mixture (80% (v/v) pine wood shavings and 20% (v/v) MgO powder). This tank was equipped with lateral sampling ports at 1, 5, 10, 15, 20, 30, 40, 60 and 80 cm depth for water sampling and measurement of physicochemical parameters. The MgO-DAS tank outflow discharges into the Monte Romero creek.

High purity limestone sand (99.9% $CaCO_3$) was purchased at a local quarry (nominal grain size 3–6 mm). Pine wood shavings (curved flakes with a maximum length of about 1.5 cm) were purchased at a local carpentry. MgO powder was supplied by Magnesitas Navarras S.A. (Zubiri, NE-Spain). The bulk chemistry of the caustic magnesia type used in this study ("LIXIMAG") can be observed in Table 1, as well as the average grain size, which is typically less than 0.1 mm.

The chemical composition and physicochemical parameter range of the AMD at the mine shaft are shown in Table 2 (see Inflow). The mean flow rate emerging from the mine shaft and flowing through the NFOL pretreatment was 1.5 L/s. In order to adjust the inflow rate to the pilot scale of the treatment, the inflow at the first limestone-DAS tank was set to 1 L/min. The residence time (assuming a porosity of 50% in the reactive tanks) was 1.5 days for T1 and T2, 3 days for

Table 1Bulk composition and particle size distribution of the caustic magnesia used in the T3 MgO-DAS tank.
Source: Magnesitas de Navarra S.A., LIXIMAG.

Compounds (%)		Granulometry (9	6)	L.O.I. (%)		
MgO	65.34	>0.5 mm	0.0	14.6		
CaO	9.80	>0.2 mm	9.0	Bulk density (kg/L)		
SiO_2	3.09	0.2-0.1 mm	22.1	1.10		
Fe_2O_3	2.45	<0.1 mm	68.9			
Al_2O_3	0.38					
SO_3	4.14					

 Table 2

 Chemical composition and physicochemical parameters of the inflow (mine shaft) and outflow (discharge of T3) in the passive treatment of Monte Romero. Data obtained from five sampling campaigns.

Major element	ts (mg/L)		·					·
	Al	Ca	Cu	Fe	Mg	Mn	Si	Zn
Inflow	70-91	193-213	2.1-3.4	237-281	219-231	13-14	30-35	342-364
Outflow	< 0.2	199-767	< 0.005	< 0.2	200-488	< 0.2	< 0.2	< 0.05
FAO	5	n.r.	0.2	5	n.r.	0.2	n.r.	2
WHO	0.2	n.r.	2	0.3	n.r.	0.4	n.r.	3
Minor element	ts (µg/L)							
,	As	Cd	Со	Cr	Ni	Pb	Sr	Ti
Inflow	395–718	447-516	591-654	3.1-6.7	581-603	123-162	187-224	7–11
Outflow	<2	<1	<1	<1	<1	<1	182-445	<1
FAO	100	10	50	100	200	5000	n.r.	n.r.
WHO	10	3	n.r.	50	70	10	n.r.	n.r.
Physicochemic	al parameters							
	рН	Eh (mV)	Conductivity	(mS/cm)	Dissolved oxygen (m	ig/L) No	et acidity (mg/L)	T (°C)
Inflow	3.55-3.61	471-503	1.63-3.76		0.88-1.55	14	104-1609	18.8-20.7
Outflow	9.33-10.35	276-335	1.45-5.26		0.81-1.86	0		21.6-26.9

n.r. = no references; FAO = irrigation water recommendations of the Food and Agriculture Organization of the United Nations, and WHO = drinking water limits according to the World Health Organization. The symbol < followed by a number corresponds to samples where the analyzed value was below the detection limit of the analytical method for that element.

each decantation pond and 0.5 days for T3; i.e. a total of 15.5 days of residence time for the whole system.

2.2. Water and solid sampling

Water samples were taken every 15 days in the most representative points of the system: mine shaft, NFOL, inflow and outflow of the reactive tanks (T1-in, T1-out, T2-in, T2-out, T3-in and T3-out), and the outflows of the decantation ponds D1 and D3. Additional water samples were taken less frequently in the depth profile of T3 reactive tank. In all sampling campaigns the samples were filtered immediately after collection through 0.1 µm Millipore filters on Millipore syringe filter holders, acidified in the field to pH<1 with Suprapure HNO₃ and stored at 4 °C in sterile polypropylene containers until analysis.

Physical–chemical parameters were measured in the field using specific portable meters. The pH and redox potential were determined using a PH25 meter (Crison®) with Crison® electrodes, which was calibrated with 3 points (4.01, 7.00, 9.21) for pH and checked using 2 points (240 and 470 mV) for redox potential with Crison® standard solutions. Electrical conductivity and temperature were measured using a CM35 meter (Crison®) with 3 calibration points (147 and 1413 μ S/cm, and 12.88 mS/cm). Dissolved O2 was measured with an auto-calibrating Hanna® meter. Gross alkalinity was determined by CHEMetrics® Total Titrets®, with a range of 10–100 or 100–1000 mg/L as CaCO3 equivalents.

The drought season in the south of Spain promotes the cease of the AMD flow from the mine shaft at the end of the summer period. At this moment, a depth profile of T3 reactive tank was dug out to study the newly-formed precipitates developed within the tank. Solid samples were taken at 0–6, 6–12, 12–20, 20–30, 30–40, 40–50, 50–60, 60–70 and 70–80 cm depth. Samples were preserved in plastic bags at room temperature in the dark during its transport to laboratory, where they were dried at room temperature to prevent mineral transformation.

2.3. Analytical techniques

Analyses of water samples were carried out in the Central Research Services of the University of Huelva by Inductively Coupled Plasma Atomic Emission Spectrometry (ICP-AES, Jobin Ybon Ultima2). The method used was designed to estimate major, minor and trace elements in waters affected by AMD (Tyler et al., 2004). Multi-elemental standard solutions prepared from single certified standards supplied by SCP SCIENCE were used for calibration. Certified Reference Materials (SRM-1640 NIST freshwater-type and inter-laboratory standard IRMM-N3 wastewater test material, European Commission Institute for Reference Materials and Measurements) were also analyzed. Detection limits were: 200 μ g/L for Al, Fe, Mn, Mg and Si; 500 μ g/L for Ca; 50 μ g/L for Zn; 5 μ g/L for Cu; 2 μ g/L for As and 1 μ g/L for Cd, Co, Cr, Ni, Pb, Sr and Ti, and the analytical error was lower than 5%.

Solid samples used for chemical analysis and X-ray diffraction (XRD) were crushed and quartered to obtain representative sub-samples, which were ground to <0.1 mm in an agate mill. XRD patterns were obtained with a Bruker D5005 X-ray Diffractometer with Cu K α radiation. Diffractometer settings were 40 kV, 30 mA, a scan range of 3–65° 2 θ , 0.02 2 θ step size and 2.4-s counting time per step. To obtain the bulk chemistry of the T3 reactive tank precipitates, the sub-samples were digested with concentrated HNO₃. Metal concentrations of the different digestions were analyzed by ICP-AES.

Semi-quantitative chemical analysis and imaging of certain areas in the precipitates were obtained by the use of a JEOL JSM 5410 scanning electron microscope with an energy dispersive system (SEM-EDS). To gain a more accurate composition of some specific precipitates at the micro-scale, quantitative chemical analysis was performed with a JEOL JXA-8200 SuperProbe Electron Probe Micro-Analyzer (EPMA), using the fitted Wavelength-Dispersive Spectroscopy (WDS) equipment.

2.4. Reactive transport modeling

A 1-D reactive transport model of the T3 tank was created using the code RETRASO (Saaltink et al., 2004) to match the experimental observations on water chemistry and precipitates, corroborate the hypothesized geochemical processes, and extrapolate its future behavior. Hydrogeochemical speciation and saturation indices were calculated using PHREEQC (Parkhurst and Appelo, 1999). Thermodynamic data for aqueous, mineral and gas phases were taken from the WATEQ4F database (Ball and Nordstrom, 1991). Data for hydrozincite was added according to Schindler et al. (1969), while brucite solubility was updated according to Xiong (2008).

3. Results and discussion

3.1. Passive treatment system hydrochemistry

The general hydrochemical behavior of the complete passive system with respect to the main major elements (Fe, Al, Ca, and Mg), alkalinity and pH is shown in Fig. 2A.

An important Fe removal (165 mg/L) occurs between the mine shaft and T1-in due to the natural pretreatment in the NFOL. This high Fe removal can be attributed to different oxidation processes such as surface catalyzed oxidation of ferrous iron (SCOOFI; Jarvis and Younger, 2001; Younger et al., 2002) and biotic and abiotic oxidation and subsequent precipitation of schwertmannite (Macías et al., 2012). Additionally and in accordance to previous studies (Macías et al., 2009), 75% of the As present in the NFOL is co-precipitated with schwertmannite. A decrease in AMD pH is observed in the transit of the water along the NFOL (Fig. 2), moving from 3.6 in the mine shaft to 2.4 in T1-in. This pH tendency can be explained by the release of protons due to Fe hydrolysis and precipitation of schwertmannite. This decrease in pH although could be understood as detrimental for the water quality is in fact beneficial because it enhances calcite dissolution in the first limestone-DAS tank (T1).

In T1, due to limestone dissolution, an increase in water pH (from 2.4 to 6), gross alkalinity (from 0 to 190 mg/L as CaCO₃ equivalents) and Ca concentration (from 221 to 608 mg/L) takes place (Fig. 2A). These changes in the water chemistry induce a strong trivalent metal removal within the tank, as it is reflected in the decrease from 100 to 20 mg/L and 90 to 6 mg/L Fe and Al, respectively (Fig. 2A). The residual As not retained in the NFOL as well as all Cu, Pb and Cr present in the AMD are completely removed within the first limestone-DAS tank due to adsorption and/or co-precipitation

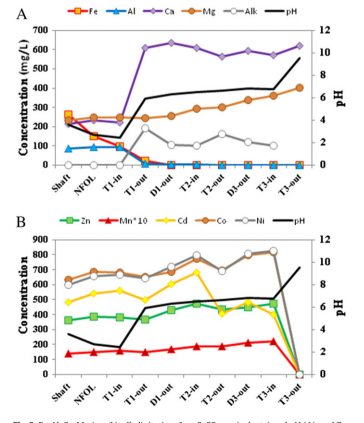


Fig. 2. Fe, Al, Ca, Mg (mg/L), alkalinity (mg/L as $CaCO_3$ equivalents) and pH (A), and Zn, Mn (mg/L), Cd, Co, Ni (μ g/L) and pH (B) distribution along the main representative points of the treatment.

with the Fe and Al precipitates. A detailed explanation of the aforementioned processes and the Fe and Al mineral phases systematically found in this step of the passive treatment system can be obtained in Caraballo et al. (2009a; 2011a).

In the following decantation ponds (D1 and D2), almost all the remaining Fe is oxidized and precipitated in the form of schwertmannite leading to some alkalinity consumption (Fig. 2A). A slight increase in water pH is also observed as a response of CO₂ degassing when the treated water contacts the atmosphere. In the second limestone-DAS tank and the third and fourth decantation ponds, the residual Fe and Al from the previous stage are now completely removed by the same processes mentioned for T1. Due to this preconditioning, the MgO-DAS tank (T3) treated an alkaline drainage with no trivalent metals and high concentration of divalent metals (Zn, Mn, Cd, Co and Ni). This absence of trivalent metals has been exposed as the best scenario to assure an optimal performance of the MgO-DAS (Caraballo et al., 2009b). As expected in the neutral pH conditions observed before T3, divalent metal concentration remained constant in the water with the exception of a slight increase in Zn concentration and a local Cd decrease within T2 (Fig. 2B). Zinc increase along the limestone sections of the system could be attributed to various factors: water evaporation in the ponds during the hot summer, high residence time of the system (inflow water does not have to match the water in the different sections of the treatment) and possible redissolution of Zn-particulate matter transported by the wind from the sulfide tailings in the proximities of the treatment. The process controlling Cd removal within T2 has been described as pH-dependent adsorption in previous studies (Caraballo et al., 2009b). As shown in Fig. 2B and Table 2, a complete Zn, Mn, Cd, Co and Ni removal was attained after T3. The hydrochemistry and mineralogy developed within the MgO-DAS reactive tank will be discussed in the next section.

In order to synthesize the enormous water quality improvement achieved in the different sections of the passive treatment, a modified Ficklin diagram has been employed. This diagram is commonly employed to classify AMD waters on the basis of pH and metal concentration (Ficklin et al., 1992; Ríos et al., 2008). The employed classification ranges from high acidity (HA) to moderate alkalinity (MAlk) and from extreme metallic (EM) to low metallic (LM). The main pollutants, based on their concentration and toxicity, present on the highly metal polluted waters of Monte Romero has been introduced in the modified Ficklin diagram (Fig. 3). As can be observed, the initial AMD belongs to the moderate acidity-extreme metallic class and after the pH decrease in the NFOL evolves to high acidity-extreme metallic. In the sections of the treatment dominated by limestone dissolution the AMD changes to near neutral-extreme metallic.

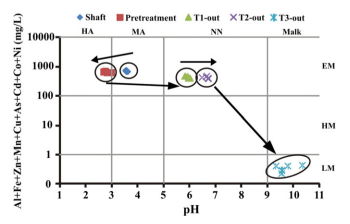


Fig. 3. Modified Ficklin diagram for the different steps of the treatment. HA: high acidity, MA: moderate acidity, NN: near neutral, MAlk: moderate alkalinity, EM: extreme metallic, HM: high metallic, LM: low metallic. Data obtained from five sampling campaigns.

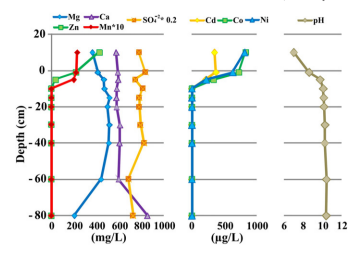


Fig. 4. Zn, Mg, Ca, Mn, SO_4 , Cd, Co, Ni and pH hydrochemical depth profile in the MgO-DAS reactive tank.

It is only after the MgO treatment when the water evolves to moderate alkalinity-low metallic. The high chemical quality obtained in the waters at the output of the system is offered in Table 2. As shown, when water chemistry after the treatment is compared with the limit values established for metal and toxic element concentration in irrigation (Ayers and Westcot, 1994) and drinking waters (WHO, 2011), all the analyzed elements easily fulfill these recommendations.

3.2. Detailed water chemistry and mineralogy of the MgO-DAS reactive tank

The inflowing drainage in T3 reactive tank showed a pH of 6.5 (Figs. 2 and 4), and after MgO powder dissolution this pH was increased to 10 in the first 10 cm of the tank profile, maintaining this value to the bottom of the reactive substrate (Fig. 4).

Slightly under the surface of the reactive material (1 cm depth), the pH value is increased to 8.6, promoting a concomitant decrease in Zn, Co and Ni concentrations by 200 mg/L, 100 μ g/L and 180 μ g/L, respectively. Manganese and Cd are not retained on the surface of the substrate. The pH value at 5 cm depth is 9.7, leading to a reduction in Zn concentration to a value of 40 mg/L from the initial 430 mg/L in the inflowing drainage, as well as for Co and Ni that are reduced from 850 and 800 μ g/L to 330 and 220 μ g/L, respectively. However, Mn and Cd concentrations do not show an important decrease. Finally, at 10 cm depth the pH is stabilized close to a value of 10, promoting that Zn, Mn, Cd, Co and Ni are abated below detection levels (Fig. 4). This profile evolution of divalent metal concentrations is consistent with the pH-dependent variations of those metal concentrations proposed by Cortina et al. (2003).

Concerning the mineralogy of the neoformed phases within T3, Fig. 5 shows a detailed view of Zn-rich precipitates distribution in the form of massive aggregates surrounding the wood shavings and

Table 3Bulk chemistry (mg/g) of the MgO-DAS filling substrate in the T3 reactive tank after operation time

Depth (cm)	Zn	Mg	Ca	Mn	Fe	S	Ni	Co	Cd
0-6	212.51	61.90	38.97	11.01	6.49	4.02	0.49	0.50	0.21
6-12	0.98	275.11	24.61	1.12	11.42	3.25	0.14	0.01	n.d.
20-30	0.14	263.10	26.59	0.67	10.44	3.41	0.11	0.01	n.d.
30-40	0.11	270.36	26.82	0.69	10.62	3.11	0.10	0.02	n.d.
40-50	0.13	251.98	24.35	0.67	9.23	3.19	0.10	0.01	n.d.
50-60	0.17	259.20	26.34	0.64	10.32	3.68	0.11	0.02	n.d.
60-70	0.27	230.57	21.91	0.61	9.18	3.44	0.10	0.02	n.d.
70-80	0.26	244.12	24.62	0.62	9.10	3.64	0.11	0.01	n.d.
Initial	n.d.	276.41	45.81	0.74	11.18	10.97	0.12	0.02	n.d.

n.d.: not detected.

the unreacted grains of the original substrate, and filling the voids left by the vegetal cells. These SEM images correspond to some samples obtained in the first 6 cm of the MgO-DAS in T3. From 6 cm depth to the bottom, the tank essentially contains unreacted MgO-DAS as can clearly be deduced if its chemical composition is compared with the one corresponding to the initial substrate (Table 3). According to these observations, T3 lifetime with an efficient performance could have been maintained during a much longer time if the mine shaft had not dried out.

The XRD pattern of the original substrate shows magnesite, ferroan magnesite, periclase, dolomite and portlandite as the main minerals detected. Gypsum was the only crystalline neoformed phase detected in the first 6 cm of the reactive material. Although Zn and Mn are the main constituents, on the basis of the bulk chemistry (Table 3), no Zn and/or Mn minerals could be precisely detected by XRD. The poor crystallinity and/or the small size of the crystallites of some neoformed phases limit the use of conventional XRD on those environments. From 10 cm to the bottom of the tank, brucite was detected as the only neoformed mineral within the tank, due to both periclase hydration and brucite precipitation.

EPMA analyses of more than 40 areas of some selected Zn-rich precipitates show a mean value of 76.22% Zn, with minor contents of Si, S, Mn, Ni and Co (Table 4). The slight Fe amount detected is probably inherited from ferroan magnesite in the initial substrate due to the absence of this element in the inflowing water. The sum of the main constituents (named as total in Table 4) quantified by EPMA presents an average of 84.88%. The chemical composition of those Zn-rich precipitates is reliably close to the expected one for hydroxy-carbonates of Zn (hydrozincite type) or Zn-hydroxides. Rötting et al. (2008) also identified these neoformed mineral phases in a column experiment treating Zn- and Mn-rich solutions using MgO-DAS.

In addition to the data presented in this study, a synchrotron light-induced micro analysis (combined μ -XRD and μ -XRF) has been recently performed to these precipitates. Hydrozincite [Zn₅(CO₃)₂(OH)₆] and a minor amount of loseyite [(Mn,Zn)₇(CO₃)₂(OH)₁₀] have been the only phases reliably identified as sinks for Zn and Mn, as

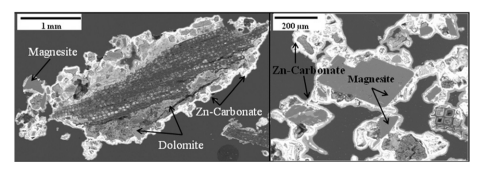


Fig. 5. SEM images of the Zn-rich precipitates developed within the MgO-DAS reactive tank.

Table 4

EPMA analysis (wt.%) of the neoformed Zn-rich phases in the first 6 cm of depth in the T3 reactive tank. A basic statistic (mean, maximum, minimum and standard deviation for 46 analysis points) is shown.

	ZnO (0.08)	SiO ₂ (0.03)	MgO (0.02)	SO ₃ (0.03)	MnO (0.08)	CaO (0.02)	NiO (0.08)	FeO (0.05)	CoO (0.08)	Total
Mean	76.22	3.72	2.51	0.92	0.60	0.48	0.17	0.16	0.14	84.88
Max	80.48	5.46	4.66	1,75	1.68	0.86	0.28	0.28	0.24	88.32
Min	68.82	1.51	1.43	0.60	0.26	0.36	0.13	0.10	0.10	79.48
Std. dev.	2.35	0.82	0.78	0.19	0.28	0.08	0.03	0.05	0.03	1.98

Detection limits are presented below each element.

well as for minor amounts of Co and Ni (Pérez-López et al., 2011). Even though no Mn³+,4+-rich mineral could be identified, oxidation of Mn²+ to Mn³+ at high pH (Morgan et al., 1967) and subsequent manganite (γ -MnOOH) cannot be discarded.

3.3. Chemical and mineralogical control on MgO-DAS performance and time evolution

In the light of the hydrochemical and mineralogical data acquired in this study, it was decided to obtain a reactive transport model to fully understand the different processes controlling the MgO-DAS performance and also to estimate the lifetime that the reactive substrate would have if the shaft had not dried out.

According to the calculations, brucite is slightly supersaturated below 10 cm depth in the tank (SI = 0.2-0.6), and pH is lightly higher than expected for brucite control. As Ca was observed to increase with depth in the T3 tank (Fig. 4) and lime (CaO) was detected as a mineral constituent of the initial substrate, it was hypothesized that some hydrated lime (Ca(OH)₂) could be dissolving within the substrate, even though equilibrium was not reached. Moreover, gypsum (another neoformed phase detected by XRD within T3) was also, to some extent, supersaturated throughout the tank, so that gypsum precipitation could explain the observed decrease of sulfate concentrations with depth (Fig. 4). Rötting et al. (2008) had also identified gypsum precipitated in laboratory MgO-DAS columns treating water with a similar composition as the T3 inflow. Therefore, the possibility of brucite dissolution/precipitation, hydrated lime dissolution and gypsum precipitation was incorporated into the reactive transport model (Table 5).

The kinetic rate law for the dissolution of brucite was derived from data of various authors summarized by Pokrovsky and Schott (2004). The dissolution rates shown in fig. 8 of their publication were approximated by the law:

$$r = 10^{-4} a_{H^{+}}^{0.45} \text{ mol} \cdot \text{m}^{-2} \cdot \text{s}^{-1}$$
 (1)

The amount m of mineral dissolved (mol/s) was calculated as:

$$m = A r(1\Omega) \tag{2}$$

where A is the reactive surface area (m^2/m^3 of porous medium) and Ω is the saturation of the solution.

A finite-element mesh with 410 linear elements and a total length of 80 cm was used to represent the depth profile of the T3 MgO-DAS

tank. Hydrozincite $(Zn_5(CO_3)_2(OH)_6)$ and β -Zn $(OH)_2$ were set as potential Zn-sinks, and manganite $(\gamma$ -MnOOH) as possible Mn-sink.

The autocatalytic oxidation of Mn²⁺ to Mn³⁺ was modeled after the kinetic law of Morgan et al. (1967), as cited in Diem and Stumm (1984):

$$\begin{split} \frac{-d \Big[M n^{+2}\Big]}{dt} &= k_1' \Big[O_{2(aq)}\Big] [OH^{-}]^2 \Big[M n^{2+}\Big] \\ &+ k_2' \Big[O_{2(aq)}\Big] [OH^{-}]^2 [M n O_x] \Big[M n^{2+}\Big] \end{split} \tag{3}$$

where $[Mn^{2+}]$, $[O_{2(aq)}]$ and $[OH^{-}]$ are dissolved concentrations (mol/L) of Mn²⁺, oxygen and OH⁻, respectively, $[MnO_x]$ is the concentration of Mn⁺³ reaction solid product (mol/L), k'_1 is the rate constant for homogeneous oxidation $(4 \cdot 10^{12} \text{ mol}^3 \text{ L}^{-3} \text{ day}^{-1})$ and k'_2 is the rate constant for heterogeneous oxidation $(10^{18} \text{ mol}^4 \text{ L}^{-4} \text{ day}^{-1})$.

All the phases were allowed to precipitate in equilibrium once saturation is reached. The input solution was set to be in equilibrium with atmospheric O_2 .

The values of the initial reactive surface areas of dissolving $Mg(OH)_2$ and $Ca(OH)_2$ were calibrated to match the measured depth profiles of Mg and Ca concentrations, and that of gypsum was adjusted to reproduce sulfate profiles (Table 5).

Using these parameters, the rest of pore-water concentrations, pH and the dissolved and precipitated solids calculated with the model match well with observations along the depth profile of the tank (Fig. 6A). Dissolved Mg concentration steeply increases in the first 4 cm of the substrate due to Mg(OH)₂ dissolution. Calcium concentration increases more slowly due to Ca(OH)₂ dissolution, causing pH to increase above Mg(OH)₂ saturation below 4 cm depth. Therefore, dissolved Mg decreases below this depth due to Mg(OH)2 reprecipitation. The pH increase also causes Zn precipitation, as hydrozincite immediately at the substrate surface and as β -Zn(OH)₂ up to approximately 3 cm depth (Fig. 6B). Concentration of Mn²⁺ starts to decrease below the depth of the Zn-precipitation front, where pH is high enough to sufficiently accelerate Mn²⁺-oxidation. The resulting Mn³⁺ immediately precipitates as manganite (Fig. 6B). As manganite has a low solubility at both alkaline and circumneutral pH (Eary, 1999), it does not redissolve even though the Mg(OH)₂-dissolution front moves downwards with time. Due to Ca increase, gypsum precipitates along the depth profile of the tank, and dissolved sulfate decreases accordingly.

The numerical model was also used to extrapolate the "lifetime" of the MgO tank. Assuming uniform flow, the MgO-DAS in the tank

Table 5Kinetic-equilibrium reactions and initial reactive surface of the MgO-DAS used in the reactive transport model.

Mineral	Initial volume fraction (m ² of mineral per m ³ of porous medium)	Initial surface area (m² mineral per m³ of porous medium)	Reaction	log K	Type of reaction
Brucite	0.2	100,000	$Mg(OH)_2 + 2 H^+ \Leftrightarrow 2 Mg^{2+} + 2 H_2O$	17.2	Kinetic
Portlandite	0.012	2.00E + 08	$Ca(OH)_2 + 2 H^+ \Leftrightarrow 2 Ca^{2+} + 2 H_2O$	22.8	Kinetic
Gypsum	0	1.00E - 12	$CaSO_4 \cdot 2H_2O \Leftrightarrow Ca^{2+} + SO_4^{2-} + 2H_2O$	-4.58	Kinetic
Hydrozincite	0	10	$Zn_5(CO_3)_2(OH)_6 + 6 H^+ \Leftrightarrow 5 Zn^{2+} + 2 CO_3^{2-} + 6 H_2O$	9.69	Equilibrium
β -Zn(OH) ₂	0	10	β -Zn(OH) ₂ +2 H ⁺ \Leftrightarrow 2 Zn ²⁺ +2 H ₂ O	11.75	Equilibrium
Manganite	0	10	γ -MnOOH + 3 H ⁺ \Leftrightarrow Mn ³⁺ + 2 H ₂ O	-0.17	Equilibrium

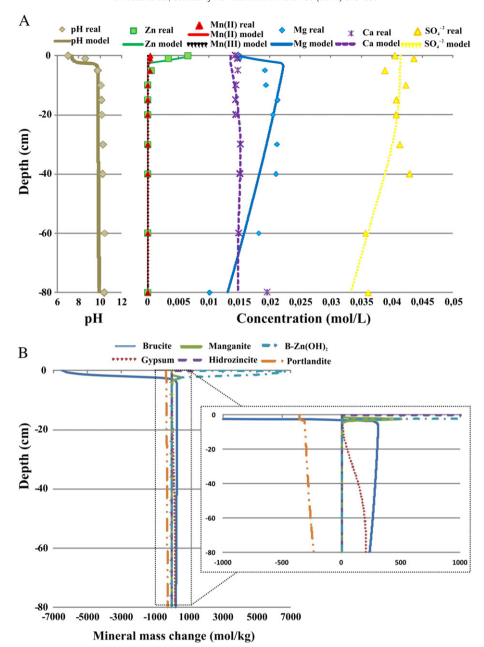


Fig. 6. Depth profiles for Zn, Mn, Mg, Ca, and SO_4 (mg/L) concentrations and pH (A), and mineral mass change (mol/kg) for brucite, gypsum, manganite, hydrozincite, β-Zn(OH)₂ and portlandite (B). Symbols indicate experimental data (real) and lines represent data obtained in the reactive transport modeling (model).

(1 m³) would have been able to treat the Monte Romero AMD at a flow rate of 0.5 L/min during approximately 3.4 years before Mn or Zn breakthrough at the tank outlet. However, this modeled hypothesis needs to be corroborated in longer field trials.

4. Conclusions

The application of the natural pretreatment NFOL to oxidize Fe, coupled with limestone-DAS for trivalent metal retention (Fe and Al mainly) and finally MgO-DAS for Zn, Mn, Cd, Co and Ni elimination, has shown to be capable to achieve the complete metal removal from a highly metal-polluted AMD.

The reactive transport model extrapolations indicate that the 1 $\,\mathrm{m}^3$ MgO-DAS tank is able to treat the Monte Romero AMD with a total acidity of 788 $\,\mathrm{mg/L}$ of $\,\mathrm{CaCO_3}$ at a flow rate of 0.5 L/min during approximately 3.4 years before Mn or Zn breakthrough the tank outlet.

The widespread use of these passive plants in specific acid discharges could improve in a few years the ecological and chemical quality of the Odiel basin. The pilot plant design tested in Monte Romero abandoned mine is being considered by the regional authorities for full scale application at this AMD discharge. The same NFOL+DAS scheme is proposed to treat other acid discharges in the Odiel basin.

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