# This is the final draft of the contribution published as:

**Pujades, E.**, Jurado, A., Orban, P., Ayora, C., Poulain, A., Goderniaux, P., Brouyère, S., Dassargues, A. (2018): Hydrochemical changes induced by underground pumped storage hydropower and their associated impacts *J. Hydrol.* **563**, 927 - 941

# The publisher's version is available at:

http://dx.doi.org/10.1016/j.jhydrol.2018.06.041

1	Hydrochemical changes induced by underground pumped storage hydropower
2	and their associated impacts
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#### 21 Abstract

Underground pumped storage hydropower (UPSH) using abandoned mines is an 22 alternative system to manage electricity production in flat regions. Water from an 23 24 underground reservoir is pumped to a surface reservoir to store electricity in the form of potential energy. Later, water is discharged through turbines into the underground 25 reservoir to produce electricity when demand increases. During this operation, the water 26 27 hydrochemistry continuously evolves. It varies in order to reach chemical equilibrium with the atmosphere (in the surface reservoir) and with the surrounding porous medium and 28 groundwater (in the underground reservoir). The hydrochemical variations may lead to 29 reactions in the reservoirs and in the surrounding porous medium, causing potentially 30 negative consequences for the environment and the system efficiency, especially when 31 pyrite is present in the surrounding porous medium. In this case, pyrite oxidation leads to 32 a decrease in pH and the precipitation of goethite or schwertmannite in the surface 33 reservoir. The decrease in pH is mitigated when calcite is present in the porous medium. 34 However, other concerns may arise, such as slight increases in pH, the precipitation of 35 ferrihydrite and calcite in the surface reservoir, and the oxidation of pyrite and dissolution 36 of calcite in the surrounding porous medium. Understanding the pH variations and the 37 precipitation/dissolution of minerals is of paramount importance in terms of the 38 environmental impacts and system efficiency. For this reason, with numerical modelling, 39 this work investigates the main hydrochemical changes and their associated 40 consequences when abandoned deep mines are used for UPSH. The main objective is 41 to highlight the importance of considering hydrochemical aspects when designing future 42 43 UPSH plants.

Keywords: Energy Storage System, Reactive Transport Modelling, Deep Mine, Pyrite,
Calcite.

## 46 **1. Introduction**

The main concern related to renewables energies, such as solar or wind energies, 47 is that the electricity production is highly variable and cannot be adapted to the demand 48 49 (Okazaki et al., 2015; Athari and Ardehali, 2016; Giordano et al., 2016; Hu et al., 2016; Mileva et al., 2016). This fact reduces their efficiency and limits their applicability. Energy 50 51 storage systems (ESSs) are a solution to manage the variable electricity production of 52 renewable sources and to increase the use of these sources (Gebretsadik et al., 2016). Such systems allow the storage of excess electricity generated during periods of low 53 energy demand and the production of electricity when the demand increases (Delfanti et 54 al., 2015; Mason, 2015). Pumped storage hydropower (PSH) is the most-used ESS 55 (EPRI, 2010). It allows for the storage and production of a large amount of electricity 56 (Steffen, 2012). PSH plants consist of two water reservoirs located at different elevations. 57 The excess electricity is used to pump water from the lower to the upper reservoir, thus 58 storing electricity in the form of potential energy. When the energy demand increases, 59 water is discharged from the upper to the lower reservoir through turbines to produce 60 electricity (Hadjipaschalis et al., 2009). Although PSH is widely used (Zhang et al., 2016), 61 such systems are constrained by topographic aspects because the reservoirs must be 62 located at different elevations, limiting the locations where plants can be constructed 63 (Mueller et al., 2015). In addition, PSH is controversial regarding its impacts on the 64 landscape, land use, environment (vegetation and wildlife) and society (relocations) 65 (Wong, 1996; Kucukali, 2014). Consequently, new types of ESSs have been evoked as 66 potential alternatives during the last years. One of them is the underground pumped 67 68 storage hydropower (UPSH) (Uddin and Asce, 2003), where the lower reservoir is underground while the upper one is located at the surface (Barnes and Levine, 2011). 69 UPSH has been considered and some projects have been planned in different parts of 70 the world. However, there are no bibliographical evidence of constructed UPSH plants. 71

For instance, a project was started in the 1980s in the Netherlands, but the plant was 72 eventually not installed for different reasons, such as unsuitable characteristics of the 73 underground area (Spriet, 2014). Similarly, Wong (1996) proposed the installation of 74 75 UPSH plants in Singapore using abandoned quarries as surface reservoirs and excavating the underground reservoirs. More recently, Severson (2011) suggested 76 77 potential sites to establish a taconite mine in Minnesota (USA) and to use the excavated cavity after the mining activities as the underground reservoir for a UPSH plant. UPSH 78 has also been investigated in Germany (Alvarado et al., 2016; Zillmann and Perau, 2015; 79 Luick et al., 2012), Belgium (Spriet, 2014), Spain (Menéndez et al., 2017) and South 80 Africa (Khan and Davidson, 2016; Winde et al., 2017). 81

The most important benefit of UPSH in comparison with PSH is that UPSH is not 82 limited by the topography and could even be used in flat regions. Consequently, more 83 possibilities exist to establish UPSH plants (Meyer, 2013). Impacts associated with 84 landscape, land use and society (relocations) are lower than those of PSH because at 85 least one of the reservoirs is located underground. Environmental impacts are different 86 than those produced by PSH and must be evaluated carefully. In addition, re-used 87 abandoned mines would provide added value to local communities after the cessation of 88 mining activities. 89

The main concern regarding the use of abandoned mines is that mine walls are 90 generally not waterproofed. Thus, UPSH plants interact with the surrounding porous 91 medium exchanging water. To date, this interaction has only been studied from a flow 92 point of view. Some studies have been focused on the impacts related to water exchange 93 94 (1) on the piezometric head field (Pujades et al., 2016; Poulain et al., 2016; Bodeux et al., 2017) and (2) on the efficiency of the plant (Pujades et al., 2017a). However, there are 95 no studies to date that have assessed the hydrochemical modifications induced by UPSH 96 and its associated impacts on the environment and the efficiency. Induced hydrochemical 97

modifications have not been reported to date given that this is an emergent EES. In fact, 98 as previously mentioned, there are no bibliographical evidence of operational UPSH 99 plants. However, the chemical particularities of mine waters and their behaviour have 100 101 been recognised as important challenges for the application of UPSH (EERA, 2016). In 102 addition, similarities between UPSH and managed aguifer recharge, in which occurred hydrochemical changes are well known (Johnson et al., 1998; Barker et al., 2016; 103 104 Antoniou et al., 2017), suggest that analogous changes may take place in UPSH systems. 105 Obviously, the hydrochemical changes would be different since water is also pumped in UPSH and it reaches the surrounding porous geological medium through an underground 106 107 cavity. For this reasons, it is meaningful to study the specific hydrochemical aspects associated to UPSH. 108

Under natural conditions, water in the underground reservoir and groundwater in 109 the surrounding porous medium are in chemical equilibrium with the porous materials. 110 Once the activity of the UPSH plants starts, water from the underground reservoir is 111 112 pumped, discharged and stored in the surface reservoir. During this operation, water is 113 aerated, and therefore, its chemical composition evolves to a new chemical equilibrium with the atmosphere. This is directly related to a variation in the dissolved O<sub>2</sub> and CO<sub>2</sub> 114 115 concentrations. When this water is subsequently discharged from the surface to the underground reservoir, it evolves again towards another chemical equilibrium with the 116 surrounding porous medium. This continuous evolution of the water chemistry may lead 117 to the precipitation and dissolution of minerals, and their associated impacts such as 118 119 variations in pH. For example, the oxidation of sulfides, which are common in coal-mined 120 environments (Younger et al., 2002), would result in groundwater with a very low pH (Bigham and Nordstrom, 2000). As a result, the UPSH activity would possibly affect the 121 surrounding groundwater quality. If part of the pumped water must be discharged in 122 123 surface water bodies because groundwater inflows progressively fill the underground

cavity (Pujades et al., 2016; Bodeux et al., 2017), the quality of those water bodies could 124 also be affected. These environmental issues are of paramount importance because 125 future UPSH plants should respect the Water Framework Directive (WFD,2000/60/EC) to 126 127 ensure a "good state" of water bodies, and especially groundwater, bodies. Hydrochemical changes are also relevant in terms of efficiency. Low pH values may 128 129 accelerate the corrosion of equipment (pipes, pumps, turbines, concrete structures) (Kapil 130 and Bhattacharyya; 2016; Sharma et al., 2011), whilst minerals precipitation may alter their mechanical efficiency (Sterpejkowicz-Wersocki, 2014). Thus, hydrochemical 131 changes may lead to a decrease of efficiency and durability of the facilities. Moreover, 132 133 periodical maintenance and cleaning tasks may be required.

For all these reasons, it is crucial to anticipate hydrochemical changes induced by 134 UPSH and their consequences on the environment and the efficiency of the system. This 135 issue is investigated in this work through numerical reactive transport models. The main 136 reactive transport processes associated with the groundwater-UPSH water interactions 137 138 are simulated considering three synthetic cases. The main objectives of this work are 1) to prove that UPSH activities can induce relevant hydrochemical changes, 2) to show that 139 these changes may entail negative consequences for the environment and the efficiency 140 141 of the system and 2) to highlight the importance of considering them in the design of future UPSH plants to ensure their feasibility. 142

143

#### 144 **2. Materials and methods**

145 2.1. Problem statement

The problem is sketched schematically in Figure 1. An underground reservoir 50 m by 50 m on 10 m of height, located in an unconfined aquifer is considered. The top and bottom of the underground reservoir are at a depth of 95 and 105 m, respectively. The surrounding porous medium is 200 m thick, and the external boundaries are located 500 m away from the underground reservoir. In natural conditions groundwater flows from the western to the eastern boundary. The head is considered to be at depths of 92.5 and 97.5 m on the west and east boundaries, respectively. Under natural conditions, the water table is located just at the top of the underground reservoir (95 m depth), which is therefore totally saturated.

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### 156 2.1.1. Chemistry of the porous medium

Abandoned coal mines are potential underground cavities to be used as 157 underground reservoirs for constructing UPSH plants. Sulfide minerals, whose oxidation 158 159 may entail important consequences for water chemistry, are common in coal deposits, and pyrite is usually the most common sulfide mineral in coal-mined environments (Akcil 160 and Koldas, 2006). Thus, Hypothesis 1 (H1) considers that the geological medium 161 contains 1% pyrite. The percentage of pyrite is not too much relevant since it is not totally 162 consumed during the simulation time. Coal mine deposits surrounded by carbonates or 163 164 containing lenses or intercalated layers of them have been worldwide reported (Sharma et al., 2013; Campaner et al., 2014; Xu et al., 2018) and carbonates counterbalance the 165 negative effects produced by pyrite oxidation. For this reason, Hypothesis 2 (H2) 166 167 considers that the geological medium contains 1% pyrite and 10% calcite. The percentage of calcite (carbonates) ensure that the neutralization acidic potential of the 168 surrounding porous medium allows counterbalancing totally the effects of pyrite oxidation 169 because the neutralization potential ratio is higher than 4 (Price et al., 1997). Hypotheses 170 1 and 2 represent the two boundary scenarios in which the neutralization capacity is null 171 172 (H1) or total (H2). Results of intermediate scenarios, in which the carbonate/pyrite ratio is lower than that considered in H2, would be comprehended by those obtained for 173 hypotheses H1 and H2. Finally, the presence of only calcite (10%) is considered in 174 175 Hypothesis 3 (H3) for two reasons. On the one hand, limestone mines are also considered

to be used as underground reservoirs for UPSH and these mines can be also 176 underground as shown the literature (Koch, 1987; Hustrulid et al., 2001). On the other 177 hand, the expected behaviour is very different to that occuring when pyrite is present. 178 179 Note that these percentages refer to the initial concentrations, and the conditions will evolve depending on the reactions that occur. It is assumed that the rest of the porous 180 181 medium is made up of silicates, whose reaction rates are very slow (White and Brantley, 182 1995) and can be neglected. Silicate dissolution and ion exchange is possible (Johnson 183 et al., 1999) and should be considered in some situations, as this has been proved in works developed in the field of aquifer managed recharge. However, the objective of this 184 185 work is to establish the main trends and highlight the importance of considering the hydrochemistry in the design of future UPSH plants. For this reason, only minerals (pyrite 186 and calcite) whose behavior is well known are selected. Obviously, the composition of 187 the geological medium in real cases is much more complex, and preliminary studies 188 performed before constructing UPSH plants must consider all the existing minerals and 189 190 all the processes.

191

### 192 2.1.2. Pumping and discharging frequencies

193 The actual pumping and discharging frequency cannot be forecasted. This depends on numerous factors such as the day, season, meteorology, electrical smart 194 grid, optimization scenarios or economic issues. Therefore, at this stage, representative 195 results are obtained assuming day/night cycles of 12 hours (i.e., water is pumped for 12 196 197 hours and discharged during the next 12 hours). The underground reservoir is considered 198 almost empty and almost full after each pumping/discharging phase. Thus, given the dimensions of the underground reservoir, pumping/discharging rates are chosen to be 199 43,000 m<sup>3</sup>/d. Note that the underground reservoir is not totally emptied at the end of each 200 201 cycle. A residual water depth of 1.4 m is intentionally left in the reservoir to keep nodes from which water is pumped and injected in a fully saturated zone. Appendix A shows the results of a sensitivity analysis in which the pumping/discharging rate is reduced by half.

#### 205 2.2. Basic concepts

The most relevant reactions that occur in the modelled hypotheses and their consequences are summarized in this section. Water aeration occurs each time that water is discharged and stored in the surface reservoir. This process induces  $CO_2$  and  $O_2$  exchanges between the water and the atmosphere. Thus,

210 
$$CO_2(aq) = CO_2(g)$$
 (R1)

211 
$$O_2(aq) = O_2(g)$$
 (R2)

The presence of pyrite is assumed in the H1 and H2 hypotheses. When the O<sub>2</sub> concentration in the water increases as a consequence of R2, pyrite is oxidized and ferrihydrite may precipitate as follows:

215 
$$FeS_2+15/4O_2(aq) + 7/2H_2O=Fe(OH)_3(s)+2SO_4^{2-}+4H^+$$
 (R3)

Ferrihydrite ( $\approx$ Fe(OH)<sub>3</sub>) precipitation induced by reaction R3 occurs if the water pH is higher than 6. Other minerals precipitate because of pyrite oxidation at low pH ranges. Goethite (FeOOH+3H<sup>+</sup>) may precipitate between pH 4 and pH 6, whilst schwertmannite (Fe<sub>8</sub>O<sub>8</sub>(OH)<sub>4.5</sub>(SO<sub>4</sub>)<sub>1.75</sub>) may precipitate at pH values lower than 4 (Sánchez-España et al., 2011). Goethite and schwertmannite precipitate as follows:

221 
$$Fe^{3+}+2H_2O = FeOOH+3H^+$$
 (R4)

222 
$$8Fe^{3+}+12.5H_2O+1.75SO_4^{2-}=Fe_8O_8(OH)_{4.5}(SO_4)_{1.75}+20.5H^+$$
 (R5)

The common characteristic of these three reactions is that they reduce the pH. These reactions are characteristic of acid mine drainage processes (Banks et al., 1997; Robb, 1994). Pyrite oxidation in the numerical model is calculated using the kinetic law described by Williamson and Rimstidt (1994) and considering a specific surface area of
 1000 m<sup>-1</sup>.

228 Calcite in the surrounding porous medium is considered for the H2 and H3 229 hypotheses. In this case, the following reaction occurs:

230

$$CaCO_3 + 2H^+ = Ca^{2+} + CO_2(aq)$$
(R6)

R6 does not occur in a unidirectional way, and if the pCO<sub>2</sub> is modified, calcite may 231 precipitate or be dissolved. For example, surface aeration of the pumped water induces 232 233 a reduction of pCO<sub>2</sub>, and, as a result, calcite should precipitate. The kinetics of calcite are implemented in the model using the kinetic law defined by Plummer et al. (1978) and 234 considering a specific surface area of 45.3 m<sup>-1</sup>. Thermodynamic data for aqueous 235 236 speciation and mineral and gas solubility are taken from the Wateg database, as implemented in the PHREEQC code (Parkhurst and Appelo, 1999). The solubility 237 constant of schwertmannite is taken from Sánchez-España et al. (2011). 238

Under the H1 hypothesis, the aeration of the water pumped into the surface 239 reservoir would increase the pO<sub>2</sub>. When this water is discharged into the underground 240 241 reservoir, it oxidizes pyrite and reduces the pH. In addition, when the water is stored in the surface reservoir, ferrihydrite, goethite or schwertmannite precipitate. In H2, pyrite 242 oxidation would induce the dissolution of calcite, which acts as a buffer, preventing a 243 244 decrease in pH. On the other hand, under the H2 and H3 hypotheses, the decrease in pCO<sub>2</sub> in the surface reservoir could produce calcite precipitation. Note that, UPSH plants 245 may act as CO<sub>2</sub> sources because pCO<sub>2</sub> increases each time that water is discharged into 246 the underground reservoir. Therefore, CO<sub>2</sub> emission is quantified under the H2 and H3 247 hypotheses. The influence of the specific surface area is assessed in a sensitivity analysis 248 249 whose results are shown in the Appendix A.

250

251 2.3. Numerical model

The problem was simulated using the code PHAST (Parkhurst et al., 1995; 252 Parkhurst and Kipp, 2002). This code solves multicomponent, reactive solute transport in 253 three-dimensional saturated groundwater flow (Parkhurst et al., 2010). Geochemical 254 255 reactions are solved by the code PHREEQC (Parkhurst, 1995; Parkhurst and Appelo, 1999). Flow and transport are computed with a modified version of HST3D (Kipp, 1987, 256 257 1997). It is possible to identify a line of symmetry crossing the domain from the west to 258 the east boundary located just in the centre of the considered problem crossing the 259 domain from the west to the east boundary (Figure 1 and Figure 2a). Therefore, the problem can be simplified by taking advantage of this symmetry, and only a half of the 260 261 domain is modelled (Figure 2a). This simplification does not affect the results and allows reducing the computation times. The modelled domain is divided into 15600 elements 262 refined in and around the underground reservoir (Figure 2b). The size of the elements 263 ranges from 2 to 100 m. 264

The flow and transport parameters are the same for the three hypotheses. For the 265 266 porous medium, the hydraulic conductivity (K) and storage coefficient (S) are chosen as 0.01 m/d and 0.05, respectively. The longitudinal ( $\alpha_L$ ) and transversal ( $\alpha_L$ ) dispersivities 267 are assumed to be 10 and 1 m, respectively. The underground reservoir (i.e., cavity) is 268 modelled by adopting a high value of  $K(10^6 \text{ m/d})$ , S of 1, and a dispersivity of  $10^4 \text{ m}$  in 269 the three directions. This value of dispersivity means that mixing of the discharged water 270 in the underground reservoir is rapid and homogeneous. This assumption is acceptable 271 because a large volume of water is discharged within a short period of time facilitating the 272 mixing. Note that the influence of the dispersivity used to simulate the underground 273 274 reservoir has been previously evaluated and is considered negligible (Pujades et al., 2017b). 275

276

277 2.3.1 Flow boundary conditions (BCs)

Dirichlet boundary conditions (BCs) are adopted at the western and eastern 278 boundaries. An initial hydraulic gradient of 0.005 is modelled by prescribing the 279 piezometric head at a depth of 92.5 and 97.5 m on the western and eastern boundaries, 280 281 respectively. Pumping and discharging rates of 21,500 m<sup>3</sup>/d are simulated by prescribing Neumann BCs in nodes located at the base of the underground reservoir. This rate allows 282 283 for the dewatering and filling of 8.6 m of the cavity in half a day. Note that the prescribed 284 flow rates are half those mentioned above because only half of the problem is modelled. Recharge through the upper and lower boundaries of the model is not considered and a 285 no-flow condition is assumed. 286

287

288 2.3.2 Hydrochemical conditions and transport boundary conditions

The initial hydrochemical conditions and transport boundary conditions are chosen 289 according to the hypotheses considered (H1, H2 or H3). Groundwater that 1) is initially 290 present in the whole modelled domain, (i.e., in the underground reservoir and in the 291 292 surrounding porous medium), 2) enters through the west boundary (during all the simulated time), and 3) initially flows out through the east domain, is in equilibrium with 293 pyrite in H1, with pyrite and calcite in H2 and with calcite in H3. In addition, it is prescribed 294 295 that groundwater under natural conditions has a  $CO_2$  partial pressure (p $CO_2$ ) of  $10^{-2}$  atm in H2 and H3. This value is chosen according to the literature (Faimon et al., 2012; Sanz 296 et al., 2011). Note that CO<sub>2</sub> is not considered under hypothesis H1 because it is not 297 relevant for the expected reactions. 298

299

300 2.3.3 Simulation strategy

The most challenging aspect of the numerical model is to simulate the evolving hydrochemistry of the discharged water during successive pumping/discharging cycles. The hydrochemistry of discharged water results in its aeration in the surface reservoir but

also depends on the chemical composition of the previously pumped water, which cannot
 be predicted in advance. To address this issue and to model the problem in a suitable
 manner, successive iterations progressively increasing the final simulated time are
 performed. The characteristics of the discharged water obtained from the previous
 iteration results are re-introduced at each iteration. The simulations are performed using
 the following steps:

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- 311

- A first iteration of 0.5 days is performed. The results of this iteration provide the hydrochemistry of water pumped to the surface reservoir from 0 to 0.5 days.
- 313 2. With the results of the first iteration, the second iteration from 0 to 1.5 days can be carried out. The input files of the model regarding the hydrochemistry 314 of the discharged water from 0.5 to 1 day are actualized using the 315 characteristics of the previously pumped water (from 0 to 0.5 day) and 316 considering two different scenarios (Sce1 and Sce2). In Sce1, it is assumed 317 that water in the surface reservoir reaches chemical equilibrium with the 318 319 atmospheric  $O_2$  and  $CO_2$ . Thus, the partial pressures of  $O_2$  (pO<sub>2</sub>) and pCO<sub>2</sub> of the stored water in the surface reservoir reach 10<sup>-0.7</sup> and 10<sup>-3.5</sup> bar, 320 321 respectively, before being discharged in the underground reservoir. In Sce2, it is considered that stored water in the surface reservoir does not reach 322 complete chemical equilibrium with the atmospheric O<sub>2</sub> and CO<sub>2</sub>. It is 323 assumed that the  $pO_2$  and  $pCO_2$  of the stored water in the surface reservoir 324 are 10<sup>-1</sup> and 10<sup>-2.29</sup> bar, respectively, before being discharged. These two 325 326 partial pressure values correspond to half of the O<sub>2</sub> and CO<sub>2</sub> concentrations if a chemical equilibrium with the atmosphere is fully achieved. These two 327 scenarios are helpful to evaluate the role of the pumping/discharging 328 329 frequency on the hydrochemical changes. In addition, it is assumed that the

precipitation of calcite, ferrihydrite, goethite and/or schwertmannite in the
 surface reservoir occurs at equilibrium. This second iteration (from 0 to 1.5
 day) provides the hydrochemistry of the pumped water from 1 to 1.5 day.

333 3. The same process as in the previous step is followed. Thus, using the 334 results of the previous iteration, the model is simulated from 0 to 2.5 days. 335 Now, the hydrochemistry of the pumped water from 1 to 1.5 days is used to 336 define the discharged water from 1.5 to 2 days. In this case, Sce1 and Sce2 337 are also considered.

338
 4. Subsequently, the fourth iteration from 0 to 3.5 days is performed using the
 characteristics of the pumped water from 2 to 2.5 day, on so on.

Note that each iteration is started at time 0, which increases the computation time. 340 Obviously, attempts are made to restart each simulation at the end of the previous one. 341 However, this was not possible due to limitations of the used code. In total, 30 days were 342 simulated because this methodology requires a long computation time (more than two 343 344 weeks are needed to simulate 30 days, and the computation time increases exponentially 345 with the simulated time). Nevertheless, 30 days are enough to reach the main objectives of evaluating the main hydrochemical reactions and their consequences in UPSH plants. 346 347 A simple code using Python (Van Rossum and Drake, 2011) was built to automate the process and facilitate the tasks. The code reads the results of each iteration, updates the 348 input files and initiates the subsequent iteration. 349

- 350
- 351 3. Results and discussion
- 352 3.1. pH evolution
- 353 3.1.1. Hypothesis 1 (only pyrite)

Figure 3a shows the calculated pH evolution in the surface and underground reservoirs and in the surrounding porous medium (at 5 and 15 m from the underground

reservoir in the downgradient direction). Figure 3a displays the numerical results for Sce1 356 (i.e., a chemical equilibrium with atmospheric O<sub>2</sub> is reached in the surface reservoir) on 357 the left, and for Sce2 (i.e., a chemical equilibrium with atmospheric O<sub>2</sub> is not totally 358 359 reached in the surface reservoir) on the right. In the case of Sce1, the pH values in the reservoirs decrease abruptly during the first cycles. Then, the decrease in pH becomes 360 361 more moderate. This behaviour is consistent with the expected reactions since the 362 increment of pO<sub>2</sub> in the surface reservoir promotes pyrite oxidation when the water is again discharged into the underground reservoir. The pH values are around 3 in the 363 underground and surface reservoirs at the end of the 30-day simulation time. The pH 364 365 values in the surrounding porous medium at 5 m from the underground reservoir also decrease as a consequence of groundwater exchanges and pyrite oxidation. The 366 decreasing trend in the pH values is similarly to that seen for the pH in the underground 367 reservoir. The decrease depends on the distance from the cavity and there is a delay for 368 larger distances. The pH decrease is less abrupt and lower at 15 m. At the end of the 369 370 simulation time, the pH values are 3.2 and 4.4 at 5 and 15 m, respectively. O<sub>2</sub> is totally consumed by pyrite oxidation each time that water is discharged. This means that water 371 aeration implies a continuous injection of O<sub>2</sub>. Consequently, the pH is expected to 372 373 decrease until pyrite is totally consumed. In the case of Sce2, the pH evolution is similar to that for Sce1, but the pH values are slightly higher. The pH values at the end of the 374 simulation are 3 in the underground reservoir, 3.2 in the surface reservoir, 3.3 at 5 m and 375 4.6 at 15 m from the underground reservoir. The computed pH distribution in the 376 surrounding porous medium after 5, 15 and 30 days is shown in Figure A1 of Appendix 377 378 Β.

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380 3.1.2. Hypothesis 2

Figure 3b shows the calculated pH evolution in the surface and underground 381 reservoirs and in the surrounding porous medium (5 and 15 m away from the underground 382 reservoir in the downgradient direction) for Sce1 (on the left) and Sce2 (on the right) under 383 384 hypothesis H2. In the case of Sce1, the pH values do not decrease like they do in H1 because calcite acts as a buffer. In fact, the pH increases instantaneously in the surface 385 386 reservoir because the CO<sub>2</sub> exsolves. Note that the instantaneous increase is not 387 observed in the figure because, prior to the first pumping, the surface reservoir is assumed empty. However, it does occur since the groundwater pH before the first 388 pumping is 7.3. The pH in the underground reservoir also increases during the early 389 390 cycles because this reservoir is filled with water from the surface one. After 2 days, however, the pH oscillates for each simulated cycle, and the pH decreases during 391 discharging and increases during pumping periods. Discharged water has higher  $O_2$ 392 concentrations that are rapidly consumed by pyrite oxidation, thus promoting a decrease 393 in the pH in the underground reservoir. When water is pumped, groundwater with higher 394 395 pH values from the porous medium enters the underground reservoir and increases the pH. The pH values in the surrounding porous medium also increase during the simulation. 396 The increase rate decay with time at 5 m and, at the end of the simulation, the pH is 397 398 quasi-stable. The pH behaviour is similar for Sce2 (Figure 3b on the right). In this case, the initial increase in pH and oscillations in the underground reservoir are less intense 399 because less CO<sub>2</sub> and O<sub>2</sub> are exchanged with the atmosphere. For both scenarios, the 400 pH increase at 15 m from the underground reservoir is not appreciable in the figure. The 401 402 distribution of the pH values in the surrounding porous medium after 5, 15 and 30 days is 403 shown in Figure A2 of Appendix B.

404

405 3.1.3. Hypothesis 3

Figure 3c shows the pH evolution in the surface and the underground reservoirs 406 and in the surrounding porous medium (at 5 and 15 m from the underground reservoir in 407 the downgradient direction) for Sce1 (on the left) and Sce2 (on the right) under hypothesis 408 409 H3. In the case of Sce1, the pH values do not decrease since this hypothesis does not 410 consider the presence of pyrite. In this case, pH increases instantaneously in the surface 411 reservoir and then remains constant. As in hypothesis H2, an instantaneous increase is 412 not observed in the figure because, prior to the first pumping, the surface reservoir is assumed emptied. However, it occurs since the groundwater pH before the first pumping 413 is 7.3. The pH value also increases in the underground reservoir because it is filled with 414 415 water from the surface reservoir. The increase is less extreme in the underground reservoir since the discharged water is mixed with the groundwater of the surrounding 416 porous medium. The pH also increases in the surrounding porous medium, but it depends 417 on the distance from the underground reservoir. The increase in pH is less as the distance 418 increases, and at a distance of 15 m from the underground reservoir, the observed pH 419 420 variation is negligible over the simulated time. The system behaves in a similar way when Sce2 is considered (Figure 3c on the right) but the maximum pH values are lower. If less 421 CO<sub>2</sub> is exchanged with the atmosphere in the surface reservoir, pH variations are smaller. 422 423 Oscillations in the pH value are observed in the underground reservoir. Each time that water is pumped, groundwater from the surrounding porous medium enters the 424 underground reservoir. Given that the pH in the surrounding porous medium is slightly 425 lower, groundwater inflows cause a decrease in the pH inside the underground reservoir. 426 427 Similarly, each time that the underground reservoir is filled, water with a low pCO<sub>2</sub> and 428 high pH enters the porous medium and mixes with the groundwater, which increases the pH values around the underground reservoir. The pH distribution in the surrounding 429 porous medium after 5, 15 and 30 days is shown in Figure A3 of Appendix B. 430

431

432 3.2. Precipitation, dissolution and oxidation of minerals

433 3.2.1. Hypothesis 1

Figure 4a shows the mass of goethite and schwertmannite precipitated in the 434 435 surface reservoir for Sce1 (on the left) and Sce2 (on the right) under hypothesis H1. Ferrihydrite does not precipitate because the pH is lower than 6, and goethite only 436 437 precipitates during the first cycles in which pH values range from 4 to 6. During the 438 subsequent cycles, the pH is lower than 4, and only schwertmannite precipitates. The results for Sce1 and Sce2 are slightly different. On the one hand, the amount of 439 precipitated goethite is lower in Sce1 because it only precipitates during two cycles whilst 440 441 goethite precipitates during the three first cycles in Sce2. On the other hand, more schwertmannite precipitates in Sce1 because more pyrite is dissolved in the surrounding 442 medium, and therefore, more minerals are available to precipitate. Minerals neither 443 precipitate in the porous medium nor in the underground reservoir. Figure 5a shows the 444 evolution of pyrite in the porous medium 5 m from the underground reservoir for Sce1 445 446 and Sce2 under hypothesis H1. The percentage of pyrite in the porous medium decreases for both scenarios because of its oxidation. However, the oxidation rate is higher in Sce1 447 because more O<sub>2</sub> is added to the system in the surface reservoir. 448

449

450 *3.2.2. Hypothesis 2* 

Figure 4b shows the precipitated mass of calcite and ferrihydrite in the surface reservoir for Sce1 (on the left) and Sce2 (on the right) under hypothesis H2. In this case, ferrihydrite precipitates instead of goethite and of schwertmannite because the pH is higher than 6. The precipitation rate of calcite is much higher during the early cycles than in the later ones. However, the precipitation rate of ferrihydrite is apparently constant, except during the initial cycles. The mass of precipitated calcite is higher than the mass

of ferrihydrite at the end of the simulations and the scenarios (Sce1 and Sce2) differ in
the total precipitated mass of both minerals, but not in the global trends.

Figure 5b displays the evolution of calcite and pyrite concentrations at a distance 459 460 of 5 m from the underground reservoir for Sce1 and Sce2 under hypothesis H2. Calcite and pyrite are continuously dissolved or oxidized in the surrounding porous medium. As 461 462 a result, their percentages in the surrounding porous medium decrease. The dissolved 463 and oxidized masses are higher for Sce1, especially for calcite. The dissolution and oxidation decrease significantly when the atmosphere-water equilibrium is not considered 464 (Sce2). In this case, the pO<sub>2</sub> in the discharged water is lower, and less pyrite and calcite 465 466 are oxidized and dissolved, respectively.

Note that the maximum precipitation rates of calcite are observed during early 467 cycles. At the beginning, all pumped water is equilibrated with the calcite of the porous 468 medium and has a pCO<sub>2</sub> of  $10^{-2}$ . As a result, a large decrease in the CO<sub>2</sub> in the surface 469 reservoir occurs, and the associated precipitations are high. In contrast, after some 470 471 cycles, the pCO<sub>2</sub>, and therefore, the mass of precipitated calcite in the surface reservoir, are lower because the discharged water (low pCO<sub>2</sub>) fills the underground reservoir and 472 only the water exchanges with the surrounding medium introduce new CO<sub>2</sub> in the pumped 473 474 water.

475

# 476 3.2.3. Hypothesis 3

Figure 4c displays the mass of precipitated calcite in the surface reservoir for Sce1 (on the left) and Sce2 (on the right) under hypothesis H3. The precipitation rate of calcite is higher during the early cycles than during the later ones. The decrease in the precipitation rate occurs during the first 5 days, and after, it is apparently constant. Obviously, if the exchange of CO<sub>2</sub> with the atmosphere is lower (Sce2), calcite precipitation decreases. When the discharged water is exchanged with the surrounding

porous medium and mixed with groundwater, its pCO<sub>2</sub> increases, and the pH decreases, 483 which promotes calcite dissolution. This fact is corroborated by the results. The pH values 484 in the surrounding porous medium are always lower than those in the surface and 485 486 underground reservoirs (Figure 3c), while the mass of calcite decreases in the surrounding porous medium (Figure 5c). Figure 5c displays the evolution of the calcite 487 percentage in the surrounding porous medium 5 m away from the underground reservoir. 488 489 Calcite is dissolved during all simulation times. The dissolution rate is higher at the beginning, and after 4-5 days, it decreases and becomes nearly constant. Because less 490 CO<sub>2</sub> is exchanged with the atmosphere for Sce2, the dissolved calcite is also lower. Note 491 492 that, as in the previous hypothesis, the maximum precipitation and dissolution rates are observed during early times. On the one hand, the variation of the pCO<sub>2</sub> in the surface 493 reservoir is higher at the beginning than at the later simulated times. On the other hand, 494 the contrast between the discharged water and groundwater is high during early cycles, 495 promoting the dissolution of calcite and its associated precipitation in the surface 496 497 reservoir.

498

499 3.3. Assessment of environmental impacts and efficiency

500 One of the objectives of the numerical modelling may be to inform about the environmental impacts identified in specific sites. Unfortunately, as mentioned above, 501 there are no bibliographical evidence of constructed UPSH plants. It is thus an emergent 502 technology, and therefore, environmental impacts have not been reported to date. In this 503 case, numerical modelling is used for predicting the potential environmental impacts of 504 505 UPSH, highlight their importance and raise awareness about the necessity of considering them for designing future UPSH plants. In addition, it allows deducing if UPSH related 506 activities may induce difficulties in the compliance of the Water Framework Directive 507 508 (WFD,2000/60/EC). This directive specifies that the states must prevent the deterioration

of groundwater and surface water bodies. The environmental objectives may be lowered 509 if the water bodies are affected by human activities with an important socio-economic 510 component, as it could be for UPSH plants. Even so, in these situations, the states must 511 512 guarantee the highest ecological and chemical status possible for surface water bodies and the least possible changes for groundwater bodies. The results show that UPSH 513 514 activity may alter the chemical properties of groundwater and surface water (when water 515 is discharged into surface water bodies), especially if pyrite is present in the geological 516 formations. For this reason, hydrochemical impacts must be assessed prior the construction of UPSH plants to ensure that the Water Framework Directive would still be 517 518 respected in the future. More specifically, in H1, pyrite oxidation leads to low pH values, and the pH increases in H2 and H3. In H2, calcite dissolution mitigates the pH decrease 519 induced by pyrite oxidation, and only a slightly increase in pH is observed when the  $pCO_2$ 520 decreases in the surface reservoir to achieve equilibrium with the atmosphere. This is the 521 same process that explains the pH increase observed for H3. The main difference 522 523 between H2 and H3 is that, after an early increase in pH, it decreases gradually for H2 while remaining constant for H3. pH variations occur also in both reservoirs, which is of 524 paramount importance in environmental terms because the excess stored water in the 525 526 surface reservoir may have to be discharged into surface water bodies, thus also modifying their pH values. 527

The porosity in the surrounding porous medium increases as a result of the dissolution of calcite and the oxidation of pyrite. The porosity increases  $2.2 \cdot 10^{-5}$  and  $1 \cdot 10^{-5}$ 5 % for Sce1 and Sce2 under hypothesis H1,  $1.4 \cdot 10^{-4}$  and  $4.2 \cdot 10^{-5} \%$  for Sce1 and Sce2 under hypothesis H2, and  $3.7 \cdot 10^{-5}$  and  $3.9 \cdot 10^{-6} \%$  for Sce1 and Sce2 under hypothesis H3, respectively. The porosity increases more for H2 because pyrite is oxidized and calcite is dissolved at the same time. Although the increase in porosity is not relevant during the simulated period, it should be considered for large activity periods. The

535 groundwater exchanges, and thus the potential environmental impacts, increase with 536 higher porosities (Pujades et al., 2016). In addition, if a pollutant was accidentally 537 discharged into the surface reservoir, it would reach faster the subsurface environment.

Table 1 summarizes the released  $CO_2$  (g/kWh) for both scenarios (Sce1 and Sce2) under hypotheses H2 and H3. Emissions are computed considering the total  $CO_2$ released during the total simulated period, and the volume of released  $CO_2$  once the emission rates are stabilized after some days of activity ( $\approx$ 10 days). The results show that UPSH plants may act as a  $CO_2$  source when carbonates are present in the surrounding porous medium. However, the emissions are much lower than those produced by other energy sources such as coal or gas (Schlömer et al., 2014).

The results are also relevant in terms of efficiency. Low pH values in the reservoirs 545 may lead to the corrosion of UPSH facilities, and the precipitation of minerals may induce 546 clogging events (e.g., in pipes). As a result, maintenance and cleaning tasks and the 547 adequate selection of materials are needed to avoid corrosion and clogging, to extend 548 549 the useful life and to maximize the efficiency of the facilities (pumps, turbines, pipes). The dissolution and oxidation of minerals in the surrounding porous medium are also relevant 550 in terms of energy production because the efficiency of pumps and turbines depends on 551 552 groundwater exchanges (Pujades et al., 2017a). Generally, their efficiency increases with greater water exchanges. The results show that the increase in porosity, and thus in 553 groundwater exchanges, is low over short periods. However, this increase should be 554 investigated because the useful life of future UPSH plants is expected to be longer. 555

556 When equilibrium between the atmosphere and water from the surface reservoir 557 in not completely reached, the behaviour of the system is similar. However, the pH 558 variations and the masses of minerals dissolved, oxidized and precipitated are lower. 559 Therefore, the comparison between Sce1 and Sce2 (for the three hypotheses) reveals

that impacts on the environment and system efficiency are smaller as the aeration during
 the discharge and storage of water in the surface reservoir is less important.

562

#### 563 **5. Conclusions and prospects**

Induced hydrochemical reactions result from the surface exposure of the pumped water in UPSH contexts. When water is discharged and stored in the surface reservoir, the concentrations of  $O_2$  and  $CO_2$  evolve to reach chemical equilibrium with the atmosphere. Thus,  $O_2$  tends to increase, and  $CO_2$  to decrease. In addition to the reactions occurring in the surface reservoir, when water is discharged into the underground reservoir, it interacts with the groundwater and the surrounding porous medium. This interaction also leads to chemical reactions.

The impacts are especially relevant if pyrite is present in the porous medium. The increase in the pO<sub>2</sub> induces pyrite oxidation. This reaction releases H<sup>+</sup>, decreasing the pH, which is relevant in terms of environmental impacts and efficiency of UPSH plants. Lower pH values may lead to the corrosion of the facilities (pipes, pumps and turbines). In addition, ferrihydrite, goethite or schwertmannite may precipitate in the surface reservoir requiring maintenance and cleaning tasks.

577 pH does not decrease when calcite is present in the porous medium. In contrast, 578 the pH slightly increases. However, other aspects must be regarded, such as the 579 precipitation of calcite and ferrihydrite in the surface reservoir. In addition, a small amount 580 of CO<sub>2</sub> is released to the atmosphere..

The results also show that UPSH may induce the dissolution/oxidation of minerals in the porous medium. This fact may be relevant in terms of efficiency and environmental impacts. The dissolution and/or oxidation of minerals promotes the increase of the transmissivity and/or the porosity. Consequently, water exchanges between the surrounding porous medium and the underground reservoir would increase.

A comparison of the numerical results obtained for Sce1 and Sce2 reveals that the hydrochemical changes induced by UPSH and their associated consequences decrease 1) when the water is stored in the surface reservoir over a shorter period and 2) when the aeration, occurred when pumped water is discharged and stored into the surface reservoir, is lower. Thus, the pumping/discharging frequency and the method to discharge and to store the pumped water in the surface reservoir determine the reactions and their associated impacts.

593 It was not possible to lengthen the simulation periods because of the long computation times required. Although an attempt was made to modify the simulation 594 595 strategy and start each iteration at the end of the previous one, it was ultimately not possible. However, the simulated period is sufficient to identify the main trends of the 596 system, the reactions that occur and their consequences. It is possible to affirm that after 597 a long active period of a UPSH plant, all the pyrite near the underground reservoir would 598 be oxidized. However, the pH would continue to decrease in the surrounding porous 599 600 medium because the discharged water with high concentrations of O<sub>2</sub> would reach downstream areas in which pyrite would still be present. The only difference would be 601 noted in the amounts of precipitated ferrihydrite, goethite and schwertmannite. The 602 603 products of the oxidized pyrite far from the underground reservoir could not be mobilized by the pumping. Similar predictions could be done for contexts in which calcite is present. 604

This work highlights the importance of considering hydrochemical aspects for the construction of future UPSH plants. In fact, preliminary studies focused on determining the hydrochemical changes and their consequences should be mandatory. The results of these preliminary studies combined with the environmental policies will allow taking decisions concerning the construction of UPSH plants using abandoned mines. In other words, these preliminary studies should be considered for selection of the potential abandoned mines to be used as underground reservoirs. Preliminary studies must consist

in a detailed hydraulic and hydrochemical characterization of the geological medium by 612 means of field tests (pumping and tracer tests), mineralogical analyses and laboratoty 613 tests. In addition, the main characteristics of the plant (volume of the underground 614 615 reservoir, flow-rates, pumping/discharging frequencies, etc) must be considered and the 616 aeration process in the surface reservoir must be investigated to precisely quantify the 617 gas exchanges. All these informations should be integrated into reactive transport models 618 for predicting the consequences of reactive transport processes induced by UPSH. 619 Monitoring will be also needed during the operational phase of UPSH plants to verify that hydrochemistry evolves as expected. In this context, hydrologists may play a specific role 620 during the selection, design and operational phases. Hydro(geo)logists should 1) 621 contribute in the characterization of the porous medium, 2) built the numerical models 622 and analyse the results, 3) design the monitoring strategies and monitor the system 623 evolution during the operational phase, and 4) establish guidelines for the selection of 624 625 potential sites for constructing UPSH plants.

The main difficulty faced with numerical models is ascertaining the characteristics of the discharged water during each cycle. This problem can be addressed by using the same modelling methodology proposed in this work. It consists of successive runs for which the maximum simulation time is progressively increased. After each pumping phase, the calculation is stopped, and the results are used to define the hydrochemical features of the subsequently discharged water.

This work considers three simplified hypothesis with respect the chemical composition of the porous medium. Obviously, hydrochemical changes induced by real UPSH plants will be different and will vary for each specific site because they depend on the chemical composition of the medium surrounding the underground reservoir. However, this work aims to highlight that the hydrochemical changes may be relevant. This is the proof that they may seriously impact the environment and the efficiency of the 638 UPSH plants. In fact, if they are not properly considered, they could put at risk the whole639 feasibility of future UPSH plants.

640

# 641 **6. Acknowledgements**

E. Pujades and A. Jurado gratefully acknowledges the financial support from the
University of Liège and the EU through the Marie Curie BeIPD-COFUND postdoctoral
fellowship programme (2014-2016 and 2015-2017 "Fellows from FP7-MSCA-COFUND,
600405"). This research was supported by the Public Service of Wallonia – Department
of Energy and Sustainable Building through the Smartwater project.

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# 861 Figure captions

Figure 1. General view of the problem. An underground reservoir (cavity) of 50 m by 50 m on 10 m height is considered. It is located below the water table.

Figure 2. a) Schematic view of the model domain. Half of the problem is modelled by taking advantage of its symmetry. b) View of the meshed numerical model.

Figure 3. pH evolutions in the surface (black continuous line) and underground (double
continuous grey line) reservoirs and in the surrounding porous medium at the
distances of 5 (dotted line) and 15 m (dashed line). The results are shown for both
scenarios (Sce1 on the left and Sce2 on the right) under the hypotheses H1 (Figure
3a), H2 (Figure 3b) and H3 (Figure 3c).

Figure 4. Computed mass of the precipitated minerals in the surface reservoir during the
simulated time. The results are shown for both scenarios (Sce1 on the left and
Sce2 on the right) under hypotheses H1 (figure 4a), H2 (Figure 4b) and H3 (Figure
4c).

Figure 5. Percentage evolution of the dissolved minerals in the surrounding porous medium at 5 m from the underground reservoir. The results are shown for Sce1 (atmosphere-water equilibrium) and Sce2 (no equilibrium) under hypotheses H1 (Figure 5a), H2 (Figure 5b) and H3 (Figure 5c).

Figure 6. CO<sub>2</sub> emissions from the surface reservoir to the atmosphere. The results are shown for Sce1 (atmosphere-water equilibrium) and Sce2 (no equilibrium) under hypotheses H2 (top) and H3 (bottom).

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1092	Figure 6

1093	Table captions
1094	Table 1. Released $CO_2$ (g/kWh) generated. The results are obtained under hypotheses
1095	H2 and H3 considering the total $CO_2$ released during the simulated time and the volume
1096	of CO₂ released once the emission rates stabilize (≈after 10 days).
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1128	Tables			
1129				
1130				
1131		Hypothesis H2	Sce1 (g of CO2 per kWh)	Sce2 (g of CO2 per kWh)
1132		Total CO <sub>2</sub> released	28	13
1133		CO2 released after	45	
1134		stabilization	15	9
1135				
1136			Sce1 (g of CO2	Sce2 (g of CO2
1100		Hypothesis H3	per kWh)	per kWh)
1137 1138		Total CO <sub>2</sub> released	per kWh) 14	per kWh) 5
1137 1138 1139		Hypothesis H3 Total CO <sub>2</sub> released CO2 released after stabilization	per kWh) 14 0.42	per kWh) 5 0.052
1137 1137 1138 1139 1140		Hypothesis H3 Total CO <sub>2</sub> released CO2 released after stabilization	per kWh) 14 0.42	per kWh) 5 0.052
1137 1137 1138 1139 1140 1141		Hypothesis H3 Total CO <sub>2</sub> released CO2 released after stabilization	per kWh) 14 0.42	per kWh) 5 0.052
1137 1137 1138 1139 1140 1141 1142	Table 1	Hypothesis H3 Total CO <sub>2</sub> released CO2 released after stabilization	per kWh) 14 0.42	per kWh) 5 0.052

## 1144 Appendix A

This appendix shows the results of a sensitivity analysis in which the specific surface area 1145 1146 (SSA) of the pyrite and the volume of pumped/discharged water are modified. Note that the effects 1147 of modifying the volume of the pumped/discharged water are analogous to those induced by the 1148 modification of the size of the underground reservoir. Results of Sce3 and Sce4 are compared with those obtained for Sce1. The SSA of pyrite is reduced one order of magnitude at Sce3 while 1149 1150 the volume of pumped/discharged water is reduced in half at Sce4. Figure A2.1 and figure A2.2 display results for hypotheses H1 and H2, respectively. Note that, hypotheses H3 is not 1151 1152 considered because significant results were not observed when the SSA of calcite was reduced. 1153 In fact, computed results considering kinetics of calcite were very similar to those obtained 1154 considering chemical equilibrium of calcite because its reaction rate is fast in comparison with our 1155 time scale. Concerning the volume of pumped/discharged water, the expected behavior is similar to that obtained for hypothesis H2. The main conclusions of this sensitivity analysis are that 1) 1156 1157 trends are the same, 2) the behavior is slightly modified, and 3) the impacts of modifying the SSA 1158 and the volume of pumped/discharged water are similar for both considered hypotheses. Main 1159 changes are described below.

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1161 Hypothesis H1:

1162 pH in the reservoirs (Figures A2.1 a and b) increases when the SSA of pyrite is reduced 1163 because less pyrite is dissolved in the surrounding porous medium and less protons are released. 1164 Larger mineral precipitations (per Kg of water) of goethite and schwertmannite are observed in 1165 Sce4 (Figure A2.1 c). Discharged water is more time in contact with the rock because the 1166 underground reservoir is not totally emptied and filled in Sce4, which may explain that more 1167 dissolved minerals are available to precipitate in the surface reservoir. Obviously, schwertmannite 1168 precipitation decreases for Sce3 since less pyrite is dissolved. Little differences are observed concerning the pH at the surrounding porous medium (5 m from the underground reservoir) 1169 1170 (Figure A2.1 d). The most remarkable aspect is that pH increases when the SSA and the pumped/discharged volume of water are reduced. Finally, the evolution of the concentration of 1171 pyrite at the surrounding medium is displayed at 5 m (Figure A2.1 e) and 1 m (Figure A2.1 f) from 1172

the underground reservoir. Apparently, more pyrite is dissolved at 5 m when the SSA is dissolved. However, this occurs because much less pyrite is dissolved in Sce3 near the underground reservoir (1 m) and more oxygen is available at 5 m. In Sce4, the dissolved pyrite is similar to that dissolved in Sce1 at 1 m whilst is much lower at 5 m because less discharged water reach this location. Globally, the volume of dissolved pyrite decrease for scenarios Sce3 and 4.

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**Figure A2.1.** Results obtained for the scenarios Sce1, Sce3 and Sce4. This results allow determining the effects of reducing the SSA (Sce3) and the volume of pumped/discharged water (Sce4) when H1 is considered.

1201 Hypothesis H2:

In general terms, the pH in the reservoirs (Figures A2.2 a and b) increases when the SSA and the volume of pumped/discharged water are decreased. The only exception is observed in the underground reservoir for Sce4. In this case, during early times the pH is lower than that computed for Sce1, after pH is equal for both scenarios. Mineral precipitations in the surface reservoir (Figure A2.1 c) increase when the volume of pumped/discharged water is reduced, while they decrease if the SSA is diminished because the dissolution of minerals in the surrounding porous medium is lowered. Concerning the pH evolution in the surrounding porous medium (Figure A2.1 d), the pH increases when the SSA is reduced because less pyrite is dissolved. In addition, the pH decreases when the volume of pumped/discharged water is reduced because the water exchanges between the underground reservoir and the surrounding porous medium are lowered. Consequently, the impact of the surrounding groundwater is smaller. Finally, as in the previous hypothesis, the mass of dissolved minerals decreases in Sce4 because the water exchanges are lowered. Moreover, it is also observed that the mass of dissolved minerals at 5 m increases when the SSA is reduced. This effect is produced because more reactants (dissolved oxygen) can reach this location since they are not previously consumed when the SSA is reduced. This fact is corroborated by the computed dissolutions at 1 m from the underground reservoir. At this location, much less minerals are dissolved for Sce3. In global terms, the volume of dissolved minerals decrease for scenarios Sce3 and Sce4.



Figure A2.2. Results obtained for the scenarios Sce1, Sce3 and Sce4. This results allow determining the effects of reducing the SSA (Sce3) and the volume of pumped/discharged water (Sce4) when H2 is considered.

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