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# Arsenic and metallic trace elements cycling in the surface water-groundwater-soil continuum down-gradient from a reclaimed mine area: Isotopic imprints

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## ABSTRACT

One decade after closure of the Salsigne mine (SW France), As contamination persisted in surface water, groundwater and soil near and down-gradient from the reclaimed ore processing site (OPS). We assess the fate of As and other associated chalcophilic MTEs, and their transport in the surface-water/groundwater/soil continuum down-gradient from the reclaimed OPS, using Sr-isotopic fingerprinting. The Sr-isotope ratio was used as a tracer of transfer processes in this hydro-geosystem and was combined to sequential extraction of soil samples to evaluate the impact of contaminated soil on the underlying phreatic groundwater. The contrast in Sr isotope compositions of the different soil fractions reflects several Sr sources in the soil. In the complex hydro-geosystem around the OPS, the transport of As and MTEs is affected by a succession of factors, such as (1) Existence of a reducing zone in the aquifer below the reclaimed OPS, where groundwater shows relatively high As and MTEs contents, (2) Groundwater discharge into the stream near the reclaimed OPS causing an increase in As and MTE concentrations in surface water; (3) Partial co-precipitation of As with Fe-oxyhydroxides, contributing to some attenuation of As contents in surface water; (4) Infiltration of contaminated stream water into the unconfined aquifer down-gradient from the reclaimed OPS; (5) Accumulation of As and MTEs in soil irrigated with contaminated stream- and groundwater; (6) Release of As and MTEs from labile soil fractions to underlying the groundwater.

## 1. Introduction

Tracking the geochemical cycles of metallic trace elements (MTEs) and arsenic in a post mining context is essential for a better orientation of environmental management, remediation strategy and neighborhood protection. Arsenic contamination is a major risk to human health and is a prominent environmental cause of cancer (Smith, 2002). Mining activities involving As ores have an impact on the environment and on human health that may persist for many decades after mine closure (Smedley and Kinniburgh, 2002).

High As concentrations are observed at numerous mining sites (Azcue et al., 1994; Azcue and Nriagu, 1995). In an As-Au mine, the

natural occurrence of As is associated with sulfide minerals, especially arsenopyrite (FeAsS), and realgar (AsS) (Cullen and Reimer, 1989; Matschullat, 2000; Moore et al., 1988; Smedley and Kinniburgh, 2002). In addition to mining effluents derived from water treatment, the oxidation of sulfides, such as FeAsS, is a common process, resulting in the release of As into the total environment (surface water, groundwater, soil and sediment) (Basu and Schreiber, 2013). Even after mine decommissioning, As sources remain in the form of tailings dams, dumps, or As-bearing ore stored on site.

Arsenic is among the most problematic and most studied metalloid elements in the environment because of its mobility over a wide range of redox conditions (Smedley and Kinniburgh, 2002). As (III) is the dominant form under reducing and acidic conditions, while As (V) is the prevalent form under oxidizing conditions. In soil, sediment and water, adsorption and desorption are the main processes controlling As transport and fate (Burton et al., 2009;

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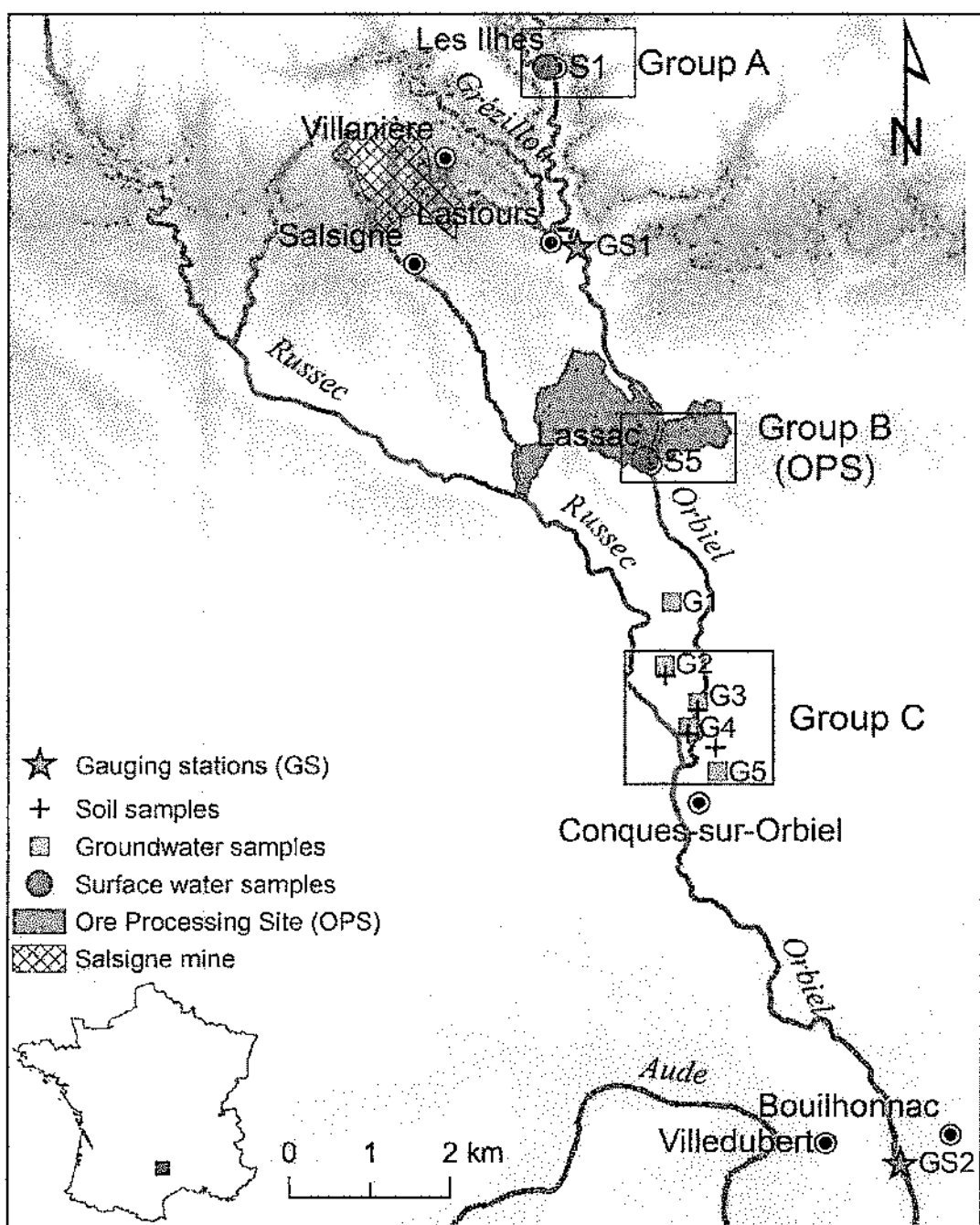


Fig. 1. Schematic map of the study area showing the three water groups, soil-sample and water-sample locations on a digital elevation model background.

streambed). These analyses were combined with isotopic fingerprinting of dissolved strontium ( $^{87}\text{Sr}/^{86}\text{Sr}$  ratio).

### 3.1. Water and soil sampling

First, groundwater was collected in the wet and dry seasons during one hydrological cycle in 2011 at five different sites (G1–G5) in the unconfined aquifer down-gradient from the reclaimed OPS. Following the preliminary investigation of groundwater and surface water made in 2011 (in wet and dry season) and owing to the interesting results, a monthly basis monitoring of surface water was decided for 2012.

A total of 24 surface water samples were collected at two sites (S1 and S5) for a monthly surface-water survey during a full year from January 2012 to January 2013.

The dominant agricultural soils irrigated with contaminated Orbiel stream water were also sampled. Samples were collected with a stainless-steel scoop, sieved at 1 mm and stored at 4 °C until analysis. For each soil-sampling site, four sub-samples were collected from the surface horizon (0–20 cm). 500 g from each soil sample were pooled and vigorously homogenized to obtain a composite soil sample.

Flow data of the Orbiel stream were extracted from the national HYDRO database ([www.hydro.eaufrance.fr](http://www.hydro.eaufrance.fr)); its volumetric flow is monitored both upstream of the reclaimed OPS, at the Lastours







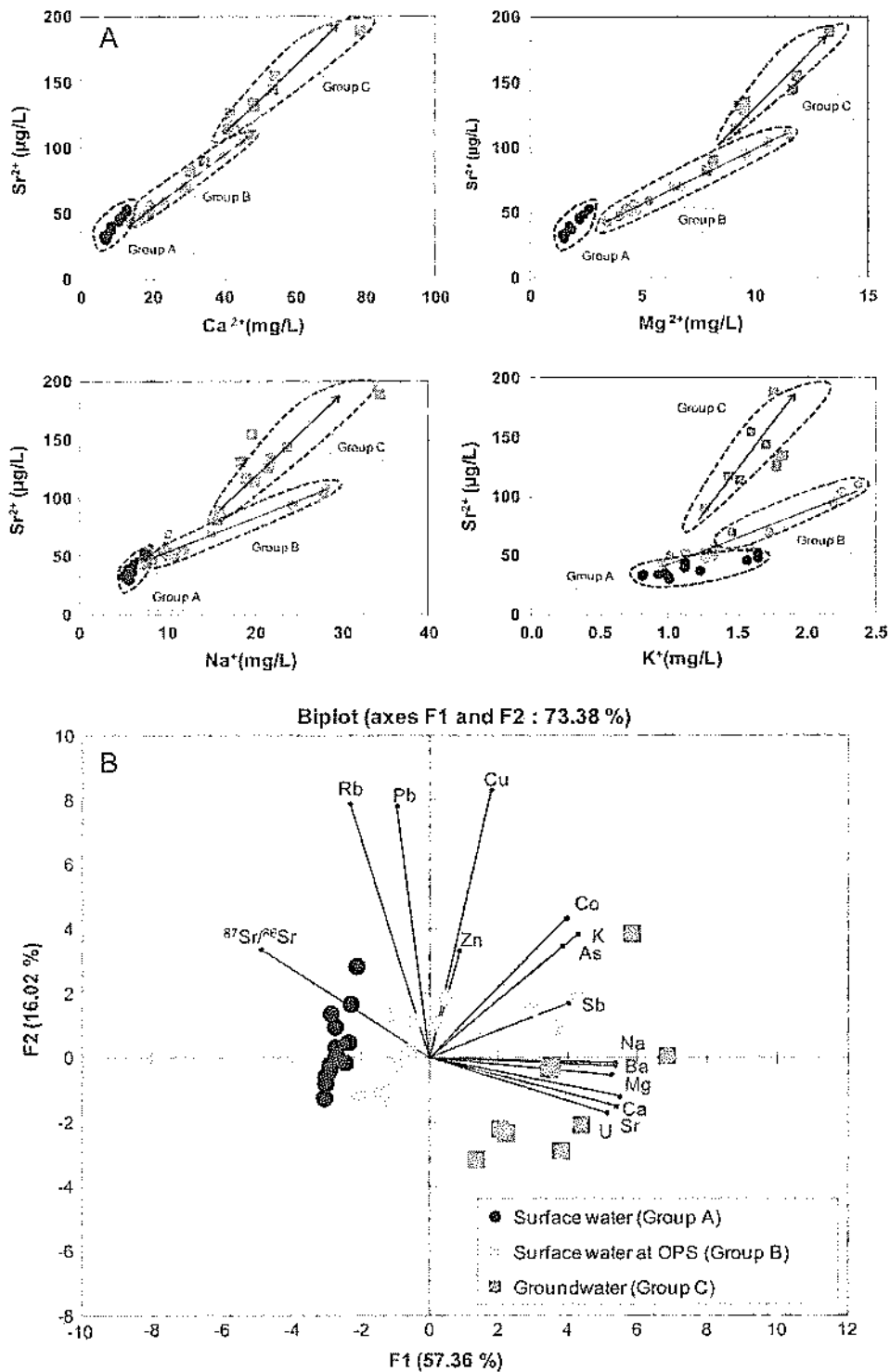


Fig. 2. (A) Binary diagrams of major cations vs. Sr showing the three water groups. (B) Results of the principal component analyses using 15 variables including major elements (Ca, Na, K, Mg), minor elements (Sr, Ba, Rb), trace elements (As, Co, Cu, Zn, Sb, Pb, U) and Sr isotopic ratios ( $^{87}\text{Sr}/^{86}\text{Sr}$ ).

4. Category IV consists of elements—including Zn, Ni and Co with <50% associated with the residual phase. The fractions associated with sulfide and oxides are important 35–50% respectively. In parallel, high As, Cu, Sr and Cd concentration were observed in the soil H<sub>2</sub>O-leachate, and in contrast, the MTEs concentrations showed low concentrations in the leachate.

#### 4.5. Environmental isotopes ( $^{87}\text{Sr}/^{86}\text{Sr}$ )

Large isotopic variations were observed in the  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios, decreasing from highly radiogenic values in surface water coming from contaminated groundwater near the reclaimed OPS (Group B) (0.71285–0.71328; Khaska et al., 2015), to much lower ratios of

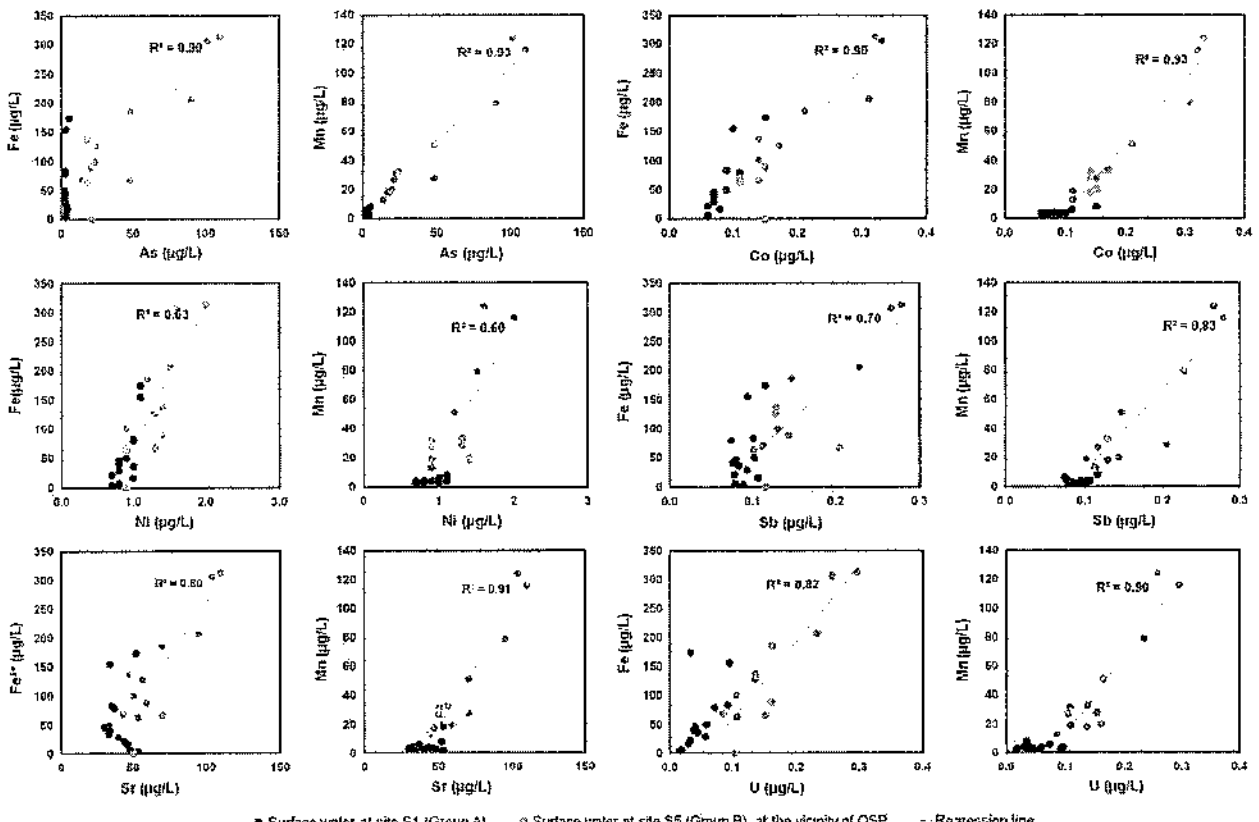


Fig. 3. Evolution of Fe, Mn, As, and MTE concentrations in surface water at the reclaimed OSP site (Group B) compared to incoming surface water, (Group A).

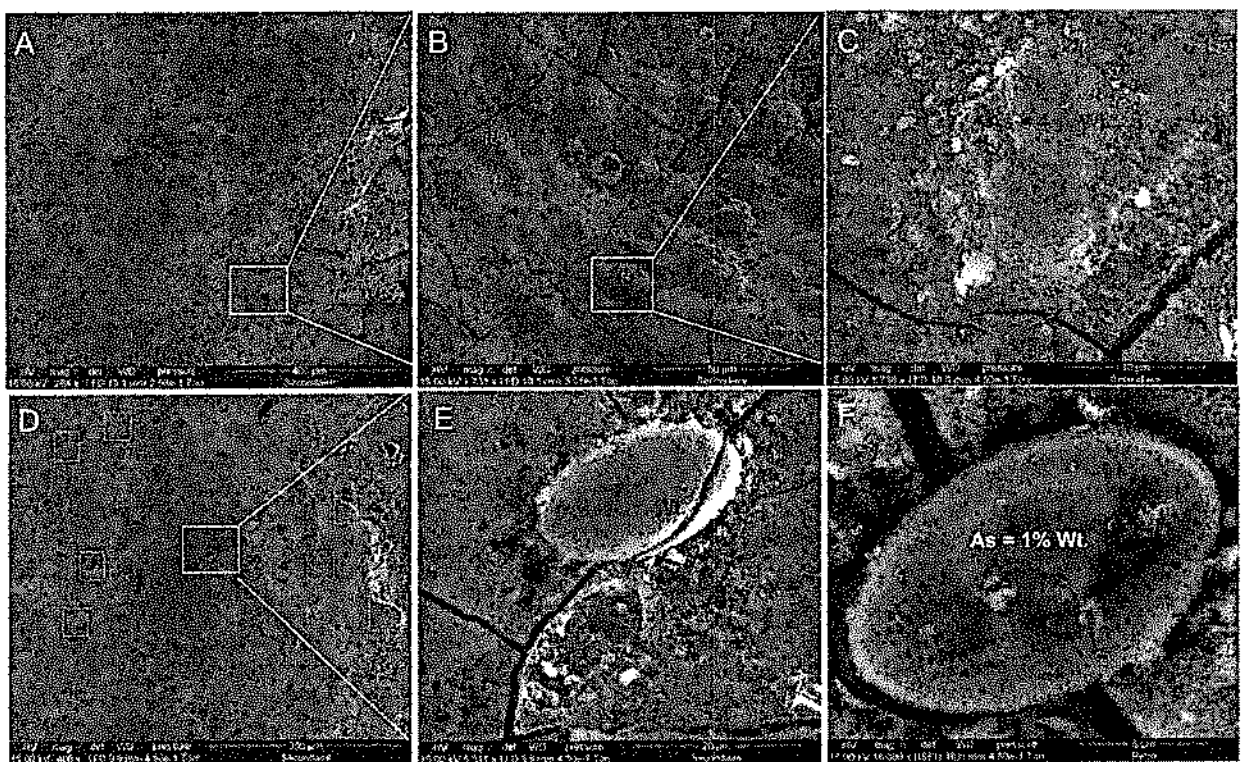


Fig. 4. A is a SEM image of Fe-oxyhydroxide precipitate coating the Orbel streambed, showing both types of As bearing Fe-oxyhydroxide precipitate. B, and C show a zoom on the abundant nodules on the streambed at site S5 with As content reaching 4.5 wt%. (Nodule is marked as "N"). E and F show a zoom on porous diatom silica shells bioadsorbing As bearing Fe-oxyhydroxide. Note the increase of arsenic content in uncoated diatom porous silica before (E) and after bioadsorption of As bearing Fe-oxyhydroxide (F).





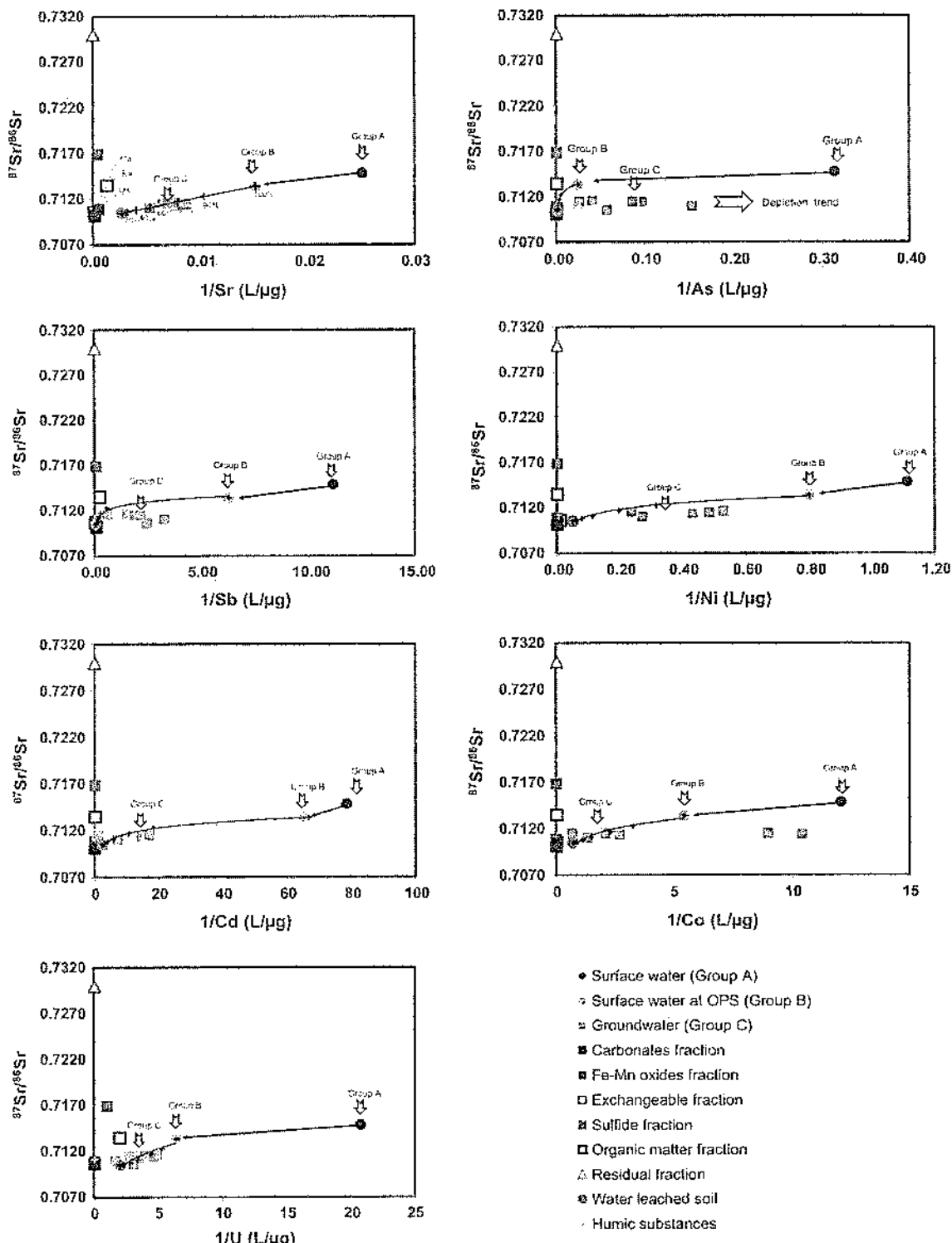


Fig. 7.  $^{87}\text{Sr}/^{86}\text{Sr}$  vs. 1/MTE and Sr (L/µg) mixing diagrams showing a decreasing trend of  $^{87}\text{Sr}/^{86}\text{Sr}$  between contaminated surface water at site S5 and labile alluvial soil. The black arrow illustrate the increase of As and MTEs at site S5 due to OPS influence, the black line represents the mixing between contaminated surface water and the labile soil fractions presented by the soil H<sub>2</sub>O leachate reflecting the equilibrium between solid labile fractions and the infiltrated water through soils.

B), with  $^{87}\text{Sr}/^{86}\text{Sr}$  ratios ranging from 0.7128 to 0.7140 (Table 1). The least radiogenic end-member has a  $^{87}\text{Sr}/^{86}\text{Sr}$  ratio similar to those of the autogenic fractions constituting the labile reservoir in the soil, thus suggesting an impact of contaminated soil on the

underlying unconfined groundwater. The soil H<sub>2</sub>O-leachate was considered as the least radiogenic end-member representing the equilibrium between infiltrated water through soil and labile soil fractions.



fractions, as the soil H<sub>2</sub>O-leachate, and the Orbiel stream water at site SS (Fig. 7), the mixing proportion of the soil labile fraction solution can be estimated to be between 25 and 40% in the unconfined groundwater. The calculated proportions of MTEs from the soil labile fraction are in the range of that calculated for Sr (40% for Ni, Cd and 60% for Cd and U) which may suggest a similar geochemical behavior during the transport of these MTEs from agricultural soil to underling groundwater.

However, to support the precision of proportion calculations, complementary experiments on soil-water kinetic equilibrium is required to better define the soil H<sub>2</sub>O leachate composition.

The geochemical processes controlling the As and MTEs release into groundwater are controlled by the interactions between the different soil constituents and water percolating in soil, such as irrigation water, effective precipitation or flood water.

The labile soil fraction as indicated by the isotopic approach includes:

1. A carbonate fraction through the dissolution process causing the release of metals sorbed and/or incorporated into the calcite crystal lattice;
2. An exchangeable fraction associated with clay minerals, organic matter or humic substances that may release MTEs through ionic exchange. It is well known that the adsorption/desorption process is pH dependent (e.g., Peters, 2008). High MTEs concentrations are usually associated with alkaline pH in water. Hence, the relatively high pH value observed in the soil promotes the release of As and MTEs into groundwater through effective precipitation (about 250 mm/y in the study area), infiltration of contaminated water during irrigation, and/or flooding events;
3. Iron oxides, from which MTEs releases is also controlled by adsorption/desorption. The high specific-surface area of iron oxides (up to 250–300 m<sup>2</sup>·g<sup>-1</sup>), and its high adsorption capacity (up to 55 g·kg<sup>-1</sup>) make it a privileged component for exchanging other MTEs with soil solution (Driehaus, 2002).

While, the mobilization of MTEs from sulfides can be due to the oxidation processes that release these elements in soil (Moore et al., 1988; Smedley et al., 1996). However, our approach shows that the dissolution of sulfides cannot explain the MTEs release from contaminated soil, as the <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the sulfide fraction are significantly higher than the required end-member. This observation is further consolidated by the reducing character of the studied soil. The observed shift in As concentrations, from the mixing curve (Fig. 7), may be explained by retention of As into the alluvial aquifer materials by oxidation and co-precipitation processes. This hypothesis is supported by the very low Fe and Mn concentrations resulting from the oxidizing conditions of the alluvial groundwater where As is adsorbed in such conditions. Moreover, the high redox potential of alluvial groundwater enhance As sorption (Gao et al., 2006). Opposite processes were observed by Hundal et al. (2013) where arsenic mobilization from alluvial soil was observed in relation to decreasing redox potential.

Also, based on the PHREEQC modeling results, the calculated MTEs total concentrations were plotted against the measured MTEs concentrations for comparison. The modelled concentrations fall globally within <16% of variation compared to of the observed data.

Based on these observations a conceptual model of the processes affecting As and MTE transport is constructed.

A conceptual model was developed integrating the results derived from the present study and that of Khaska et al. (2015) (Fig. 8). The hydrodynamic and hydrochemical processes affecting As and MTE transport in the studied hydrological system are summarized below:

- 1) A reducing zone exists in the aquifer near the reclaimed OPS, which is enriched in As- and MTEs via the reductive dissolution process;
- 2) Discharge of groundwater down-gradient of the reclaimed OPS, leading to an increase in As and MTE concentrations in surface water;
- 3) Precipitation of the Fe-oxyhydroxide, with only low partial precipitation ( $K_d = 0.36$ ) of dissolved As content from the surface water;
- 4) The contaminated surface water is then reintroduced into the unconfined aquifer farther downgradient of the reclaimed OPS;
- 5) Release of As and MTEs from the contaminated labile soil fractions linked to carbonate, Fe-Mn oxides and exchangeable sites.

## 6. Conclusions

We assessed the fate of As and other associated chalcophile MTEs and their transport mechanism in the surface-water/groundwater/soil continuum in a reclaimed mining context. Sr-isotope ratios used as tracers of As and MTEs transfer and their combination with sequential soil-extraction work, have proved that increasing As concentrations in surface water are explained by As released from unconfined groundwater associated with the occurrence of a reducing zone down-gradient of the reclaimed OPS. Such reducing conditions contribute only slightly to the immobilization of As in surface water, where the oxidation of Fe(II) to Fe(III) is confirmed by As-rich Fe-oxyhydroxide precipitate. The major part of As in surface water is reintroduced into the unconfined aquifer downstream of the contamination plume through surface-water infiltration. Study of the <sup>87</sup>Sr/<sup>86</sup>Sr ratios identified the soil fractions influencing the underlying groundwater. Moreover, our data show that the mobilization of As and MTEs in contaminated soil varies significantly as a function of the MTEs categories. It varies between 5% for Cr in Category 2, and 52% for As in Category 1. The labile fractions are mainly bound to carbonate, Fe-Mn oxides, and exchangeable and humic substance fractions.

This study, based on an original methodology, has strong implications in characterizing the cycling of MTEs and arsenic in a post-mining context, where pollution by these elements persists even after mine closure. Our study also has led to a better understanding of the key geochemical processes that occur during interactions between groundwater and surface water, and their implications on the fate of As and MTEs. For post-closure management and environmental issues, it is of outstanding importance to determine the origins of pollution (e.g. soils, sediments, tailing dams, runoff, streams, or groundwater) in the local hydrogeological context that was strongly modified during mine exploitation, the different pollutant transfer processes, and the impact of pollutant mobilization on the underlying unconfined aquifer especially when such groundwater is used for irrigation of agricultural products and private vegetable gardens.

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