Understanding the fate of H2S injected in basalts by means of time-domain induced polarization geophysical logging

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Abstract

To help meet emission standards, hydrogen sulfide (H2S) from geothermal production may be injected back into the subsurface, where basalt offers, in theory, the capacity to mineralize H2S into pyrite. Ensuring the viability of this pollution mitigation technology requires information on how much H2S is mineralized, at what rate and where. To date, monitoring efforts of field-scale H2S reinjection have mostly occurred via mass balance calculations, typically capturing less than 5% of the injected fluid. While these studies, along with laboratory experiments and geochemical models, conclude effective H2S mineralization, their extrapolation to quantify mineralization and its persistence over time leads to considerable uncertainty. Here, a geophysical methodology, using time-domain induced polarization (TDIP) logging in two of the injection wells (NN3 and NN4), is developed to follow the fate of H2S re-injected at Nesjavellir geothermal site in south-west Iceland. Results show a strong chargeability increase at +40 days, corresponding to precipitation of up to $1\\%$ in NN4 and $2\\%$ in NN3 according to laboratory-based relationships. A uniform increase is observed along NN4, whereas it is localized below 450 in NN3. Changes are more pronounced with the larger electrode spacing, indicating that pyrite precipitation takes place away from the wells. Furthermore, a chargeability decrease is observed at later monitoring rounds in both wells, suggesting that pyrite is either passivated or re-dissolved after precipitating. These results highlight the ability of TDIP logging to monitor pyrite mineralization and have implications for understanding the fate of H2S upon subsurface storage in basaltic environments.

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¹⁰ Key Points:

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11	•	${\cal H}_2 S$ reinjection in basalt and mineralization into pyrite was monitored using TDIP
12		logging in two of the injection wells
13	•	Chargeability increases observed at +40 days (first monitoring round) correspond
14		to $1-2\%$ pyrite precipitation with great spatial variability
15	•	Subsequent monitoring rounds show a decrease in chargeability, suggesting that
16		pyrite is either passivated or re-dissolved

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17 Abstract

To help meet emission standards, hydrogen sulfide (H_2S) from geothermal production 18 may be injected back into the subsurface, where basalt offers, in theory, the capacity to 19 mineralize H_2S into pyrite. Ensuring the viability of this pollution mitigation technol-20 ogy requires information on how much H_2S is mineralized, at what rate and where. To 21 date, monitoring efforts of field-scale H_2S reinjection have mostly occurred via mass bal-22 ance calculations, typically capturing less than 5% of the injected fluid. While these stud-23 ies, along with laboratory experiments and geochemical models, conclude effective H_2S 24 mineralization, their extrapolation to quantify mineralization and its persistence over 25 time leads to considerable uncertainty. Here, a geophysical methodology, using time-domain 26 induced polarization (TDIP) logging in two of the injection wells (NN3 and NN4), is de-27 veloped to follow the fate of H_2S re-injected at Nesjavellir geothermal site in south-west 28 Iceland. Results show a strong chargeability increase at +40 days, corresponding to pre-29 cipitation of up to 1% in NN4 and 2% in NN3 according to laboratory-based relation-30 ships. A uniform increase is observed along NN4, whereas it is localized below 450 in NN3. 31 Changes are more pronounced with the larger electrode spacing, indicating that pyrite 32 precipitation takes place away from the wells. Furthermore, a chargeability decrease is 33 observed at later monitoring rounds in both wells, suggesting that pyrite is either pas-34 sivated or re-dissolved after precipitating. These results highlight the ability of TDIP 35 logging to monitor pyrite mineralization and have implications for understanding the fate 36 of H_2S upon subsurface storage in basaltic environments. 37

³⁸ Plain Language Summary

High-temperature geothermal production is responsible for air pollution due to hy-39 drogen sulfide (H_2S) present in the magmatic fluid. To help meet emission standards, 40 H_2S may be injected back into the subsurface, where basalt offers the capacity to trans-41 form H_2S into pyrite. Transformation into an immobile mineral prevents further trans-42 port into the atmosphere, sea, surface streams, or lakes. However, ensuring the viabil-43 ity of this pollution mitigation technology requires information such as how much H_2S 44 is mineralized, at what rate and where, which are highly uncertain due to the hetero-45 geneity and inaccessibility of subsurface processes. Here, a geophysical monitoring method-46 ology is developed and tested during the re-injection of H_2S at Nesjavellir geothermal 47 site in south-west Iceland. According to laboratory studies, pyrite precipitation is ex-48 pected to increase the electrical capacitance, "chargeability", of the subsurface. Using the 49 so-called time-domain induced-polarization (TDIP) method embedded in a wireline log-50 ging tool, the chargeability of a 2m-wide cylinder around the injection wells is measured 51 with high spatial-resolution before and during H_2S injection. A strong chargeability in-52 crease at +40 days indicates that up to 2% pyrite is formed. A subsequent decrease at 53 later rounds raises questions on whether pyrite is re-dissolved or passivated by other sec-54 ondary minerals. 55

⁵⁶ 1 Introduction

Geothermal energy production emits an estimated 0.2 Mt/yr of hydrogen sulfide 57 (H_2S) globally, with Iceland alone emitting 30 kt/yr (Marieni et al., 2018). These anthro-58 pogenic H_2S emissions are sourced from the geothermal production steams, which con-59 tain significant concentrations of H_2S (Prikryl et al., 2018; Stefánsson et al., 2011). H_2S 60 emissions are toxic to humans and can be fatal in concentrations as low as 320 ppm (World 61 Health Organization - Regional Office for Europe, 2000). H_2S emissions also pose a threat 62 to the environment, oxidizing when exposed to atmospheric oxygen to form acid rain (Greaver 63 et al., 2012). Recent air quality regulations, both internationally and within Iceland, have 64 limited the amount of H_2S atmospheric emissions to 50-150 µg/m3 per 24 hours (Aradóttir 65

et al., 2015; World Health Organization - Regional Office for Europe, 2000; Iceland Ministry of the Environment, Energy and Climate, 2010).

To reduce emissions from geothermal energy production in Iceland, H_2S can be dis-68 solved into the geothermal wastewater and injected into the basalt subsurface (Mamrosh 69 et al., 2014). To help meet emission standards, H_2S from the geothermal production steam 70 may be captured at the power plant, dissolved into geothermal wastewater, and injected 71 into the basalt subsurface (Mamrosh et al., 2014). Geochemical and reactive transport 72 models suggest that such a H_2S -injection results in effective sulfide mineralization, with 73 74 pyrite being the dominant sulfur bearing mineral (Prikryl et al., 2018; Stefánsson et al., 2011; Marieni et al., 2018; Bacon et al., 2014), due to the high reactivity of basalt and 75 its divalent cations content (up to 25 wt.% Ca, Mg, Fe) (Matter et al., 2016; Gysi & Ste-76 fánsson, 2008). However, these findings are based on simulations and laboratory exper-77 iments with limited evidence from field tests (Robin et al., 2020). In practice, the H_2S 78 mineral storage process requires careful monitoring to identify any adverse effects of the 79 injection, such as the acidification of the shallow groundwater system and the mobiliza-80 tion of toxic metals from the basaltic rocks (Floor et al., 2011; Cuoco et al., 2013; Delmelle 81 et al., 2015; Galeczka et al., 2016; Flaathen & Gislason, 2007). 82

Current monitoring practices for H_2S mineralization into pyrite are similar to those 83 for CO_2 mineralization into carbonates. They consist mainly of quantification by (i) mass 84 balance calculations using tracer tests (Matter et al., 2016), (ii) transport models (Ratouis 85 et al., 2022), and (iii) steady-state "reaction path" geochemical models based on chem-86 ical monitoring. Chemical and mineral analyses on precipitates formed on metallic sur-87 faces within monitoring wells also bring qualitative mineralization evidence (Matter et 88 al., 2016). More recently, the triple sulfur isotope systematics (32S, 33S, and 34S) of geother-89 mal fluids were investigated as a possible tool to trace H_2S sequestration by sulfide min-90 eralization in the geothermal reservoir (Robin et al., 2020). One major pitfall of these 91 monitoring practices is that they typically capture less than 5% of the flow and extrap-92 olate the results (Matter et al., 2016), whereas the remaining 95% may include undetected 93 upward migration through fractures and springs. In addition, a key assumption behind 94 the quantification is that the amount of dissolved gas not captured by the measurement 95 downstream, compared to expectations, has mineralized and that mineralization is per-96 manent (Gunnarsson et al., 2018; Matter et al., 2016). While physical verification of these 97 assumptions, e.g., through post-injection core-drilling, has not yet been possible (Carbfix, 98 2022), other studies focusing on CO_2 indicate that additional processes to carbonate min-99 eralization can happen, namely the formation of unstable organic carbon due to micro-100 bial activity (Trias et al., 2017). Therefore, quantifying mineralization with these meth-101 ods, whether sulfide or carbonate, leads to considerable uncertainty (White et al., 2020). 102

Time-lapse geophysics is a group of potentially suitable methods to obtain in-situ 103 information on the spatial distribution and magnitude of H_2S mineralization, as well as 104 its evolution over time. Time-lapse geophysics as a monitoring technique can, in theory, 105 capture dynamic processes with high spatial and temporal resolutions (Hermans et al., 106 2023; Lévy et al., 2022). Geo-electrical methods are particularly suitable for pyrite de-107 tection. Underground massive metallic ore deposits, as well as veinlets and disseminated 108 sulfides, have been detected and discriminated with induced polarization (IP) field mea-109 surements (Pelton et al., 1978; Placencia-Gómez, 2015; Börner et al., 2018). Petrophys-110 ical laboratory measurements also indicate that the presence of pyrite can be detected 111 in altered volcanic rocks and quantified to some extent, with induced polarization (Lévy, 112 Gibert, et al., 2019). The polarizability, also called chargeability, describes the ampli-113 tude of electrical polarization in rocks, which is particularly strong at interfaces between 114 fluid and semi-conductors, such as pyrite or magnetite (Bücker et al., 2018; Gurin et al., 115 2015; Abdulsamad et al., 2017). The reorganization of charges at the interface between 116 pore fluid and metallic particles creates local electrical "displacement" currents, which 117 add up until a maximum voltage is reached. The time-delay between maximum voltage 118

and maximum current, also known as the phase-angle for sinusoidal signals, is caused 119 by the polarization (Bücker et al., 2018; Chelidze & Gueguen, 1999; Olhoeft, 1985). On 120 the other hand, the conductivity quantifies the amplitude of the electric conduction phe-121 nomenon in rocks, which is in particular enhanced by the presence of smectite, as well 122 as high porosity, salinity, and temperature (Flóvenz et al., 2005; Kristinsdóttir et al., 2010; 123 Lévy et al., 2018; Waxman & Smits, 1968). Geo-electrical field investigations carried out 124 at a geothermal site in North-East Iceland confirmed that the conductivity and IP sig-125 nals of naturally present pyrite, smectite, and magnetite can be picked up down to 200 126 m depth (Lévy, Maurya, et al., 2019). 127

In January 2021, the geothermal power company ON Power started a small-scale 128 injection of H_2S and CO_2 dissolved in seal water from liquid-ring vacuum pumps at Nes-129 javellir, South-West Iceland. This small-scale injection runs alongside larger-scale injec-130 tion of separated geothermal fluid from the power plant that has been part of standard 131 operations since 1990, shortly after the power plant was installed. The injection takes 132 place in eight shallow wells ranging from 311 m to 660 m deep. We use this injection ex-133 periment to investigate the subsurface induced polarization response over time and as-134 sess (i) the sensitivity of time-lapse geophysics to pyrite precipitation and (ii) the sus-135 tainability of pyrite mineralization. As surface geophysical measurements lose resolution 136 with depth, wireline logging was considered a relevant intermediate scale between lab-137 oratory and surface geophysical measurements to assess pyrite precipitation in the vicin-138 ity of the injection wells. It is thus the focus of this study. 139

¹⁴⁰ 2 Theory and Background

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2.1 Electrical response of pyrite with frequency-domain IP (FDIP)

In an electrical context, geological formations can be thought of as a combination 142 of resistors and capacitors (Lévy, 2019). Ions in pore water and those connected to clay 143 minerals are charge carriers in aquifers. Conductive water and clay minerals can be rep-144 resented by conductors with high conductivity, as opposed to freshwater and pure quartz 145 sand or gravel, which have lower conductivity. Semi-conducting metallic particles (e.g., 146 pyrite or magnetite) can cause electronic conduction beyond 20% per volume if the par-147 ticles are interconnected (e.g., veinlets) (Slater et al., 2005, 2006; Revil, Abdel Aal, et 148 al., 2015), but disseminated particles mainly cause electrical polarization, an effect sim-149 ilar to that of a capacitor in an electric circuit. Polarization caused by semi-conductors 150 is usually several orders of magnitude stronger than sediment/water interfaces (Revil et 151 al., 2017; Slater et al., 2005, 2006). This is due to the specifics of the crystal structure 152 of semi-conducting minerals, such as pyrite and magnetite, and the possibility for elec-153 trons and holes to "jump" and rapidly redistribute within the metallic particle in response 154 to the external field (Shuey, 2012; Pridmore & Shuey, 1976). This redistribution causes 155 the accumulation of ions at the interface between fluid and metallic particles. 156

Figure 1 illustrates the large polarization, represented by the phase angle parameter, of volcanic samples containing pyrite, as opposed to a volcanic sample of similar geology and full of smectite but without pyrite; the latter being more conductive but much less polarizable, as described in details by Lévy, Gibert, et al. (2019) and Lévy, Weller, and Gibert (2019). It is also visible in Figure 1 that the grain size distribution of metallic particles affects the frequency at which the phase angle peak occurs.

Several models have been developed to describe how ions reversibly accumulate at the interface between pore fluid and metallic particles, causing polarization. One of the most widespread phenomenological relaxation models to interpret FDIP data is the Pelton model (Pelton et al., 1978; Tarasov & Titov, 2013; Weller & Slater, 2022), given by equation 1.



Figure 1. Laboratory results for naturally altered volcanic samples from the Krafla geothermal field (north-east Iceland) containing varying amounts of pyrite with different grain sizes. Top panels: FDIP data for four different samples, left = impedance modulus, right = phase angle, as a function of frequency. Middle panels: scanning electron microscope (SEM) images for the three samples containing pyrite. Pyrite grains are highlighted in red. Bottom panels: distribution of pyrite grain size, based on the SEM images of the three samples.

$$\rho^*(f) = \rho_{DC} \left[1 - m\left(1 - \frac{1}{1 + (i2\pi f\tau)^c}\right)\right] \tag{1}$$

with four fitting parameters: the DC resistivity ρ_{DC} , the chargeability m, the Pelton time constant τ and the so-called Cole–Cole exponent c.

However, the Pelton model, presented in equation 1, is only valid for spectra measured on samples with a single grain size fraction. Variations of this model for two grain
size fractions exist but the fitting parameters are not directly comparable to each other,
and even the two grain size fractions model may not represent the complexity of naturally occurring pyrite crystals in volcanic rocks (Lévy, Gibert, et al., 2019).

A more general approach to analyzing FDIP data is the Debye decomposition (Nordsiek Weller, 2008). With this approach, the frequency-dependent complex electrical resistivity $\rho^*(f)$ describes an IP spectrum that can be fitted by a superposition of N Debye terms (Pelton models with c = 1), as shown in equation 2.

$$\rho^*(f) = \rho_{DC} \left[1 - \sum_{k=1}^N m_k \left(1 - \frac{1}{1 + i2\pi f \tau_k}\right)\right]$$
(2)

where m_k and τ_k are pairs of partial chargeability and relaxation time of a single Debye model. The function $m_k = f(\tau_k)$ is also referred to as relaxation time distribution (RTD). The N chargeability values m_k , which are related to a set of pre-defined relaxation times τ_k , and the value of DC resistivity ρ_{DC} are the result of a fitting procedure described by Nordsiek and Weller (2008). Two parameters can be integrated from the Debye decomposition to simplify the interpretations: the total chargeability m_{tot} (equation 3) and the mean relaxation time and τ_{mean} (equation 4).

$$m_{tot} = \sum_{k=1}^{N} m_k \tag{3}$$

$$\tau_{mean} = exp(\frac{\sum_{k=1}^{N} m_k ln(\tau_k)}{m_{tot}}) \tag{4}$$

The main empirical petrophysical relationship linking the chargeability to pyrite volume in the absence of background polarization was developed by Revil, Florsch, and Mao (2015) (equation 5).

$$m = \frac{9}{2}p_v \tag{5}$$

where p_v is the volume fraction of pyrite (or other metallic particles) and m is the chargeability in V/V (dimensionless). In their study, the chargeability parameter is fitted to frequency-domain IP (FDIP) data by a Pelton model (Cole & Cole, 1941; Pelton et al., 1978). According to Martin and Weller (2023), the chargeability m fitted by a Pelton model (Pelton et al., 1978) and m_{tot} fitted by Debye decomposition (Nordsiek & Weller, 2008) are similar for sand-pyrite mixtures.

While the influence of temperature on the polarization effect is still poorly understood, recent laboratory experiments indicate that the phase angle in the presence of metallic particles is temperature independent in the range 5-50°C (Revil et al., 2018). 2.2 Frequency-domain and time-domain induced polarization (FDIP and TDIP)

In the field, the Time Domain Induced Polarization (TDIP) method is often preferred to FDIP, primarily due to its easier and faster implementation. TDIP consists of injecting a direct current pulse of finite duration (DC) through two current electrodes (A and B) and recording the resulting voltage at two potential electrodes (M and N) with a fine-enough sampling rate to capture the exponential charge and discharge of the voltage, during the on-time and off-time, respectively (Sumner, 1976; Telford et al., 1990). Further details on the TDIP methods can be found in Binley and Slater (2020).

In general, voltage discharge measured by TDIP is expected to have a stretched 207 shape, which describes the relaxation behavior in a similar manner as Cole-Cole and other 208 related FDIP models (Alvarez et al., 1991; Cole & Cole, 1941; Davidson & Cole, 1951; 209 Havriliak & Negami, 1966). More details are described in Appendix Appendix A. While 210 quantitatively connecting the TDIP and FDIP responses of rocks containing metallic par-211 ticles is not trivial (Alvarez et al., 1991), the Debye decomposition is a common method 212 for analyzing both TDIP and FDIP data (Nordsiek & Weller, 2008; Tarasov & Titov, 213 2007). An equivalence was found by Martin et al. (2021) between the Debye decompo-214 sition of FDIP and TDIP data, provided that the current injection time in TDIP was 215 long enough (ideally 64 seconds, but decent comparisons were found beyond 2 seconds). 216 With 2-second current injection, as is the case in this study, the coefficients from the De-217 by decomposition of TDIP data are slightly underestimated compared to FDIP data. 218 Despite this limit, this approach was considered the most relevant one. As opposed to 219 the integral chargeability, which tends to smoothen out noise in the data, the Debye de-220 composition tends to amplify the noise and, thus, requires careful data processing (Martin 221 et al., 2021). 222

- ²²³ 3 Materials and Methods
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3.1 H_2S production and re-injection at Nesjavellir geothermal field, Iceland

The methodology developed in this study for monitoring H_2S injection was tested 226 at the Nesjavellir geothermal field (SW Iceland), where a geothermal power station has 227 been operated since 1990 (Figure 2). Nesjavellir is a high-temperature geothermal field 228 $(> 200^{\circ}C \text{ above 1 km depth})$ characterized by a relatively high permeability (Zakharova 229 & Spichak, 2012). Nesjavellir is located just north of the Hengill central volcano, which 230 is situated on a ridge-ridge-transform triple junction (Foulger, 1988). The area is com-231 prised of mainly late Quaternary hyaloclastites and post-glacial age hyaloclastites and 232 basalt flows (Árnason et al., 1969; Foulger & Toomey, 1989). This study focuses on shal-233 low depths northeast of the geothermal production area and above the low-permeability 234 clay cap layer (approximately 500 m depth) (Schiffman & Fridleifsson, 1991; Gómez-Díaz 235 et al., 2022; Gunnarsdottir et al., 2020; Franzson & Gunnlaugsson, 2020). Intrusions are 236 rare at these shallow depths, and none have been identified in the upper 300 m (Franzson, 237 1988). 238

The Nesjavellir power station currently produces 120 MWe of electricity and 300 239 MWth of thermal energy for district heating. The hydrothermal production fluid (260-240 300°C) is sourced at 1000-1500 m depth (Snæbjörnsdóttir et al., 2020). Following a con-241 trolled pressure-decrease, the hydrothermal fluid is divided into (i) steam and (ii) geother-242 mal separated water at 192°C. Electricity generation at Nesjavellir uses 240 kg/s of steam 243 to power four turbines. The steam is then condensed, using 2,000 l/s of cold ground-244 water (5-7°C), to form condensate wastewater, composed primarily of distilled water. 245 In parallel, heat exchangers use the $192^{\circ}C$ geothermal separated water to heat cold ground-246 water to 87°C, which is then pumped to the capital Reykjavík, 27 km away. The sep-247



Figure 2. Aerial view of Nesjavellir valley (south-west Iceland). The white building shows the geothermal power plant. The two wells, NN3 and NN4, investigated in this study are indicated by white circles, while other wells in the area are shown with small black rectangle signs. A buried power line (132 kV), most likely responsible for the 50 Hz background noise, is shown in blue. Pipelines are shown in pink and roads in black.

Table 1. Details on the Nesjavellir NN injection wells in this study. The borehole fluid conductivity was measured with a QL40-FTC logging probe (*Mount Sopris QL40-FTC*, 2014) in July 2022 (+540 days). Temperature and caliper logs are presented in Figures B1 and B2 in Appendix Appendix A.

Well ID	NN-3	NN-4
Coordinates	64°06'38.23" N, 21°15'20.27" W	64°06'50.03" N, 21°15'0.06" W
Depth (m)	563	422
Deviation	0°	0°
Casing Depth (m)	205	201
Borehole mean diameter (mm)	405 (min=317 - max=680)	403 (min=324 - max=636)
Average flow Rate (kg/s)	11	113
Injected fluid temperature (°C)	65	85
Borehole temperature while measuring (°C)	63-69	57-71
Reservoir temperature (°C)	15-20	20-40
Borehole fluid conductivity (mS/m)	60-100	100
pH	8.5	9.1
H_2S (ppm)	74.42	76.81
$CO_2 (\text{ppm})$	13.5	8.35
SiO_2 (ppm)	435.08	428.09

arated water contains the dissolved solids from the hydrothermal fluid and thus has a 248 high potential for chemical pollution and corrosion. Starting in 2004 and prior to 2021, 249 wastewater composed of condensate and separation water has been disposed of in drainage 250 wells. This initial wastewater had negligible concentrations of dissolved CO_2 and H_2S . 251 Starting on January 29th, 2021, a third type of fluid, called seal water, is now mixed into 252 the condensate and separation water before injection. Seal water contains the non-condensable 253 fraction of gases, primarily H_2S , that pose corrosion problems to turbines over time. It 254 is formed by dissolving the CO_2 and H_2S gases from liquid ring vacuum pumps at the 255 condensers in cold groundwater or condensate water. The currently injected wastewa-256 ter contains, on average, 75 ppm of H_2S (Table 1), with the intention to sequester H_2S 257 through mineral storage. 258

Two injection wells are used for monitoring: NN3 and NN4, drilled into fresh lavas 259 to 563 m (NN3) and 422 m (NN4) depth in 2001, northeast of the power station. Their 260 specific lithology and alteration stage at the time of drilling (i.e., prior to large-scale warm 261 wastewater injection) are presented in Helgadóttir (2021), emphasizing an increase in the 262 amount of clay minerals below 400 m in NN3. Between 2004 and 2021, injection of the 263 initial wastewater into the cold groundwater system took place through these wells and 264 has probably been enhancing basaltic rock alteration compared to natural conditions. 265 Starting from January 2021, H_2S has been continuously injected through the final wastew-266 ater. Reaction path modeling from the literature predicts that basaltic glass dissolution 267 is the main source of iron over short time durations, with magnetite dissolution contribut-268 ing less iron due to the slower dissolution kinetics and more limited amount. Pyrite pre-269 cipitation is expected to be rapid (Prikryl et al., 2018; Stefánsson et al., 2011). An overview 270 of the area, including roads, boreholes, and buried infrastructure, is presented in Fig-271 ure 2. Information on the wells and the injection are given in Table 1. Temperature logs 272 in NN3 and NN4 measured the days of TDIP logging are presented in Figures B1 in Ap-273 pendix Appendix B, together with caliper logs measured after drilling (Figure B2). Fur-274 ther details on the wells can be found in Gómez-Díaz et al. (2022) and Hafstað (2003). 275

3.2 Field monitoring with QL40-IP logging tool

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The QL40-ELOG/IP (Mount Sopris QL40-IP, 2020) logging tool measures the elec-277 trical resistivity and time-domain IP (TDIP) response with electrodes made of stainless 278 steel. The tool has a diameter of 43 mm and uses the "normal" electrode configuration 279 with 16- and 64" electrode spacings, defined as the distance between the current elec-280 trode (A) and the potential electrode (N) (Helander, 1983). The current generator, built 281 into the tool, sends current at electrode A in all directions and the shielding of the log-282 ging wireline serves as the current sink. The reference potential electrode (M) is located 283 on top of an 8m-long isolation bridle, such that the distance between the potential elec-284 trodes (M and N) is considered infinite in comparison to the much smaller electrode spac-285 ing AN. Moreover, grounding of the logging unit (truck) is achieved by clamping the truck 286 ground onto the casing for security reasons (Advanced Logic Technology, 2021). The 64" 287 measurements are sensitive to polarization effects over a larger measurement volume, deeper 288 into the formation, compared to the 16". The radial investigation characteristic, as de-289 fined by Roy and Dhar (1971), considers electrostatic potentials of individual cylindri-290 cal shells of varying radii integrated to obtain the total contribution to the measured sig-291 nal. According to the formula by Roy and Dhar (1971), 75% of the measured signal with 292 the 64" spacing is caused by the cylindrical volume comprised in a 2.5-m radius from the 293 borehole central line. As a comparison, 75% of the measured signal with the 16" spac-294 ing comes from within 0.5 m, which is about twice the boreholes' mean radius (Table 295 1). Here, measurements took place every 25 cm, with a logging speed of 1.8 m/min. 296

A total of eight measuring rounds took place but only four measuring rounds are presented in this study for NN-3 and NN-4 due to mainly one problem encountered onsite, related to the inability to inject current higher than 200 mA, happening randomly at certain places and dates (Table 2).

Traditional applications of the QL40-IP tool, e.g. in the mining industry, are typ-301 ically fulfilled with 250 or 500 msec current injections (on-time) and a fixed number of 302 400 sample points in the whole cycle $T_{on,+}; T_{off,+}; T_{on,-}; T_{off,-}$. Two important differ-303 ences between traditional applications and the present study led to the development of 304 a new processing board. First, Martin et al. (2021) show that FDIP and TDIP data mea-305 sured on the same samples decently overlap when current injections last at least 2 sec-306 onds during TDIP data acquisition. This overlap is key to quantifying pyrite volumes 307 by Debye decomposition since these empirical relationships have only been established 308 with FDIP data. Second, the 132 kV buried power line connected to the Nesjavellir power 309 plant (Figure 2) is responsible for a strong 50 Hz signal in the TDIP data, as revealed 310 by initial tests in NN-4 prior to this monitoring study (December 2019). These tests were 311 carried out with the original QL40-IP tool, using 250-msec and 500-msec current injec-312 tions, corresponding to full cycles of 1 and 2 seconds, respectively, and thus sampling rates 313 of 400 and 200 Hz (400 sample points over the whole cycle). Using 2-second current in-314 jections, the full cycle lasts 8 seconds, which leads to a sampling rate of 50 Hz with 400 315 sample points and aliases the 50 Hz noise. Further investigations also showed that both 316 the amplitude and exact frequency of the "50 Hz" noise varied over time. Therefore, it 317 was clear that TDIP data acquisition with 2-second current injection required a larger 318 number of sample points to model the noise and remove it before data analysis. This was 319 achieved by increasing the memory of the processing board and led to an updated ver-320 sion of the QL40-IP tool used in the rest of this study. 321

From September 2020 (baseline) and on, the following IP acquisition settings were used: square-wave current injection $(T_{on,+}; T_{off,+}; T_{on,-}; T_{off,-}]$ for 2 seconds each, 8 seconds total), 450 Hz sampling rate of the voltage during the whole cycle (i.e. 3600 sample points), and wireline speed of 1.8 m/min. The 50 Hz noise is further illustrated in Figure 3 (first and second columns).



Figure 3. Full waveform (FW) TDIP data measured in NN3 with 16" and 64" electrode spacings (left column). The 50 Hz noise is further illustrated by zooming on the positive decay for the 64" data in the second column (signal in black), where the FW signal after removing the 50 Hz noise is also shown (in red). Further processing of the FW data into polarizability curves is shown in the third (linear scale) and fourth (log-log scale) columns. Results are shown at two different dates (baseline and +40 days) and two different depths (263 m and 470 m), showing varying levels of background noise.

Measuring date	NN-3	NN-4
December 2019 (preliminary)	no meas.	404 sample points
September 2020 (baseline)	good	good
March 2021 $(+40 \text{ days})$	good	good
November 2021 ($+270$ days)	good	current too low
February 2022 $(+380 \text{ days})$	current too low	good
July 2022 $(+540 \text{ days})$	good	good
October 2022 $(+630 \text{ days})$	current too low	current too low

 Table 2.
 TDIP monitoring timeline and evaluation of data quality. "Good" means data quality was sufficient for further analysis.

In parallel to the TDIP monitoring, the changes in electrical resistivity were also monitored to evaluate the possible contribution of processes other than pyrite precipitation to the changes in the IP response. Indeed, changes in borehole fluid composition, temperature, and precipitation of clay minerals are reflected by changes in resistivity. Similarly, the formation of connected clusters of pyrite particles that would lose their polarization properties while becoming more conductive would also cause changes in the resistivity logs.

3.3 Data processing from full waveform voltage signal to polarizability decays

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The apparent chargeability $M_{app}(t)$, also called polarizability and noted $\eta(t)$ in Martin et al. (2021) and Tarasov and Titov (2007), corresponds to the ratio between the decaying voltage during the off-time, $V_{decay}(t)$, and the maximum voltage reached during the on-time, V_{DC} . It is calculated by stacking positive and negative decays (equation 6).

$$M_{app}(t) = \eta(t) = \frac{V_{decay}(t)}{V_{DC}} = \frac{1}{2} \left(\frac{V_{decay,+}(t)}{V_{DC,+}} + \frac{V_{decay,-}(t)}{V_{DC,-}} \right)$$
(6)

A series of processing steps are carried out to clean the full waveform signal, optimize the signal-to-noise ratio, and maintain three time-decades of signal for the polarizability. The term polarizability is used in the rest of the paper to refer to the apparent chargeability defined in equation 6 in order to avoid confusion with the integral chargeability or the total chargeability parameter calculated from Debye decomposition.

First, a time-invariant self-potential (SP) voltage affects both V(t) and V_{DC} in the positive and negative decays. The SP offset at each depth and each date is calculated by averaging the positive and negative DC voltages, $V_{DC,+}$ and $V_{DC,-}$ and removed from V(t) and V_{DC} .

Second, harmonics of 50 Hz noise, due to a 132 kV buried power line, affect the volt-349 age full waveform measurements, as mentioned in the previous section and illustrated 350 in Figure 3. A power line noise model, based on the algorithm proposed by Larsen et 351 al. (2022) and applied in a similar manner by Olsson et al. (2016), is used to subtract 352 the harmonic noise from the signal. As shown by Larsen et al. (2022), the fundamen-353 tal frequency varies over time, rapidly oscillating between 49.9 and 50.1 Hz, and must 354 first be determined. The 2-second decaying voltage is thus decomposed into 200 msec 355 segments, where the frequency is fitted in every segment. Once the exact frequency is 356 determined, a model of the background noise, including harmonics up to Nyquist fre-357 quency, is removed from the signal. The denoised voltage after applying this 50 Hz fil-358 ter is presented in Figure 3 (panels in the second column) as the red curve on top of the 359 FW voltage (in black) during the positive off-time section of the full cycle. 360

Third, the voltage decays are re-gated into log-spaced time windows. This step trans-361 forms the full waveform voltage decay signal with 900 linearly spaced data points into 362 a gated voltage signal with 37 logarithmically spaced gates. More accurately, the re-gated 363 voltage consists of 10 gates in the first 22 msec (2.21 msec width, i.e., the original sam-364 pling step) and then exponentially larger gate widths (from 4.42 to 141.44 msec), over 365 which an exponentially increasing number of data points (from 2 to 64) are averaged. 366 Given the strong exponentially decaying shape in the early times, it appeared best to 367 keep all data points in the first 22 msec. Since the original data points are kept as the 368 10 first gates, it corresponds to a "square-gate" with one data point averaged. To be con-369 sistent, the square gating strategy is kept throughout the decay. This is especially rel-370 evant because the signal can be locally approximated by linear functions in the later times. 371 At some depths, the first gate (centered at 1.1 msec) is affected by electromagnetic (EM) 372 coupling (see, for example, at +263 m in Figure 3). Therefore, it was discarded from all 373 the re-gated discharge curves, and the first effective gate is at 3.32 msec: visual inspec-374 tion of all the decays suggests that EM coupling is negligible at this gate, which is con-375 sistent with observations by Olsson et al. (2016). 376

Fourth, the positive and negative voltage decays are divided by the DC voltage and 377 stacked to obtain the polarizability in mV/V (equation 6). The stacking allows remov-378 ing extra noise with the same sign in the positive and negative decays. These four pro-379 cessing steps eventually provide the processed polarizability curves over almost three decades 380 (in the range of 3-2000 msec), as presented in Figure 3 with a linear scale (third column) 381 and log-log scale (fourth column). The linear scale allows better comparison to the full 382 waveform signal presented in the same figure, while the log-log scale is used in the rest 383 of this study. 384

Fifth, polarizability decays are averaged over 2-meter thick sections (Figure 4), cor-385 responding to 8 decays averaged together (0.25 m spatial resolution). This is done to fa-386 cilitate the comparison between measuring rounds since the exact depth of each single 387 measurement varies at different dates. It also allows smoothing out short-wavelength sig-388 nal oscillations that would prevent meaningful comparison between measuring rounds. 389 The standard deviation corresponding to this average also captures noisy areas where 390 the signal varies a lot over short distances. The averaging over 2-meter sections also takes 391 into consideration the fact that logging measurements at one depth are, in reality, sen-392 sitive to a sphere around this given depth. Finally, averaging over 2-meter sections helps 393 to identify and mitigate remaining harmonic noise. 394

Some polarizability curves remain noisy after the processing (non-monotoneously or non-exponentially decaying), e.g., at 263 m depth and +40 days shown in Figure 3 and at 382-384m and +40 days shown in Figure 4. Noisy processed polarizability curves are further addressed in the next subsection and discussed at the end of the paper.

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3.4 Estimations of pyrite precipitation from polarizability decays

A simple way to represent TDIP data (polarizability decays) as a function of depth is to integrate the decays to obtain the so-called "integral chargeability", M_{int} (Mao et al., 2016; Telford et al., 1990). It represents the area enclosed by the discharge curve, V(t), and its zero asymptote, in a given time-window, [t1:t2], divided by the primary voltage V_{DC} and the time-window width, as described by equation 7 (Bertin & Loeb, 1976; Sumner, 1976). An error on M_{int} , $Err(M_{int}9$, is calculated following equation 8 at every date d.

$$M_{int} = \frac{1}{t_2 - t_1} \int_{t_1}^{t_2} \frac{V(t)}{V_{DC}} dt$$
(7)



Figure 4. Resulting polarizability decays after the five processing steps. An example of noisy decay in NN3 is shown at 382-384 m depth, with larger error bars and a non-exponentially decaying shape.

$$Err(M_{int}(d)) = \frac{M_{int}(d)}{\overline{\eta}} \sqrt{\frac{1}{n} \sum_{i=1}^{n} std(\eta_i)^2}$$
(8)

where $std(\eta_i)$ is the standard deviation of the polarizability of the *i*-th gate within the averaged 2-meter thick section, *n* is the number of time gates in the polarizability decays and $\overline{\eta}$ is the average polarizability over the *n* gates.

The integral chargeability presents the advantage of smoothing out the noise. It is calculated here in the time interval ranging from the second gate (3.32 msec) to the last gate (1926 msec). M_{int} is used in this study as a qualitative assessment tool of the change in polarization over time at a given depth, the trend corresponding to pyrite formation or dissolution. The error bar of the relative difference Δ_{rel} is calculated with equation 9 for the integral chargeability M_{int} and the resistivity ρ .

$$Err(\Delta_{rel}M_{int}(d)) = \sqrt{(M_{int}(d) \cdot \frac{Err(M_{int}(d_0))}{M_{int}(d_0)^2})^2 + (\frac{Err(M_{int}(d))}{M_{int}(d_0)})^2}$$
$$Err(\Delta_{rel}\rho(d)) = \sqrt{(\rho(d) \cdot \frac{Err(\rho(d_0))}{\rho(d_0)^2})^2 + (\frac{Err(\rho(d))}{\rho(d_0)})^2}$$
(9)

where d and d_0 correspond to the current monitoring date and the baseline date, respectively, $Err(\rho(d))$ is the standard deviation within the averaged 2-meter thick section, and $Err(M_{int}(d))$ is calculated as per equation 8.

The integral chargeability calculated from TDIP data cannot be converted to pyrite 419 volume as M_{int} is not equivalent to the "Cole-Cole" or "Pelton" m in equation 5. Sev-420 eral approaches have been developed to relate TDIP and FDIP data, considering there 421 is no strict equivalence between Pelton models in frequency-domain and stretched ex-422 ponential functions in time-domain (Alvarez et al., 1991). As described in section 2, there 423 is an equivalence between the Debye decomposition calculated with FDIP and TDIP data, 424 with a slight underestimation of m_{tot} from TDIP data when the current on-time is 2 sec-425 onds (Martin et al., 2021). Therefore, the conversion to pyrite volume fraction is car-426 ried out here by using Debye decomposition. The polarizability $M_{app} = \eta(t)$ is fitted 427 by a non-negative linear least-square procedure, where the fitting function is given in equa-428 tion 10 and the optimization problem is expressed in equation 11. 429

$$\eta(t) = \sum_{k=1}^{N} m_k e^{\frac{-t}{\tau_k}} \tag{10}$$

$$\min_{x \ge 0} \|Cx - d\|_2^2 \tag{11}$$

where x is a vector with the m_k model parameters, d is a vector with the measured data $\eta(t_i)$ and C is the linear multiplier defined as $C_{i,k} = e^{\frac{-t_i}{\tau_k}}$, where t_i are the center times of the p gates (i=1..p) and τ_k are the pre-defined N relaxation times (k=1..N). Here we define the $\tau_{k,k=1..N}$ as a set of N = 300 logarithmically-spaced values of relaxation times in the range 3-2000, covering approximately three decades. During the procedure, most of the coefficients m_k are kept to 0 due to the non-negative constraints. The fit eventually results in 10-20 coefficients being non-zeros.

⁴³⁷ The total chargeability m_{tot} is calculated by summing all the m_k coefficients (equa-⁴³⁸ tion 3). The difference of m_{tot} at different measuring rounds, Δm_{tot} , is converted into ⁴³⁹ a difference in pyrite volume fraction p_v , Δp_v , following equation 12, which is based on ⁴⁴⁰ equation 5.

$$\Delta p_v(d)_{\%} = \frac{200}{9} [m_{tot}(d) - m_{tot}(d_0)]$$
(12)

where d and d_0 are the monitoring date and the baseline, respectively. Only the difference is converted, not the absolute pyrite volume fraction, in order to subtract the contribution of the background polarization that may be partly due to iron-oxides or other polarizable material present before the start of H_2S injection.

As opposed to the integral chargeability, the Debye decomposition seems to act as 445 a noise amplifier, similar to observations by Martin et al. (2021). As mentioned previ-446 ously, some polarizability curves remained noisy despite the numerous processing steps, 447 with a resulting non-decaying shape. Such curves should not be processed with Debye 448 decomposition. An additional processing step is introduced to ensure that only exponentially-449 decaying curves are processed with Debye decomposition. The criterion for ensuring this 450 shape is based on the Kohlrausch-Williams-Watts relaxation model for TDIP data (Alvarez 451 et al., 1991), presented in Appendix Appendix A. In practice, the polarizability curves 452 $\eta(t)$ are fitted with a stretched exponential function, presented in equation 13, before ap-453 plying the optimization procedure described in equations 10 and 11. 454

$$f(t) = \alpha e^{-\left(\frac{t}{\tau_{KWW}}\right)^{\beta}} \tag{13}$$

The fitting parameters α , β , and τ_{KWW} , further described in Appendix Appendix 455 A, are not used for interpretation in this study. Instead, the stretched exponential fit serves 456 two purposes: (i) calculating a deviation to a stretched exponential to remove outliers 457 polarizability curves where $R^2 < 0.998$ and (ii) smoothing out the polarizability decays 458 that are reasonably deviating to ensure the Debye decomposition is not fitting noise. In 459 order to confirm that this fitting procedure is not introducing a significant bias, the m_{tot} 460 and τ_{mean} resulting from Debye decomposition before and after the fitting procedure are 461 compared in Figures A1 and A2 in Appendix Appendix A. While other types of curves 462 may reflect IP effects, with e.g. negative polarizability values having a physical expla-463 nation in the context of dipole-dipole acquisition with sharp layer boundaries (Fiandaca 464 et al., 2022), the polarizability curves discarded here did not belong to these "hetero-465 dox transients", but rather had a distorted shape, similar to the curve in NN3 at +40466 days at 382-384 m depth, presented in Figure 4. 467

An error is calculated on the resulting values of m_{tot} (equation 3), taking into ac-468 count both the residuals to the stretched exponential fit and the standard deviation within 469 the averaged 2-meter sections, to help discriminate between variations that fall within 470 an uncertainty interval from significant variations over time. First, an error is calculated 471 for each gate of the polarizability curve as the sum of the 2-meter standard deviation 472 (corresponding to the variation within the averaged 2-meter section, see also equation 473 8) and the residuals to the stretched exponential fit (equation 14): if the residual is pos-474 itive, it is added to the positive error bar err_+ , if the residual is negative, it is added to 475 the negative error bar err_{-} . 476

$$Err_{+}(\eta_{i,fit}) = \begin{cases} std(\eta_{i}) + residuals(\eta_{i}, \eta_{i,fit}), & \text{if residuals}(\eta_{i}, \text{fit}) > 0\\ std(\eta_{i}), & \text{otherwise} \end{cases}$$
$$Err_{-}(\eta_{i,fit}) = \begin{cases} std(\eta_{i}) - residuals(\eta_{i}, \eta_{i,fit}), & \text{if residuals}(\eta_{i}, \text{fit}) < 0\\ std(\eta_{i}), & \text{otherwise} \end{cases}$$
(14)

where $\eta_{i,fit}$ is the value of the stretched exponential fit of the polarizability at the i-th gate, and Err_+ and Err_- are the positive and negative error bars on $\eta_{i,fit}$ at each gate.

Similarly to the error on M_{int} , a relative error is calculated for each decay using the root mean square of the error on all the gates divided by the average polarizability $\overline{\eta}$. This relative error is then applied to the total chargeability m_{tot} , then to Δm_{tot} (change from the baseline), and finally to the pyrite volume fraction change Δp_v , following equation 15.

$$Err_{rel} = \frac{1}{\overline{\eta}} \sqrt{\frac{1}{n} \sum_{i=1}^{n} \left(\frac{Err_{+}(\eta_{i,fit}) + Err_{-}(\eta_{i,fit})}{2}\right)^{2}}$$
$$Err(m_{tot}) = Err_{rel} \cdot m_{tot}$$
$$Err(\Delta m_{tot}(d)) = \sqrt{Err(m_{tot}(d))^{2} + Err(m_{tot}(d_{0}))^{2})}$$
$$Err(\Delta p_{v}(d)) = \frac{2}{90} \cdot Err(\Delta m_{tot}(d))$$
(15)

485 4 Results

486

4.1 Baseline resistivity and integral chargeability

In order to evaluate the amplitude and meaning of the changes in the following sec-487 tions, the resistivity and integral chargeability measured in the baseline are first presented 488 in Figure 5. The average resistivity of NN3 and NN4 is about 30 Ωm , with a clear thresh-489 old at 450 m in NN3, separating the well into two sections, above and below 30 Ωm . Both 490 in NN3 and NN4, a few intervals have resistivity values significantly higher than 30 Ωm , 491 up to 200 Ωm . These high-resistivity intervals coincide with low-chargeability intervals 492 (gray rectangles in Figure 5). The average integral chargeability is around 20-30 mV/V, 493 with generally higher values in NN4. 494

Intervals of higher resistivity correspond to more compact, less porous, litholog-495 ical layers, where less hydrothermal fluid flow and thus alteration is expected (Lévy et 496 al., 2018). Therefore, the relatively high chargeability in the baseline, as well as the co-497 incidence of locally high resistivity with locally low chargeability, suggest that hydrother-498 mal alteration had been occurring in the wells prior to the start of H_2S injection, espe-499 cially in intervals of low resistivity. Small amounts of magnetite are expected in unal-500 tered zones, typically 2-4 wt.% (Lévy et al., 2018) and may contribute to the chargeabil-501 ity in high resistivity layers (Lévy, Gibert, et al., 2019; Peshtani et al., 2022). Alteration 502 minerals, such as pyrite and smectite, are especially expected in permeable, fractured 503 layers (Lévy, Gibert, et al., 2019; Gudmundsson et al., 2010; Lévy et al., 2020) and may 504 be responsible for the higher chargeability in low resistivity intervals, particularly be-505 low 450 m in NN3 and below 375 m in NN4. Note that smectite only contributes to po-506 larization when present in small quantities, typically lower than 10% (Lévy, Weller, & 507 Gibert, 2019; Telford et al., 1990; Mendieta et al., 2021), while it tends to decrease the 508 overall polarization response at larger quantities (Lévy, Gibert, et al., 2019), as illustrated in Figure 1. Based on these observations, the interval below 450 m in NN3 is where ex-510 tensive pyrite precipitation is expected following H_2S injection, which is also consistent 511 with the observation that most of the "natural" hydrothermal alteration happens be-512 low 450 m in the area (Helgadóttir, 2021). 513

In both wells, higher chargeability is observed with the 64" spacing compared to the 16". Considering that the fluid is not chargeable in this frequency-range, this illustrates that the 64" images more volume of the chargeable properties of the rock formation, and thus more changes are also expected with the 64" spacing. On the other hand,



Figure 5. Resistivity (blue/red) and integral chargeability (green/black) logs in NN3 (left panels) and NN4 (right panels) using the 16- and 64" spacing (colors). All plots are based on an averaged signal every 2 meters, and the error bars correspond to the standard deviation within these 2-meter intervals. The dashed line shows the resistivity threshold of 30 Ωm . The gray rectangles indicate intervals with locally high resistivity and low integral chargeability.

similar resistivities are observed with both spacings, which reflects the fact that the fluid and the formation have resistivities in the same range. Based on fluid conductivity logging, the borehole fluid resistivity is around 10 Ωm in both wells (Table 1), i.e., only three times lower than the average 30 Ωm measured for the formation.

522

4.2 Changes in resistivity and integral chargeability over time

Logging results at +40 days (NN3 and NN4), +270 days (NN3), +380 days (NN4), 523 and +540 days (NN3 and NN4) are presented as relative differences (in percentage), com-524 pared to the baseline in Figure 6. The average value within a 2-meter thick interval is 525 taken for each round to smooth out rapid vertical variations. Error bars, calculated from 526 the standard deviation of this 2-meter average (equation 9), allow discriminating signif-527 icant changes over time from changes within the local variability. The resistivity after 528 baseline is further corrected to remove the temperature effect, following equation 16 and 529 using temperature logs shown in Figure B1 in order to assess resistivity changes reflect 530 changes in porosity or clay content. 531

$$\frac{\rho(T)}{\rho(T_0)} = 1 + \alpha_T (T_0 - T)$$
(16)

where T and T_0 are the current temperature and the reference temperature (corresponding to the baseline temperature), respectively. And $\alpha_T = 0.02^{\circ}C^{-1}$ is based on Arps (1953).

It can first be observed that changes depend on the electrode spacing for both the resistivity ρ and the integral chargeability M_{int} : relative differences are larger with the 64" spacing, indicating that more changes are happening in the rock formation than near the borehole wall (Figure 6). Resistivity variations are in many places within error bars, and thus, less focus is given here to interpret these variations.

Significant changes in the integral chargeability are observed. In NN4, M_{int} increases 540 at +40 days by 10-20% with the 64" spacing and 5-10% with the 16" spacing (Figure 541 6). At the next monitoring date, i.e., +380 days, M_{int} decreases back to the baseline in 542 most places with both spacings (note that data above 330 m were discarded for this date 543 due to low current injected). At the last monitoring date, +540 days, M_{int} remains at 544 baseline values down to 300 m, but the 64" signal decreases below baseline values be-545 tween 300 m and the bottom of the well (380 m). The 16" signal mostly remains at base-546 line values all along the well. A similar trend is observed in the bottom part of NN3, be-547 low 450 m: first M_{int} increases at +40 days by up to 50% with the 64" and up to 20% 548 with the 16" spacing. Then, M_{int} decreases for both spacings but remains above base-549 line values everywhere, unlike in NN4. At +540 days, below 450 m, slightly higher M_{int} 550 values are observed compared to +270 days for the 64" spacing. The 16" signal follows 551 the same trends everywhere, with lower amplitude. The maximum M_{int} increase is smaller 552 in NN4 than in NN3, but the increase is more uniformly spread along the borehole in 553 NN4. In particular, a significant increase is observed at +40 days in the whole well, i.e., 554 between 200 and 380 m. In NN3, the most important changes are located between 450 555 and 550 m, with only minimal changes down to 450 m. 556

A consistent observation links the 16- and 64" spacing in NN3 and NN4: M_{int} does not follow a monotonous increase as would have been expected if pyrite had been precipitating continuously or even if pyrite had precipitated up to a certain amount and remained in place. To illustrate further the non-monotonous pattern observed in the four panels showing M_{int} variations in Figure 6, the evolution of M_{int} with time in certain depth ranges is presented in Figure 7 for NN3 and NN4, using only the 64" spacing (larger variations) and showing absolute differences in mV/V.

Resistivity changes in NN3 with the 64" spacing show a spatial correlation with 564 M_{int} changes: a decrease of resistivity is observed in the upper part (above 450 m), and 565 an increase is observed in the bottom part. As opposed to the time-evolution observed 566 for M_{int} , the resistivity increase at the bottom of NN3 seems monotonous (Figure 6). 567 In NN4, a resistivity decrease is observed at +380 days above 300 m, and an increase 568 is observed between 300 and 350 m. However, an almost uniform resistivity decrease is 569 observed at +540 days, both with the 64" and 16" spacing, which indicates that the fluid 570 conductivity may have changed due to a change in fluid composition. Since the fluid con-571 ductivity was only measured at the last round (+540 days), further interpretation of the 572 resistivity changes regarding porosity, cementation, or clay content would be uncertain. 573

Overall, this qualitative analysis of the monitoring results indicates that (i) the integral chargeability strongly increases at +40 days in both wells and tends to decrease at the following monitoring rounds, going back or close to the baseline, (ii) the integral chargeability increase at +40 days is rather uniform over the whole depth interval in NN4 (200-400 m), while the increase in NN3 is localized between 450 and 550 m and (iii) the resistivity tends to increase monotonously over time at the bottom (below 450 m) of NN3,



Figure 6. Relative difference of the temperature-corrected resistivity ρ (left) and integral chargeability M_{int} (right) in NN3 (top) and NN4 (bottom), for both 64" and 16" electrode spacing. Colors indicate the monitoring rounds.



Figure 7. Evolution of the integral chargeability (absolute difference in mV/V) over time in NN3 (top) and NN4 (bottom) in different depth intervals, represented by the color scale. The dashed black line indicates the baseline level. Note the different scales for NN3 and NN4. Only data measured with 64" spacing are shown.

which may correspond to a decrease of porosity related to intense pyrite and secondary mineral precipitation, although impossible to quantify further.

In the following section, spectral information is extracted from TDIP monitoring results to allow a first-order quantification of pyrite precipitation in areas with an increase in M_{int} .

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4.3 Extraction of spectral information through Debye decomposition

Since Debye decomposition tends to amplify the noise in the polarizability curves, decays that could not be fitted by a stretched exponential function (equation 13) with $R^2 \ge 0.998$ were discarded. This also smoothed out the decays that were kept and avoided fitting noise with the Debye decomposition. The theoretical justification and illustration of using a stretched exponential model for TDIP data is further described in Appendix Appendix A (Figures A1 and A2). Figure 4 also illustrates that the stretched exponential fit has minimal difference with the polarizability data in most cases.

⁵⁹³ Debye decompositions, illustrated in Figure 8 for two sets of decays in NN3 and ⁵⁹⁴ NN4 (64" spacing), show a similar trend as observed in Figures 6 and 7. The polariz-⁵⁹⁵ ability decays at +40 days are above the baseline as well as the later monitoring rounds, ⁵⁹⁶ both in NN3 and NN4. This trend is also reflected in the coefficients m_k , especially at ⁵⁹⁷ long τ_k values, and the resulting m_{tot} , larger at +40 days. All Debye decompositions are ⁵⁹⁸ available as figure sets S1 to S4 in Supplementary Information.

As can be seen on Figure A1, only 70% of the decays passed all the processing steps. Most affected datasets were at +380 days in NN4 (above 330 m) and at +40 days in NN3 (above 450 m). The noisy data above 450 m at +40 days is attributed to two factors that can be observed in Figure 3: (i) significantly stronger 50 Hz noise above 450 m than be-



Figure 8. Debye decomposition for two sets of decays measured with the 64" spacing in a) NN3 at 492-494 m and b) NN4 at 362-364 m. Left: polarizability data and fit by the RTD function (equation 10); right: coefficients m_k for all the τ_k at the four measuring rounds.

low at all dates (reason unknown) and (ii) stronger 50 Hz noise at +40 days compared
to baseline, at all depths. As can be observed in Figure 3, the signal level is always better with the 16" (blue) than with the 64" (red) electrode spacing, which is expected considering that the measured voltage decreases for increasing spacing with this configuration. This explains that fewer 16" data were discarded during the Debye decomposition
filtering procedure (A1). However, the 64" data involve a larger depth of penetration around
the borehole and are thus more relevant for conversion to pyrite volume change.

On the one hand, the integral chargeability presents the advantage of smoothing 610 out noise and providing a qualitative overview of the trends for the whole dataset. On 611 the other hand, Debye decomposition allows extracting spectral information and further 612 quantifying pyrite volume fraction change. Ideally, a unique relation would exist between 613 m_{tot} values obtained by Debye decomposition and the integral chargeability M_{int} , which 614 would allow to convert M_{int} into pyrite volume fraction. However, we find that differ-615 ent relations exist between m_{tot} and M_{int} , depending on the mean relaxation time. In-616 vestigating further these relations is beyond the scope of this study. Thus, pyrite esti-617 mations in the next section are calculated on a reduced portion of the dataset, where De-618 by decomposition was carried out. 619

4.4 Quantification of pyrite volume fraction change

The absolute difference in total chargeability, Δm_{tot} , further converted into Δp_v 621 are presented in Figure 9 for the three monitoring rounds. While the same trends are 622 observed as in Figure 6, this conversion allows estimating the extent of pyrite precipi-623 tation or dissolution. Error bars also differentiate between significant differences in pyrite 624 volume fraction versus differences too uncertain to be interpreted further. Overall, pre-625 cipitation of up to 2% and 1% are observed with the 64" and 16" electrode spacing, re-626 spectively. Figure 9 illustrates that more volume fraction changes are observed when tak-627 ing into consideration a larger volume around the boreholes, which emphasizes once more 628 that changes are not limited to the very near vicinity of the injection boreholes. 629

In NN4, an increase in pyrite volume fraction at +40 days (+0.5-1% compared to 630 baseline) is observed with both the 16" and 64" electrode spacings at all depths, followed 631 by a decrease at +380 and +540 days. In addition, the 64" data indicate that the de-632 crease at +540 days extends below baseline values (-0.2-0.5% compared to baseline). In 633 NN3, changes are more depth-dependent than in NN4. Between 200 and 450 m, no sig-634 nificant changes are observed at any rounds. However, it is important to note that, at 635 +40 days, most of the 64" data in this depth-range were discarded during the Debye de-636 composition filtering procedure. Below 450 m, all the way down to 550 m, +1-2% in the 637 pyrite volume fraction is suggested by the 64" data and +0.5-1% by the 16" data. At 638 +380 and +540 days, the 64" and 16" data, where available, suggest a decrease towards 639 baseline values, yet never extending below baseline values, as opposed to NN4. 640

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4.5 Analysis of the mean relaxation time

Overall, relaxation times are shorter in NN4 than in NN3, with a maximum of 300 642 msec in NN4 and above 1000 msec in NN3 (Figure 10). It can also be noted that the vari-643 ations of the average relaxation times across the different monitoring rounds follow the 644 variations of m_{tot} . In NN3 below 450 m, τ_{mean} increases at +40 days and decreases back 645 at +270 days. In NN4 across the whole well, τ_{mean} increases at +40 days and decreases 646 back at +270 days. In NN3, significantly longer relaxation times are observed below 450647 m, which reflects the different shape of the polarizability decay curves, with most of the 648 decay happening at late times, as opposed to depths above 450 m. Similarly, in NN4, 649 decays at 346-348 m and at 380-382 m have a different shape and correspond to min-650 imum and maximum values of τ_{mean} over the whole well, 40 and 400 msec, respectively. 651



Figure 9. Estimations of pyrite fraction change (absolute difference of pyrite vol-%) in NN3 (top) and NN4 (bottom), based on TDIP logging with 64" and 16" electrode spacing, using Debye decomposition of the decays.



Figure 10. Left: mean relaxation times, τ_{mean} , from Debye decomposition in NN3 (top) and NN4 (bottom), for both the 64" and 16" spacings, at different monitoring rounds. Polarizability curves for four selected depths for each well, using the 64" spacing. The corresponding depths are shown on the depth- τ_{mean} plots with gray rectangles.

As was observed in Figure 1, longer relaxation times, corresponding to lower fre-652 quencies, can be associated with either larger metallic grain sizes or connected metal-653 lic grains, e.g., forming veins. In this case, relatively small pyrite grain sizes are expected 654 due to the short time available, especially after +40 days (around 10 μm according to 655 laboratory experiments presented by (Prikryl et al., 2018)). Therefore, long relaxation 656 times in NN3, below 450 m, may indicate connected pyrite particles in fractures. On the 657 other hand, shorter relaxation times in NN4 may be associated with small, disseminated 658 pyrite particles. 659

It is important to note here that TDIP data are only capturing polarization processes associated with relaxation times larger than 3 msec, which is above the typical range of relaxation times investigated by FDIP laboratory studies, such as Martin and Weller (2023) and (Revil, Abdel Aal, et al., 2015). Therefore, converting the relaxation time distributions into grain size distribution would be uncertain.

5 5 Discussion

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5.1 Interpreting subsurface processes behind the monitored resistivity and TDIP responses

First, the justification for converting total chargeability into pyrite volume frac-668 tion changes is discussed here. This conversion assumes that pyrite precipitation is the 669 process governing the increase in chargeability. Based on reaction path modeling car-670 ried out in similar conditions, it is likely that some amount of smectite precipitates (Galeczka 671 et al., 2022). Still, smectite precipitation is expected to have little influence on the IP 672 signal in the presence of pyrite and could not alone explain the observed chargeability 673 increases (Lévy, Gibert, et al., 2019; Lévy, Weller, & Gibert, 2019; Revil et al., 2017). 674 Precipitation of iron-(oxyhydro)oxides following basalt dissolution is also possible, but 675 only magnetic iron-oxides, such as magnetite or ilmenite, yield a strong polarization re-676 sponse (Peshtani et al., 2022). In these conditions, only non-magnetic Fe(III)-bearing 677 phases are expected, such as goethite or hematite, based on laboratory experiments (Menefee 678 et al., 2018; Andreani et al., 2009), field-scale modeling (Galeczka et al., 2022), and the 679 general low-temperature alteration sequence of Icelandic basalt (Crovisier et al., 1992; 680 Arnórsson et al., 1983; Gunnlaugsson & Arnòrsson, 1982; Stefánsson & Gíslason, 2001). 681 In addition, amorphous silica and other silica-based minerals, such as quartz and zeo-682 lites, are expected (Daval et al., 2011; Galeczka et al., 2022). Amorphous silica, zeolites, 683 and Fe(III)-bearing phases further have a strong tendency to clog the pore space (Menefee 684 et al., 2018; Andreani et al., 2009), without creating a distinctive polarization signal (Revil 685 et al., 2002). Finally, the influence of microbial activity on the IP response is still poorly 686 understood but seems to yield polarization of much smaller amplitude than the observed 687 changes (Mellage et al., 2018) and will therefore not be considered here. 688

With these considerations in mind, TDIP monitoring brings three types of insights: spatial variability, long-term evolution, and quantity of pyrite volume changes.

The spatial variability of pyrite precipitation is assessed as a function of depth, vari-691 ability between NN3 and NN4, and distance from injection well. First, it is clear that 692 more pyrite precipitation occurs in the depth range 450-550 m in NN3 than in the rest 693 of the well, while in NN4, rather uniform pyrite precipitation is inferred (Figures 6 and 694 9). This can be understood in light of the baseline resistivity (Figure 5), showing resis-695 tivity lower than 30 Ωm at levels deeper than 450 m in NN3. In this 450-550 m inter-696 val, more clay content was found at the time of drilling (Helgadóttir, 2021), indicating 697 preferential hydrothermal flow. While the injection rate is ten times larger in NN4 than 698 in NN3 (Table 1), more precipitation seems to occur at the bottom of NN3 compared 699 to NN4. This may be due to fracture flow transporting the H_2S -rich fluid rapidly away 700 in NN4, not allowing time for pyrite precipitation. This would be consistent with the over-701

⁷⁰² all lower resistivity in NN4 in the common depth interval 200-380 m, compared to NN3 ⁷⁰³ (Figure 5), and would mean that there exists an optimal permeability that maximizes ⁷⁰⁴ the volumetric change in pyrite, ensuring at the same time ample fluid supply (H_2S sup-⁷⁰⁵ ply) and sufficient fluid residence time (basalt dissolution).

Second, it can be inferred that more precipitation occurs away from the well than on the borehole walls, based on the comparison between the 64" and 16" datasets in both wells (Figure 6).

The time-evolution of resistivity, integral chargeability, total chargeability and mean 709 relaxation time show that subsurface processes may be more complex than the expected 710 continuous pyrite precipitation suggested by previous studies (Clark et al., 2020; Gun-711 narsson et al., 2018). Integral chargeability time-series presented in Figure 7 show that 712 in both wells, at most depths, the initial increase at +40 days is followed by a decrease, 713 extending below baseline values in NN4 at depths below 330 m. This trend, which is com-714 mon to the 64" and 16" spacings (Figure 6), is also observed with the total chargeabil-715 ity and relaxation time (Figures 10, 9 and A1). Here, three different interpretations are 716 explored: (i) pyrite is still precipitating after +40 days, but the IP signal decreases any-717 way, (ii) previously formed pyrite grains are "passivated" by a coating layer of secondary 718 minerals preventing polarization while no more pyrite is forming and (iii) pyrite is be-719 ing re-dissolved. 720

Interpretation (i) would be consistent with the observation by Martin and Weller (2023) that the same amount of pyrite, with increasing grain size, yields a lower IP signal. However, the relaxation time, which is the main indicator for particle size (Figure 1) also starts decreasing after +40 days (Figure 10), which indicates that pyrite particles, if they are still there, are not growing in size. This interpretation is, therefore, ruled out.

Interpretation (ii) is based on the fact that strong polarization effects are caused 727 by metallic particles only when they have a clean surface. If secondary minerals isolate 728 the pyrite surface from the fluid, the particles are passivated, and the polarization de-729 creases (Gurin et al., 2019). Amorphous silica and Fe(III)-bearing minerals are common 730 coating layers, which may coat pyrite grain surfaces, as well as reduce basalt dissolution 731 rates by coating fresh basalt and clogging the pore space (Menefee et al., 2018; Andreani 732 et al., 2009; Daval et al., 2011). Corrosion of pyrite surface in the presence of little amounts 733 of oxygen may also lead to a similar passivation effect, without significantly dissolving 734 the pyrite particle (Placencia-Gómez et al., 2013). 735

Interpretation (iii) contradicts the common idea that pyrite mineralization is sta-736 ble in this context once formed. However, practical field conditions differ from typical 737 hypotheses in reactive transport models, which are constrained by the availability of suit-738 able frameworks and databases. Firstly, a high disequilibrium near the borehole, where 739 H_2S -rich fluids are constantly injected, makes conventional "steady-state" assumptions 740 irrelevant. Secondly, previous experiments on CO_2 injection in basalt at Hellisheiði (sim-741 ilar set-up as the Nesjavellir field case presented here) triggered a bloom of iron-oxidizing 742 bacteria, such as *Gallionellaceae*, and more generally sulfur- and iron- oxidation mark-743 ers (Trias et al., 2017). At Nesjavellir, provided that oxygen was present in the system 744 at some point (which could be the case due to e.g., leakage in the pipe system happen-745 ing at the beginning of the H_2S injection), specific microbial communities may have de-746 veloped and triggered pyrite oxidation (Mielke et al., 2003; Percak-Dennett et al., 2017). 747 The fracture networks that are ubiquitous in basaltic rocks could also be a pathway for 748 intermittent oxygen delivery, which would extend the depth of the habitable zone for iron-749 oxidizing bacteria (Bochet et al., 2020). Overall, biologically-driven mechanisms, which 750 could explain an unexpected pyrite dissolution over time, are not taken into account in 751 previous studies claiming that pyrite mineralization is stable due to the difficulty of rep-752 resenting them correctly in reactive transport simulations. While high-temperature geother-753

mal systems may prevent the development of most microbial communities, the shallow
system studied here has a relatively low reservoir temperature, although the injected fluid
temperature is higher (Table 1).

The pyrite volume fraction change estimated in Figure 9 assumes, to simplify, that 757 all the decrease of chargeability is explained by pyrite dissolution. In practice, there may 758 be a mix of passivation and re-dissolution and, thus, a more conservative interpretation 759 of the vol.% presented in Figure 9 would be the change in pyrite volume fraction con-760 tributing to the IP signal. While it is impossible from geophysical data alone to deter-761 mine whether passivation (interpretation ii) or re-dissolution (interpretation iii) are re-762 sponsible for the observed signals, both could be further investigated by laboratory stud-763 ies, as well as by installing a more continuous geophysical monitoring infrastructure. 764

In addition to the uncertainty on the source of polarization decrease, the quantifi-765 cation of pyrite volume presents additional uncertainty. It should first be noted that the 766 influence of background polarization not related to sulfide minerals in the baseline (Fig-767 ure 5) is managed by only converting the difference of total chargeability between dif-768 ferent monitoring rounds into the difference in pyrite volume fraction. However, equa-769 tion 5 proposed by Revil, Florsch, and Mao (2015) and used here presents other limits, 770 as Martin and Weller (2023) found that the exact relation between m_{tot} and p_v depend 771 on the grain size of pyrite particles. Furthermore, the use of an equation calibrated with 772 FDIP measurements to convert TDIP measurements is made possible by "long" current 773 injections (2 seconds), where the TDIP and FDIP responses are supposed to overlap (Martin 774 et al., 2021). However, 2-second current injections remain short, compared to 64"second 775 used by Martin et al. (2021) to show the equivalence with laboratory measurements. There-776 fore, the pyrite difference calculated here from 2-second TDIP data is certainly under-777 estimated. It was also assumed here that, at a given depth, precipitation occurs uniformly 778 over a certain (undetermined) lateral extent, considered representative of near-borehole 779 conditions. Further modeling of the contribution from the different cylindrical layers around 780 the borehole to the measured signal, and thus of the pyrite volume in each of these lay-781 ers, could be achieved with inversion. 782

Finally, it is interesting to note that the mean relaxation time is not only time-dependent but also depth-dependent and that an overall consistency of the decay shapes at a given depth is seen over time (Figure 10). This suggests that the local lithology and fracture network influence the relaxation time, most likely by influencing the patterns for pyrite precipitation. Although this is purely speculative, it may have implications for a better understanding of fracture networks in subsurface storage systems.

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5.2 Requirements and added-value of IP-logging compared to current practices

As mentioned in the Introduction, current monitoring of field-scale H_2S reinjec-791 tion, and similarly of CO_2 reinjection, have major pitfalls: (i) tracer tests capture less 792 than 5% of the gas migration flow (Matter et al., 2016), (ii) mass balance calculations 793 assume that the amount of sulfur not captured by downstream measurements, compared 794 to expectations based on tracer tests, has mineralized into sulfide minerals (Gunnarsson 795 et al., 2018) and (iii) long-term predictions by reactive transport models are not calibrated 796 by field monitoring and thus present significant uncertainty (Ratouis et al., 2022; Aradót-797 tir et al., 2012; White et al., 2020). Little has been researched on the long-term stabil-798 ity of the newly formed minerals in a context where the continuous injection of H_2S (and similarly, CO_2) creates a constant disequilibrium in the system, in terms of acidity and 800 redox conditions in particular. Overall, current monitoring approaches are still indirect 801 and cannot physically confirm the presence, location, and quantity of expected miner-802 als in the rock matrix, nor their long-term stability. The results presented here show the 803 feasibility of monitoring pyrite precipitation with logging TDIP chargeability measure-804

ments within injection wells. However, several conditions/constraints/obstacles need to be overcome.

To start with, an appropriate tool is required. Compared to more conventional ap-807 plications of TDIP, typically in mining exploration, the expected polarization signal is 808 weaker (smaller volumes of sulfide minerals). At the same time, the presence of power 809 lines due to the proximity to the power plant creates a significant 50 Hz background noise. 810 Therefore, large current injection is essential to ensure a sufficient signal-to-noise ratio, 811 and the acquisition of voltage with a high sampling rate is essential to separate the con-812 813 tribution from 50 Hz noise and from the rock to the voltage waveform. In addition, the need for quantification of pyrite precipitation, and thus comparison to frequency-domain 814 IP calibrations, requires that current injection and voltage acquisition happen over a suf-815 ficiently long time to capture the polarization information needed for the interpretation. 816 All these specifications required the development of new hardware in the existing QL40-817 IP instrument. In particular, the need for both longer injection/acquisition time and a 818 higher sampling rate means that more data needs to be collected and recorded in the in-819 strument's memory. The instrument went from recording 100 samples per cycle (depth) 820 to 980. Nevertheless, a trade-off was adopted, and the current waveform, deemed less 821 critical for the analysis, was not recorded; only the average current at each depth was 822 recorded. In addition, the maximum injection/acquisition times were 2 seconds. Ideally, 823 recording the current full waveform, injecting during longer times, and combining dif-824 ferent injection times would allow a more accurate comparison to FDIP calibrations. 825

Field conditions are also important to consider to carry out successful TDIP log-826 ging. The instrument has a temperature limit of $70^{\circ}C$, and its exposure to corrosive flu-827 ids shall be limited. This can be challenging in the context of CO_2 and H_2S injection 828 wells at geothermal sites. On the other hand, injection wells are the location with strong 829 changes in mineralogy. At the Nesjavellir site, the need to stop the injection and cool 830 down the wells a few days before the measurements represented the main obstacle to re-831 peating the measurements more regularly. In addition, the method requires an opera-832 tor on-site to operate the instrument. In this context, where data redundancy is limited 833 while TDIP measurements are easily subject to noise, data loss is a critical aspect to man-834 age, especially for quantitative conversion. 835

Despite field limitations and remaining uncertainties, IP-logging monitoring brings 836 novel insights into the processes at play, as emphasized in the previous subsection. A pos-837 sible way forward would be to carry out the logging in dedicated monitoring wells rel-838 atively close to the injection wells to avoid the financial and environmental cost of stop-839 ping the injection during monitoring. Another possibility that would also fulfill the need 840 for more continuous data acquisition to overcome episodic data losses would be the in-841 stallation of permanent electrodes in dedicated wells. The electrodes would connect to 842 an instrument remaining at the surface during the whole monitoring period, and only 843 batteries would need to be changed periodically. Furthermore, this approach could al-844 low 2D or 3D monitoring if several wells are used together in a cross-borehole electrical 845 tomography approach. External data acquisition would also be helpful to further elu-846 cidate the physical processes behind the observed IP response. Monitoring the fluid con-847 ductivity over time appears essential to assess changes in fluid composition and further 848 interpret resistivity changes. Complementary logging methods may include sonic and den-849 sity logging for assessment of porosity changes, acoustic televiewer, and spinner logs for 850 mapping fractures and flow paths, and color optical camera for direct observations of the 851 precipitation of pyrite and biofilms at the borehole wall. 852

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5.3 Can the IP-logging method be applied for monitoring other reactive processes?

Due to the strong signal caused by sulfides and other metallic minerals (Pelton et 855 al., 1978), mapping ore deposits is still the main industrial application of IP methods 856 today. Here, we present the first field example where reactive processes involving the pre-857 cipitation and/or dissolution of metallic minerals are monitored with IP-logging. Other 858 field examples for monitoring, with IP methods, reactive processes involving metallic min-859 erals include the immobilization of uranium contamination by stimulating iron and sulfate-860 reducing microorganisms (Flores Orozco et al., 2011) as well as the injection of zero-valent-861 iron amendment to aid bioremediation of chlorinated solvents (Flores Orozco et al., 2015). 862 In these examples, surface TDIP is used, and the limited spatial and temporal resolu-863 tion complicates the interpretation. There, IP-logging could help improve the understand-864 ing of physical processes at play. 865

Studies on the link between IP and bacterial activity often find that microbially-866 mediated metallic minerals cause an IP response (Williams et al., 2005; Flores Orozco 867 et al., 2011; Slater et al., 2007). However, recent studies suggest that polarization of bac-868 terial cells themselves may control the IP response in certain cases (Mellage et al., 2018; 869 Strobel et al., 2023). Therefore, in addition to monitoring the bioremediation of contam-870 inants, IP-logging could also be applied to the study of microbial life in a dynamic nat-871 ural environment, such as intermittent oxic-anoxic fluid mixing in fractured rocks (Bochet 872 et al., 2020), provided that upscaling issues can be overcome (Mellage et al., 2018). 873

Finally, we evaluate to what extent the method developed here could be applied 874 to monitoring CO_2 injection in basalts. Laboratory studies suggest that calcite precip-875 itation causes significant polarization increase (Wu et al., 2010). However, others observe 876 limited polarization increase, or even a decrease, in response to calcite precipitation (Wu et al., 2011; Zhang et al., 2012; Saneiyan et al., 2018) as well as no decrease associated 878 with calcite dissolution (Halisch et al., 2018). Recent studies attribute these contradic-879 tory findings to the dominating influence of pore water chemical changes or dissolution 880 changes on the IP response (Rembert, 2021; Izumoto et al., 2020). Therefore, the cur-881 rent petrophysical knowledge on the electrical response of calcite is insufficient to allow 882 direct mapping of calcite in the field with IP methods. Nevertheless, H_2S is often as-883 sociated with CO_2 in emissions from geothermal or coal power plants (Wang et al., 2011; 884 Fridriksson et al., 2016). H_2S is often considered an impurity and is only injected with 885 CO_2 to minimize the cost of separation. Still, the formation of pyrite could be used as 886 a marker of mineralization processes, given that pyrite and calcite precipitations are pri-887 marily controlled by basalt dissolution rates (Prikryl et al., 2018; Stefánsson et al., 2011) 888 and also share similar kinetics (Plummer et al., 1979; Williamson & Rimstidt, 1994). 889

⁸⁹⁰ 6 Conclusions

To date, monitoring efforts of an implemented field-scale H_2S reinjection system have only occurred via concentration measurements, mass balance calculations, and analysis of precipitates on submersible pumps in injection and downgradient boreholes. These studies, along with geochemical numerical models and laboratory simulations, concluded effective H_2S sequestration through pyrite formation. Borehole sampling of fluids, although effective at identifying the occurrence of sulfide mineralization, provides limited quantitative, spatial, and temporal information on in-situ sulfide mineralization.

In order to bring insights into the subsurface physical processes controlling mineralization in the context of H_2S injection in basalts, as well as quantitative information on the sulfide mineralization in a specific case, a "shallow" (200-550 m) injection of H_2S at Nesjavellir geothermal field was monitored by means of resistivity and time-domain induced polarization wireline logging. Pyrite precipitation was expected to affect primarily the induced polarization signal, while other changes (e.g., fluid conductivity, temperature, clay minerals precipitation) were expected to affect the resistivity.

The QL40-IP logging tool, developed by the company Advanced Logic Technology, was modified to allow longer current injection and a larger sampling rate than previously. This was needed to filter a strong 50 Hz background signal due to the proximity of a buried power line and convert the TDIP data into sulfide volumes using a laboratory-based relationship.

The first logging measurements were carried out in the summer of 2019, and the baseline with the modified tool was measured in September 2020. H_2S injection began on January 29th, 2021. After the start of injection, five logging monitoring rounds (+40, +270, +380, +540, +630 days) were carried out in wells NN3 and NN4. Out of these five rounds, only three remained for each well due to too low current injected in the other rounds for unknown reasons.

Monitoring results show that polarization strongly increases at +40 days in both 916 wells and tends to decrease at the following monitoring rounds, going back close to the 917 baseline or below. The polarization increase is rather uniform over the whole depth in-918 terval in NN4 (200-400 m), while the increase is clearly localized between 450 and 550 919 m in NN3, where more hydrothermal alteration was also found prior to the injection of 920 any wastewater. Conversion of chargeability absolute difference into pyrite volume frac-921 tion change indicates precipitation of up to 1% in NN4 and 2% in NN3 at +40 days. In 922 both wells, changes are more pronounced with the larger electrode spacing (64"), indi-923 cating that pyrite precipitation takes place away from the well. 924

The decrease of chargeability observed in both wells at +270 and +380 days sug-925 gests that pyrite is either passivated or re-dissolved after precipitating. The presence of 926 specific microbial communities that can produce oxygen at these depths should be fur-927 ther investigated. Recommendations for future monitoring include: (i) installation of per-928 manent monitoring systems to better understand the trends, (ii) considering the possi-929 bility of pyrite re-dissolution in reactive transport modeling of H_2S sequestration, and 930 (iii) validation of geophysical results by additional in-situ measurements and observa-931 tion, such as color camera and multi-level water sampling in wells. 932

933 7 Open Research

The TDIP data used for monitoring H_2S fate in the study, together with the resistivity, fluid conductivity and temperature data, will be available on the following Zenodo repository at the time of publication (Lévy, 2023): https://doi.org/10.5281/zenodo .10145643 with Creative Commons Attribution 4.0 International license. For review purposes, data are shared as supplementary information.

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Appendix A Added-value and justification of the stretched exponential fitting procedure

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The stretched exponential function $f(t) = \alpha e^{-\left(\frac{t}{\tau_{KWW}}\right)^{\beta}}$, also named Kohlrausch-1373 Williams-Watts (KWW), and the Havriliak-Negami (HN) relaxation function $g(\omega) = \frac{1}{(1+(i\omega\tau_{HN})^{\alpha}_{HN})^{\gamma}_{HN}}$ have been widely used to describe the relaxation behavior of glass-forming liquids and complex systems, in time- and frequency-domain respectively (Alvarez et al., 1374 1375 1376 1991). The HN relaxation function is a generalization of the Pelton model with one more 1377 parameter (Havriliak & Negami, 1966). It also corresponds to the "Generalized Cole-1378 Cole" model (Pelton et al., 1983). A relationship among the parameters of the KWW 1379 and HN models is suggested by the fact that both models yield an accurate description 1380 of real data. An empirical and mathematical relation between the coefficients of the KWW 1381 and HN functions is found by Alvarez et al. (1991). This relationship is not analytical 1382 since the HN and the KWW relaxation functions are not exactly Fourier transforms of 1383

each other. In particular, the relation between the relaxation times in both functions depends on the exponent beta in the KWW function, β . For example, for β close to 1, $\tau_{HN} \simeq \tau_{KWW}$, and for $\beta = 0.1$, $\tau_{HN} = 300\tau_{KWW}$. The KWW has one less parameter, so it can always be transformed into an HN model, but the contrary is not always true. The product $\alpha_{HN}\gamma_{HN}$ is close to β . In addition, even though the HN model is a generalized version of the Pelton model, the fitted chargeability parameter cannot be used to quantify pyrite as it is not strictly comparable to the chargeability in the Pelton model.

That's why the Debye decomposition is used here to relate the measured TDIP data with petrophysical relationships calibrated on FDIP data.

Nevertheless, based on the theory described by Alvarez et al. (1991), the stretched 1393 exponential function is used to automatically discriminate noisy discharge curves that 1394 do not follow the stretched exponential shape. Debye decomposition is only applied to 1395 discharge curves that can be reasonably fitted by a stretched exponential. Moreover, the 1396 stretched exponential fit allows smoothing out the "accepted" discharge curves that may 1397 still contain some noise, which in turn results in more meaningful outcomes from the De-1398 by decomposition. This is illustrated for the total chargeability, M_{tot} , and the mean re-1399 laxation time, τ_{mean} in Figures A1 and A2. In these figures, Debye decomposition car-1400 ried out on the stretched exponential fitted function is compared to Debye decomposi-1401 tion carried out on the polarizability data themselves (before the fitting step). The com-1402 parison clearly indicates less noise in the curves of SE fitted spectra. 1403

¹⁴⁰⁴ Appendix B Temperature and borehole diameter in NN3 and NN4



♦ +380 days ♦ +540 days Baseline +40 days ÷ ÷

Figure A1. Total chargeability from Debye decomposition in NN3 (top) and NN4 (bottom), with 64" (left) and 16" (right) spacing. The results are shown for Debye decompositions carried out on stretched exponential fit ("SE fit") and on original data. Note that the Debye decomposition on original data includes more discharge curves since it happens before the filtering procedure related to the SE fit.



Figure A2. Weighted-average relaxation time from Debye decomposition in NN3 (top) and NN4 (bottom), with 64" (left) and 16" (right) spacing. The results are shown for Debye decompositions carried out on stretched exponential fit ("SE fit") and on original data. Note that the Debye decomposition on original data includes more discharge curves since it happens before the filtering procedure related to the SE fit.



Figure B1. Temperature measured in NN3 and NN4 at the four monitoring rounds.



Figure B2. Borehole diameter (caliper) below the casing, measured in 2001 after drilling, in NN3 and NN4.

Understanding the fate of H_2S injected in basalts by means of time-domain induced polarization geophysical logging

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¹⁰ Key Points:

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11	•	${\cal H}_2 S$ reinjection in basalt and mineralization into pyrite was monitored using TDIP
12		logging in two of the injection wells
13	•	Chargeability increases observed at +40 days (first monitoring round) correspond
14		to $1-2\%$ pyrite precipitation with great spatial variability
15	•	Subsequent monitoring rounds show a decrease in chargeability, suggesting that
16		pyrite is either passivated or re-dissolved

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17 Abstract

To help meet emission standards, hydrogen sulfide (H_2S) from geothermal production 18 may be injected back into the subsurface, where basalt offers, in theory, the capacity to 19 mineralize H_2S into pyrite. Ensuring the viability of this pollution mitigation technol-20 ogy requires information on how much H_2S is mineralized, at what rate and where. To 21 date, monitoring efforts of field-scale H_2S reinjection have mostly occurred via mass bal-22 ance calculations, typically capturing less than 5% of the injected fluid. While these stud-23 ies, along with laboratory experiments and geochemical models, conclude effective H_2S 24 mineralization, their extrapolation to quantify mineralization and its persistence over 25 time leads to considerable uncertainty. Here, a geophysical methodology, using time-domain 26 induced polarization (TDIP) logging in two of the injection wells (NN3 and NN4), is de-27 veloped to follow the fate of H_2S re-injected at Nesjavellir geothermal site in south-west 28 Iceland. Results show a strong chargeability increase at +40 days, corresponding to pre-29 cipitation of up to 1% in NN4 and 2% in NN3 according to laboratory-based relation-30 ships. A uniform increase is observed along NN4, whereas it is localized below 450 in NN3. 31 Changes are more pronounced with the larger electrode spacing, indicating that pyrite 32 precipitation takes place away from the wells. Furthermore, a chargeability decrease is 33 observed at later monitoring rounds in both wells, suggesting that pyrite is either pas-34 sivated or re-dissolved after precipitating. These results highlight the ability of TDIP 35 logging to monitor pyrite mineralization and have implications for understanding the fate 36 of H_2S upon subsurface storage in basaltic environments. 37

³⁸ Plain Language Summary

High-temperature geothermal production is responsible for air pollution due to hy-39 drogen sulfide (H_2S) present in the magmatic fluid. To help meet emission standards, 40 H_2S may be injected back into the subsurface, where basalt offers the capacity to trans-41 form H_2S into pyrite. Transformation into an immobile mineral prevents further trans-42 port into the atmosphere, sea, surface streams, or lakes. However, ensuring the viabil-43 ity of this pollution mitigation technology requires information such as how much H_2S 44 is mineralized, at what rate and where, which are highly uncertain due to the hetero-45 geneity and inaccessibility of subsurface processes. Here, a geophysical monitoring method-46 ology is developed and tested during the re-injection of H_2S at Nesjavellir geothermal 47 site in south-west Iceland. According to laboratory studies, pyrite precipitation is ex-48 pected to increase the electrical capacitance, "chargeability", of the subsurface. Using the 49 so-called time-domain induced-polarization (TDIP) method embedded in a wireline log-50 ging tool, the chargeability of a 2m-wide cylinder around the injection wells is measured 51 with high spatial-resolution before and during H_2S injection. A strong chargeability in-52 crease at +40 days indicates that up to 2% pyrite is formed. A subsequent decrease at 53 later rounds raises questions on whether pyrite is re-dissolved or passivated by other sec-54 ondary minerals. 55

⁵⁶ 1 Introduction

Geothermal energy production emits an estimated 0.2 Mt/yr of hydrogen sulfide 57 (H_2S) globally, with Iceland alone emitting 30 kt/yr (Marieni et al., 2018). These anthro-58 pogenic H_2S emissions are sourced from the geothermal production steams, which con-59 tain significant concentrations of H_2S (Prikryl et al., 2018; Stefánsson et al., 2011). H_2S 60 emissions are toxic to humans and can be fatal in concentrations as low as 320 ppm (World 61 Health Organization - Regional Office for Europe, 2000). H_2S emissions also pose a threat 62 to the environment, oxidizing when exposed to atmospheric oxygen to form acid rain (Greaver 63 et al., 2012). Recent air quality regulations, both internationally and within Iceland, have 64 limited the amount of H_2S atmospheric emissions to 50-150 µg/m3 per 24 hours (Aradóttir 65

et al., 2015; World Health Organization - Regional Office for Europe, 2000; Iceland Ministry of the Environment, Energy and Climate, 2010).

To reduce emissions from geothermal energy production in Iceland, H_2S can be dis-68 solved into the geothermal wastewater and injected into the basalt subsurface (Mamrosh 69 et al., 2014). To help meet emission standards, H_2S from the geothermal production steam 70 may be captured at the power plant, dissolved into geothermal wastewater, and injected 71 into the basalt subsurface (Mamrosh et al., 2014). Geochemical and reactive transport 72 models suggest that such a H_2S -injection results in effective sulfide mineralization, with 73 74 pyrite being the dominant sulfur bearing mineral (Prikryl et al., 2018; Stefánsson et al., 2011; Marieni et al., 2018; Bacon et al., 2014), due to the high reactivity of basalt and 75 its divalent cations content (up to 25 wt.% Ca, Mg, Fe) (Matter et al., 2016; Gysi & Ste-76 fánsson, 2008). However, these findings are based on simulations and laboratory exper-77 iments with limited evidence from field tests (Robin et al., 2020). In practice, the H_2S 78 mineral storage process requires careful monitoring to identify any adverse effects of the 79 injection, such as the acidification of the shallow groundwater system and the mobiliza-80 tion of toxic metals from the basaltic rocks (Floor et al., 2011; Cuoco et al., 2013; Delmelle 81 et al., 2015; Galeczka et al., 2016; Flaathen & Gislason, 2007). 82

Current monitoring practices for H_2S mineralization into pyrite are similar to those 83 for CO_2 mineralization into carbonates. They consist mainly of quantification by (i) mass 84 balance calculations using tracer tests (Matter et al., 2016), (ii) transport models (Ratouis 85 et al., 2022), and (iii) steady-state "reaction path" geochemical models based on chem-86 ical monitoring. Chemical and mineral analyses on precipitates formed on metallic sur-87 faces within monitoring wells also bring qualitative mineralization evidence (Matter et 88 al., 2016). More recently, the triple sulfur isotope systematics (32S, 33S, and 34S) of geother-89 mal fluids were investigated as a possible tool to trace H_2S sequestration by sulfide min-90 eralization in the geothermal reservoir (Robin et al., 2020). One major pitfall of these 91 monitoring practices is that they typically capture less than 5% of the flow and extrap-92 olate the results (Matter et al., 2016), whereas the remaining 95% may include undetected 93 upward migration through fractures and springs. In addition, a key assumption behind 94 the quantification is that the amount of dissolved gas not captured by the measurement 95 downstream, compared to expectations, has mineralized and that mineralization is per-96 manent (Gunnarsson et al., 2018; Matter et al., 2016). While physical verification of these 97 assumptions, e.g., through post-injection core-drilling, has not yet been possible (Carbfix, 98 2022), other studies focusing on CO_2 indicate that additional processes to carbonate min-99 eralization can happen, namely the formation of unstable organic carbon due to micro-100 bial activity (Trias et al., 2017). Therefore, quantifying mineralization with these meth-101 ods, whether sulfide or carbonate, leads to considerable uncertainty (White et al., 2020). 102

Time-lapse geophysics is a group of potentially suitable methods to obtain in-situ 103 information on the spatial distribution and magnitude of H_2S mineralization, as well as 104 its evolution over time. Time-lapse geophysics as a monitoring technique can, in theory, 105 capture dynamic processes with high spatial and temporal resolutions (Hermans et al., 106 2023; Lévy et al., 2022). Geo-electrical methods are particularly suitable for pyrite de-107 tection. Underground massive metallic ore deposits, as well as veinlets and disseminated 108 sulfides, have been detected and discriminated with induced polarization (IP) field mea-109 surements (Pelton et al., 1978; Placencia-Gómez, 2015; Börner et al., 2018). Petrophys-110 ical laboratory measurements also indicate that the presence of pyrite can be detected 111 in altered volcanic rocks and quantified to some extent, with induced polarization (Lévy, 112 Gibert, et al., 2019). The polarizability, also called chargeability, describes the ampli-113 tude of electrical polarization in rocks, which is particularly strong at interfaces between 114 fluid and semi-conductors, such as pyrite or magnetite (Bücker et al., 2018; Gurin et al., 115 2015; Abdulsamad et al., 2017). The reorganization of charges at the interface between 116 pore fluid and metallic particles creates local electrical "displacement" currents, which 117 add up until a maximum voltage is reached. The time-delay between maximum voltage 118

and maximum current, also known as the phase-angle for sinusoidal signals, is caused 119 by the polarization (Bücker et al., 2018; Chelidze & Gueguen, 1999; Olhoeft, 1985). On 120 the other hand, the conductivity quantifies the amplitude of the electric conduction phe-121 nomenon in rocks, which is in particular enhanced by the presence of smectite, as well 122 as high porosity, salinity, and temperature (Flóvenz et al., 2005; Kristinsdóttir et al., 2010; 123 Lévy et al., 2018; Waxman & Smits, 1968). Geo-electrical field investigations carried out 124 at a geothermal site in North-East Iceland confirmed that the conductivity and IP sig-125 nals of naturally present pyrite, smectite, and magnetite can be picked up down to 200 126 m depth (Lévy, Maurya, et al., 2019). 127

In January 2021, the geothermal power company ON Power started a small-scale 128 injection of H_2S and CO_2 dissolved in seal water from liquid-ring vacuum pumps at Nes-129 javellir, South-West Iceland. This small-scale injection runs alongside larger-scale injec-130 tion of separated geothermal fluid from the power plant that has been part of standard 131 operations since 1990, shortly after the power plant was installed. The injection takes 132 place in eight shallow wells ranging from 311 m to 660 m deep. We use this injection ex-133 periment to investigate the subsurface induced polarization response over time and as-134 sess (i) the sensitivity of time-lapse geophysics to pyrite precipitation and (ii) the sus-135 tainability of pyrite mineralization. As surface geophysical measurements lose resolution 136 with depth, wireline logging was considered a relevant intermediate scale between lab-137 oratory and surface geophysical measurements to assess pyrite precipitation in the vicin-138 ity of the injection wells. It is thus the focus of this study. 139

¹⁴⁰ 2 Theory and Background

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2.1 Electrical response of pyrite with frequency-domain IP (FDIP)

In an electrical context, geological formations can be thought of as a combination 142 of resistors and capacitors (Lévy, 2019). Ions in pore water and those connected to clay 143 minerals are charge carriers in aquifers. Conductive water and clay minerals can be rep-144 resented by conductors with high conductivity, as opposed to freshwater and pure quartz 145 sand or gravel, which have lower conductivity. Semi-conducting metallic particles (e.g., 146 pyrite or magnetite) can cause electronic conduction beyond 20% per volume if the par-147 ticles are interconnected (e.g., veinlets) (Slater et al., 2005, 2006; Revil, Abdel Aal, et 148 al., 2015), but disseminated particles mainly cause electrical polarization, an effect sim-149 ilar to that of a capacitor in an electric circuit. Polarization caused by semi-conductors 150 is usually several orders of magnitude stronger than sediment/water interfaces (Revil et 151 al., 2017; Slater et al., 2005, 2006). This is due to the specifics of the crystal structure 152 of semi-conducting minerals, such as pyrite and magnetite, and the possibility for elec-153 trons and holes to "jump" and rapidly redistribute within the metallic particle in response 154 to the external field (Shuey, 2012; Pridmore & Shuey, 1976). This redistribution causes 155 the accumulation of ions at the interface between fluid and metallic particles. 156

Figure 1 illustrates the large polarization, represented by the phase angle parameter, of volcanic samples containing pyrite, as opposed to a volcanic sample of similar geology and full of smectite but without pyrite; the latter being more conductive but much less polarizable, as described in details by Lévy, Gibert, et al. (2019) and Lévy, Weller, and Gibert (2019). It is also visible in Figure 1 that the grain size distribution of metallic particles affects the frequency at which the phase angle peak occurs.

Several models have been developed to describe how ions reversibly accumulate at the interface between pore fluid and metallic particles, causing polarization. One of the most widespread phenomenological relaxation models to interpret FDIP data is the Pelton model (Pelton et al., 1978; Tarasov & Titov, 2013; Weller & Slater, 2022), given by equation 1.



Figure 1. Laboratory results for naturally altered volcanic samples from the Krafla geothermal field (north-east Iceland) containing varying amounts of pyrite with different grain sizes. Top panels: FDIP data for four different samples, left = impedance modulus, right = phase angle, as a function of frequency. Middle panels: scanning electron microscope (SEM) images for the three samples containing pyrite. Pyrite grains are highlighted in red. Bottom panels: distribution of pyrite grain size, based on the SEM images of the three samples.

$$\rho^*(f) = \rho_{DC} \left[1 - m\left(1 - \frac{1}{1 + (i2\pi f\tau)^c}\right)\right] \tag{1}$$

with four fitting parameters: the DC resistivity ρ_{DC} , the chargeability m, the Pelton time constant τ and the so-called Cole–Cole exponent c.

However, the Pelton model, presented in equation 1, is only valid for spectra measured on samples with a single grain size fraction. Variations of this model for two grain
size fractions exist but the fitting parameters are not directly comparable to each other,
and even the two grain size fractions model may not represent the complexity of naturally occurring pyrite crystals in volcanic rocks (Lévy, Gibert, et al., 2019).

A more general approach to analyzing FDIP data is the Debye decomposition (Nordsiek Weller, 2008). With this approach, the frequency-dependent complex electrical resistivity $\rho^*(f)$ describes an IP spectrum that can be fitted by a superposition of N Debye terms (Pelton models with c = 1), as shown in equation 2.

$$\rho^*(f) = \rho_{DC} \left[1 - \sum_{k=1}^N m_k \left(1 - \frac{1}{1 + i2\pi f \tau_k}\right)\right]$$
(2)

where m_k and τ_k are pairs of partial chargeability and relaxation time of a single Debye model. The function $m_k = f(\tau_k)$ is also referred to as relaxation time distribution (RTD). The N chargeability values m_k , which are related to a set of pre-defined relaxation times τ_k , and the value of DC resistivity ρ_{DC} are the result of a fitting procedure described by Nordsiek and Weller (2008). Two parameters can be integrated from the Debye decomposition to simplify the interpretations: the total chargeability m_{tot} (equation 3) and the mean relaxation time and τ_{mean} (equation 4).

$$m_{tot} = \sum_{k=1}^{N} m_k \tag{3}$$

$$\tau_{mean} = exp(\frac{\sum_{k=1}^{N} m_k ln(\tau_k)}{m_{tot}}) \tag{4}$$

The main empirical petrophysical relationship linking the chargeability to pyrite volume in the absence of background polarization was developed by Revil, Florsch, and Mao (2015) (equation 5).

$$m = \frac{9}{2}p_v \tag{5}$$

where p_v is the volume fraction of pyrite (or other metallic particles) and m is the chargeability in V/V (dimensionless). In their study, the chargeability parameter is fitted to frequency-domain IP (FDIP) data by a Pelton model (Cole & Cole, 1941; Pelton et al., 1978). According to Martin and Weller (2023), the chargeability m fitted by a Pelton model (Pelton et al., 1978) and m_{tot} fitted by Debye decomposition (Nordsiek & Weller, 2008) are similar for sand-pyrite mixtures.

While the influence of temperature on the polarization effect is still poorly understood, recent laboratory experiments indicate that the phase angle in the presence of metallic particles is temperature independent in the range 5-50°C (Revil et al., 2018). 2.2 Frequency-domain and time-domain induced polarization (FDIP and TDIP)

In the field, the Time Domain Induced Polarization (TDIP) method is often preferred to FDIP, primarily due to its easier and faster implementation. TDIP consists of injecting a direct current pulse of finite duration (DC) through two current electrodes (A and B) and recording the resulting voltage at two potential electrodes (M and N) with a fine-enough sampling rate to capture the exponential charge and discharge of the voltage, during the on-time and off-time, respectively (Sumner, 1976; Telford et al., 1990). Further details on the TDIP methods can be found in Binley and Slater (2020).

In general, voltage discharge measured by TDIP is expected to have a stretched 207 shape, which describes the relaxation behavior in a similar manner as Cole-Cole and other 208 related FDIP models (Alvarez et al., 1991; Cole & Cole, 1941; Davidson & Cole, 1951; 209 Havriliak & Negami, 1966). More details are described in Appendix Appendix A. While 210 quantitatively connecting the TDIP and FDIP responses of rocks containing metallic par-211 ticles is not trivial (Alvarez et al., 1991), the Debye decomposition is a common method 212 for analyzing both TDIP and FDIP data (Nordsiek & Weller, 2008; Tarasov & Titov, 213 2007). An equivalence was found by Martin et al. (2021) between the Debye decompo-214 sition of FDIP and TDIP data, provided that the current injection time in TDIP was 215 long enough (ideally 64 seconds, but decent comparisons were found beyond 2 seconds). 216 With 2-second current injection, as is the case in this study, the coefficients from the De-217 by decomposition of TDIP data are slightly underestimated compared to FDIP data. 218 Despite this limit, this approach was considered the most relevant one. As opposed to 219 the integral chargeability, which tends to smoothen out noise in the data, the Debye de-220 composition tends to amplify the noise and, thus, requires careful data processing (Martin 221 et al., 2021). 222

- ²²³ 3 Materials and Methods
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3.1 H_2S production and re-injection at Nesjavellir geothermal field, Iceland

The methodology developed in this study for monitoring H_2S injection was tested 226 at the Nesjavellir geothermal field (SW Iceland), where a geothermal power station has 227 been operated since 1990 (Figure 2). Nesjavellir is a high-temperature geothermal field 228 $(> 200^{\circ}C \text{ above 1 km depth})$ characterized by a relatively high permeability (Zakharova 229 & Spichak, 2012). Nesjavellir is located just north of the Hengill central volcano, which 230 is situated on a ridge-ridge-transform triple junction (Foulger, 1988). The area is com-231 prised of mainly late Quaternary hyaloclastites and post-glacial age hyaloclastites and 232 basalt flows (Árnason et al., 1969; Foulger & Toomey, 1989). This study focuses on shal-233 low depths northeast of the geothermal production area and above the low-permeability 234 clay cap layer (approximately 500 m depth) (Schiffman & Fridleifsson, 1991; Gómez-Díaz 235 et al., 2022; Gunnarsdottir et al., 2020; Franzson & Gunnlaugsson, 2020). Intrusions are 236 rare at these shallow depths, and none have been identified in the upper 300 m (Franzson, 237 1988). 238

The Nesjavellir power station currently produces 120 MWe of electricity and 300 239 MWth of thermal energy for district heating. The hydrothermal production fluid (260-240 300°C) is sourced at 1000-1500 m depth (Snæbjörnsdóttir et al., 2020). Following a con-241 trolled pressure-decrease, the hydrothermal fluid is divided into (i) steam and (ii) geother-242 mal separated water at 192°C. Electricity generation at Nesjavellir uses 240 kg/s of steam 243 to power four turbines. The steam is then condensed, using 2,000 l/s of cold ground-244 water (5-7°C), to form condensate wastewater, composed primarily of distilled water. 245 In parallel, heat exchangers use the $192^{\circ}C$ geothermal separated water to heat cold ground-246 water to 87°C, which is then pumped to the capital Reykjavík, 27 km away. The sep-247



Figure 2. Aerial view of Nesjavellir valley (south-west Iceland). The white building shows the geothermal power plant. The two wells, NN3 and NN4, investigated in this study are indicated by white circles, while other wells in the area are shown with small black rectangle signs. A buried power line (132 kV), most likely responsible for the 50 Hz background noise, is shown in blue. Pipelines are shown in pink and roads in black.

Table 1. Details on the Nesjavellir NN injection wells in this study. The borehole fluid conductivity was measured with a QL40-FTC logging probe (*Mount Sopris QL40-FTC*, 2014) in July 2022 (+540 days). Temperature and caliper logs are presented in Figures B1 and B2 in Appendix Appendix A.

Well ID	NN-3	NN-4
Coordinates	64°06'38.23" N, 21°15'20.27" W	64°06'50.03" N, 21°15'0.06" W
Depth (m)	563	422
Deviation	0°	0°
Casing Depth (m)	205	201
Borehole mean diameter (mm)	405 (min=317 - max=680)	403 (min=324 - max=636)
Average flow Rate (kg/s)	11	113
Injected fluid temperature (°C)	65	85
Borehole temperature while measuring (°C)	63-69	57-71
Reservoir temperature (°C)	15-20	20-40
Borehole fluid conductivity (mS/m)	60-100	100
pH	8.5	9.1
H_2S (ppm)	74.42	76.81
$CO_2 (\text{ppm})$	13.5	8.35
SiO_2 (ppm)	435.08	428.09

arated water contains the dissolved solids from the hydrothermal fluid and thus has a 248 high potential for chemical pollution and corrosion. Starting in 2004 and prior to 2021, 249 wastewater composed of condensate and separation water has been disposed of in drainage 250 wells. This initial wastewater had negligible concentrations of dissolved CO_2 and H_2S . 251 Starting on January 29th, 2021, a third type of fluid, called seal water, is now mixed into 252 the condensate and separation water before injection. Seal water contains the non-condensable 253 fraction of gases, primarily H_2S , that pose corrosion problems to turbines over time. It 254 is formed by dissolving the CO_2 and H_2S gases from liquid ring vacuum pumps at the 255 condensers in cold groundwater or condensate water. The currently injected wastewa-256 ter contains, on average, 75 ppm of H_2S (Table 1), with the intention to sequester H_2S 257 through mineral storage. 258

Two injection wells are used for monitoring: NN3 and NN4, drilled into fresh lavas 259 to 563 m (NN3) and 422 m (NN4) depth in 2001, northeast of the power station. Their 260 specific lithology and alteration stage at the time of drilling (i.e., prior to large-scale warm 261 wastewater injection) are presented in Helgadóttir (2021), emphasizing an increase in the 262 amount of clay minerals below 400 m in NN3. Between 2004 and 2021, injection of the 263 initial wastewater into the cold groundwater system took place through these wells and 264 has probably been enhancing basaltic rock alteration compared to natural conditions. 265 Starting from January 2021, H_2S has been continuously injected through the final wastew-266 ater. Reaction path modeling from the literature predicts that basaltic glass dissolution 267 is the main source of iron over short time durations, with magnetite dissolution contribut-268 ing less iron due to the slower dissolution kinetics and more limited amount. Pyrite pre-269 cipitation is expected to be rapid (Prikryl et al., 2018; Stefánsson et al., 2011). An overview 270 of the area, including roads, boreholes, and buried infrastructure, is presented in Fig-271 ure 2. Information on the wells and the injection are given in Table 1. Temperature logs 272 in NN3 and NN4 measured the days of TDIP logging are presented in Figures B1 in Ap-273 pendix Appendix B, together with caliper logs measured after drilling (Figure B2). Fur-274 ther details on the wells can be found in Gómez-Díaz et al. (2022) and Hafstað (2003). 275

3.2 Field monitoring with QL40-IP logging tool

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The QL40-ELOG/IP (Mount Sopris QL40-IP, 2020) logging tool measures the elec-277 trical resistivity and time-domain IP (TDIP) response with electrodes made of stainless 278 steel. The tool has a diameter of 43 mm and uses the "normal" electrode configuration 279 with 16- and 64" electrode spacings, defined as the distance between the current elec-280 trode (A) and the potential electrode (N) (Helander, 1983). The current generator, built 281 into the tool, sends current at electrode A in all directions and the shielding of the log-282 ging wireline serves as the current sink. The reference potential electrode (M) is located 283 on top of an 8m-long isolation bridle, such that the distance between the potential elec-284 trodes (M and N) is considered infinite in comparison to the much smaller electrode spac-285 ing AN. Moreover, grounding of the logging unit (truck) is achieved by clamping the truck 286 ground onto the casing for security reasons (Advanced Logic Technology, 2021). The 64" 287 measurements are sensitive to polarization effects over a larger measurement volume, deeper 288 into the formation, compared to the 16". The radial investigation characteristic, as de-289 fined by Roy and Dhar (1971), considers electrostatic potentials of individual cylindri-290 cal shells of varying radii integrated to obtain the total contribution to the measured sig-291 nal. According to the formula by Roy and Dhar (1971), 75% of the measured signal with 292 the 64" spacing is caused by the cylindrical volume comprised in a 2.5-m radius from the 293 borehole central line. As a comparison, 75% of the measured signal with the 16" spac-294 ing comes from within 0.5 m, which is about twice the boreholes' mean radius (Table 295 1). Here, measurements took place every 25 cm, with a logging speed of 1.8 m/min. 296

A total of eight measuring rounds took place but only four measuring rounds are presented in this study for NN-3 and NN-4 due to mainly one problem encountered onsite, related to the inability to inject current higher than 200 mA, happening randomly at certain places and dates (Table 2).

Traditional applications of the QL40-IP tool, e.g. in the mining industry, are typ-301 ically fulfilled with 250 or 500 msec current injections (on-time) and a fixed number of 302 400 sample points in the whole cycle $T_{on,+}; T_{off,+}; T_{on,-}; T_{off,-}$. Two important differ-303 ences between traditional applications and the present study led to the development of 304 a new processing board. First, Martin et al. (2021) show that FDIP and TDIP data mea-305 sured on the same samples decently overlap when current injections last at least 2 sec-306 onds during TDIP data acquisition. This overlap is key to quantifying pyrite volumes 307 by Debye decomposition since these empirical relationships have only been established 308 with FDIP data. Second, the 132 kV buried power line connected to the Nesjavellir power 309 plant (Figure 2) is responsible for a strong 50 Hz signal in the TDIP data, as revealed 310 by initial tests in NN-4 prior to this monitoring study (December 2019). These tests were 311 carried out with the original QL40-IP tool, using 250-msec and 500-msec current injec-312 tions, corresponding to full cycles of 1 and 2 seconds, respectively, and thus sampling rates 313 of 400 and 200 Hz (400 sample points over the whole cycle). Using 2-second current in-314 jections, the full cycle lasts 8 seconds, which leads to a sampling rate of 50 Hz with 400 315 sample points and aliases the 50 Hz noise. Further investigations also showed that both 316 the amplitude and exact frequency of the "50 Hz" noise varied over time. Therefore, it 317 was clear that TDIP data acquisition with 2-second current injection required a larger 318 number of sample points to model the noise and remove it before data analysis. This was 319 achieved by increasing the memory of the processing board and led to an updated ver-320 sion of the QL40-IP tool used in the rest of this study. 321

From September 2020 (baseline) and on, the following IP acquisition settings were used: square-wave current injection $(T_{on,+}; T_{off,+}; T_{on,-}; T_{off,-}]$ for 2 seconds each, 8 seconds total), 450 Hz sampling rate of the voltage during the whole cycle (i.e. 3600 sample points), and wireline speed of 1.8 m/min. The 50 Hz noise is further illustrated in Figure 3 (first and second columns).



Figure 3. Full waveform (FW) TDIP data measured in NN3 with 16" and 64" electrode spacings (left column). The 50 Hz noise is further illustrated by zooming on the positive decay for the 64" data in the second column (signal in black), where the FW signal after removing the 50 Hz noise is also shown (in red). Further processing of the FW data into polarizability curves is shown in the third (linear scale) and fourth (log-log scale) columns. Results are shown at two different dates (baseline and +40 days) and two different depths (263 m and 470 m), showing varying levels of background noise.

Measuring date	NN-3	NN-4
December 2019 (preliminary)	no meas.	404 sample points
September 2020 (baseline)	good	good
March 2021 $(+40 \text{ days})$	good	good
November 2021 ($+270$ days)	good	current too low
February 2022 $(+380 \text{ days})$	current too low	good
July 2022 $(+540 \text{ days})$	good	good
October 2022 $(+630 \text{ days})$	current too low	current too low

 Table 2.
 TDIP monitoring timeline and evaluation of data quality. "Good" means data quality was sufficient for further analysis.

In parallel to the TDIP monitoring, the changes in electrical resistivity were also monitored to evaluate the possible contribution of processes other than pyrite precipitation to the changes in the IP response. Indeed, changes in borehole fluid composition, temperature, and precipitation of clay minerals are reflected by changes in resistivity. Similarly, the formation of connected clusters of pyrite particles that would lose their polarization properties while becoming more conductive would also cause changes in the resistivity logs.

3.3 Data processing from full waveform voltage signal to polarizability decays

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The apparent chargeability $M_{app}(t)$, also called polarizability and noted $\eta(t)$ in Martin et al. (2021) and Tarasov and Titov (2007), corresponds to the ratio between the decaying voltage during the off-time, $V_{decay}(t)$, and the maximum voltage reached during the on-time, V_{DC} . It is calculated by stacking positive and negative decays (equation 6).

$$M_{app}(t) = \eta(t) = \frac{V_{decay}(t)}{V_{DC}} = \frac{1}{2} \left(\frac{V_{decay,+}(t)}{V_{DC,+}} + \frac{V_{decay,-}(t)}{V_{DC,-}} \right)$$
(6)

A series of processing steps are carried out to clean the full waveform signal, optimize the signal-to-noise ratio, and maintain three time-decades of signal for the polarizability. The term polarizability is used in the rest of the paper to refer to the apparent chargeability defined in equation 6 in order to avoid confusion with the integral chargeability or the total chargeability parameter calculated from Debye decomposition.

First, a time-invariant self-potential (SP) voltage affects both V(t) and V_{DC} in the positive and negative decays. The SP offset at each depth and each date is calculated by averaging the positive and negative DC voltages, $V_{DC,+}$ and $V_{DC,-}$ and removed from V(t) and V_{DC} .

Second, harmonics of 50 Hz noise, due to a 132 kV buried power line, affect the volt-349 age full waveform measurements, as mentioned in the previous section and illustrated 350 in Figure 3. A power line noise model, based on the algorithm proposed by Larsen et 351 al. (2022) and applied in a similar manner by Olsson et al. (2016), is used to subtract 352 the harmonic noise from the signal. As shown by Larsen et al. (2022), the fundamen-353 tal frequency varies over time, rapidly oscillating between 49.9 and 50.1 Hz, and must 354 first be determined. The 2-second decaying voltage is thus decomposed into 200 msec 355 segments, where the frequency is fitted in every segment. Once the exact frequency is 356 determined, a model of the background noise, including harmonics up to Nyquist fre-357 quency, is removed from the signal. The denoised voltage after applying this 50 Hz fil-358 ter is presented in Figure 3 (panels in the second column) as the red curve on top of the 359 FW voltage (in black) during the positive off-time section of the full cycle. 360

Third, the voltage decays are re-gated into log-spaced time windows. This step trans-361 forms the full waveform voltage decay signal with 900 linearly spaced data points into 362 a gated voltage signal with 37 logarithmically spaced gates. More accurately, the re-gated 363 voltage consists of 10 gates in the first 22 msec (2.21 msec width, i.e., the original sam-364 pling step) and then exponentially larger gate widths (from 4.42 to 141.44 msec), over 365 which an exponentially increasing number of data points (from 2 to 64) are averaged. 366 Given the strong exponentially decaying shape in the early times, it appeared best to 367 keep all data points in the first 22 msec. Since the original data points are kept as the 368 10 first gates, it corresponds to a "square-gate" with one data point averaged. To be con-369 sistent, the square gating strategy is kept throughout the decay. This is especially rel-370 evant because the signal can be locally approximated by linear functions in the later times. 371 At some depths, the first gate (centered at 1.1 msec) is affected by electromagnetic (EM) 372 coupling (see, for example, at +263 m in Figure 3). Therefore, it was discarded from all 373 the re-gated discharge curves, and the first effective gate is at 3.32 msec: visual inspec-374 tion of all the decays suggests that EM coupling is negligible at this gate, which is con-375 sistent with observations by Olsson et al. (2016). 376

Fourth, the positive and negative voltage decays are divided by the DC voltage and 377 stacked to obtain the polarizability in mV/V (equation 6). The stacking allows remov-378 ing extra noise with the same sign in the positive and negative decays. These four pro-379 cessing steps eventually provide the processed polarizability curves over almost three decades 380 (in the range of 3-2000 msec), as presented in Figure 3 with a linear scale (third column) 381 and log-log scale (fourth column). The linear scale allows better comparison to the full 382 waveform signal presented in the same figure, while the log-log scale is used in the rest 383 of this study. 384

Fifth, polarizability decays are averaged over 2-meter thick sections (Figure 4), cor-385 responding to 8 decays averaged together (0.25 m spatial resolution). This is done to fa-386 cilitate the comparison between measuring rounds since the exact depth of each single 387 measurement varies at different dates. It also allows smoothing out short-wavelength sig-388 nal oscillations that would prevent meaningful comparison between measuring rounds. 389 The standard deviation corresponding to this average also captures noisy areas where 390 the signal varies a lot over short distances. The averaging over 2-meter sections also takes 391 into consideration the fact that logging measurements at one depth are, in reality, sen-392 sitive to a sphere around this given depth. Finally, averaging over 2-meter sections helps 393 to identify and mitigate remaining harmonic noise. 394

Some polarizability curves remain noisy after the processing (non-monotoneously or non-exponentially decaying), e.g., at 263 m depth and +40 days shown in Figure 3 and at 382-384m and +40 days shown in Figure 4. Noisy processed polarizability curves are further addressed in the next subsection and discussed at the end of the paper.

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3.4 Estimations of pyrite precipitation from polarizability decays

A simple way to represent TDIP data (polarizability decays) as a function of depth is to integrate the decays to obtain the so-called "integral chargeability", M_{int} (Mao et al., 2016; Telford et al., 1990). It represents the area enclosed by the discharge curve, V(t), and its zero asymptote, in a given time-window, [t1:t2], divided by the primary voltage V_{DC} and the time-window width, as described by equation 7 (Bertin & Loeb, 1976; Sumner, 1976). An error on M_{int} , $Err(M_{int}9$, is calculated following equation 8 at every date d.

$$M_{int} = \frac{1}{t_2 - t_1} \int_{t_1}^{t_2} \frac{V(t)}{V_{DC}} dt$$
(7)



Figure 4. Resulting polarizability decays after the five processing steps. An example of noisy decay in NN3 is shown at 382-384 m depth, with larger error bars and a non-exponentially decaying shape.

$$Err(M_{int}(d)) = \frac{M_{int}(d)}{\overline{\eta}} \sqrt{\frac{1}{n} \sum_{i=1}^{n} std(\eta_i)^2}$$
(8)

where $std(\eta_i)$ is the standard deviation of the polarizability of the *i*-th gate within the averaged 2-meter thick section, *n* is the number of time gates in the polarizability decays and $\overline{\eta}$ is the average polarizability over the *n* gates.

The integral chargeability presents the advantage of smoothing out the noise. It is calculated here in the time interval ranging from the second gate (3.32 msec) to the last gate (1926 msec). M_{int} is used in this study as a qualitative assessment tool of the change in polarization over time at a given depth, the trend corresponding to pyrite formation or dissolution. The error bar of the relative difference Δ_{rel} is calculated with equation 9 for the integral chargeability M_{int} and the resistivity ρ .

$$Err(\Delta_{rel}M_{int}(d)) = \sqrt{(M_{int}(d) \cdot \frac{Err(M_{int}(d_0))}{M_{int}(d_0)^2})^2 + (\frac{Err(M_{int}(d))}{M_{int}(d_0)})^2}$$
$$Err(\Delta_{rel}\rho(d)) = \sqrt{(\rho(d) \cdot \frac{Err(\rho(d_0))}{\rho(d_0)^2})^2 + (\frac{Err(\rho(d))}{\rho(d_0)})^2}$$
(9)

where d and d_0 correspond to the current monitoring date and the baseline date, respectively, $Err(\rho(d))$ is the standard deviation within the averaged 2-meter thick section, and $Err(M_{int}(d))$ is calculated as per equation 8.

The integral chargeability calculated from TDIP data cannot be converted to pyrite 419 volume as M_{int} is not equivalent to the "Cole-Cole" or "Pelton" m in equation 5. Sev-420 eral approaches have been developed to relate TDIP and FDIP data, considering there 421 is no strict equivalence between Pelton models in frequency-domain and stretched ex-422 ponential functions in time-domain (Alvarez et al., 1991). As described in section 2, there 423 is an equivalence between the Debye decomposition calculated with FDIP and TDIP data, 424 with a slight underestimation of m_{tot} from TDIP data when the current on-time is 2 sec-425 onds (Martin et al., 2021). Therefore, the conversion to pyrite volume fraction is car-426 ried out here by using Debye decomposition. The polarizability $M_{app} = \eta(t)$ is fitted 427 by a non-negative linear least-square procedure, where the fitting function is given in equa-428 tion 10 and the optimization problem is expressed in equation 11. 429

$$\eta(t) = \sum_{k=1}^{N} m_k e^{\frac{-t}{\tau_k}} \tag{10}$$

$$\min_{x \ge 0} \|Cx - d\|_2^2 \tag{11}$$

where x is a vector with the m_k model parameters, d is a vector with the measured data $\eta(t_i)$ and C is the linear multiplier defined as $C_{i,k} = e^{\frac{-t_i}{\tau_k}}$, where t_i are the center times of the p gates (i=1..p) and τ_k are the pre-defined N relaxation times (k=1..N). Here we define the $\tau_{k,k=1..N}$ as a set of N = 300 logarithmically-spaced values of relaxation times in the range 3-2000, covering approximately three decades. During the procedure, most of the coefficients m_k are kept to 0 due to the non-negative constraints. The fit eventually results in 10-20 coefficients being non-zeros.

⁴³⁷ The total chargeability m_{tot} is calculated by summing all the m_k coefficients (equa-⁴³⁸ tion 3). The difference of m_{tot} at different measuring rounds, Δm_{tot} , is converted into ⁴³⁹ a difference in pyrite volume fraction p_v , Δp_v , following equation 12, which is based on ⁴⁴⁰ equation 5.

$$\Delta p_v(d)_{\%} = \frac{200}{9} [m_{tot}(d) - m_{tot}(d_0)]$$
(12)

where d and d_0 are the monitoring date and the baseline, respectively. Only the difference is converted, not the absolute pyrite volume fraction, in order to subtract the contribution of the background polarization that may be partly due to iron-oxides or other polarizable material present before the start of H_2S injection.

As opposed to the integral chargeability, the Debye decomposition seems to act as 445 a noise amplifier, similar to observations by Martin et al. (2021). As mentioned previ-446 ously, some polarizability curves remained noisy despite the numerous processing steps, 447 with a resulting non-decaying shape. Such curves should not be processed with Debye 448 decomposition. An additional processing step is introduced to ensure that only exponentially-449 decaying curves are processed with Debye decomposition. The criterion for ensuring this 450 shape is based on the Kohlrausch-Williams-Watts relaxation model for TDIP data (Alvarez 451 et al., 1991), presented in Appendix Appendix A. In practice, the polarizability curves 452 $\eta(t)$ are fitted with a stretched exponential function, presented in equation 13, before ap-453 plying the optimization procedure described in equations 10 and 11. 454

$$f(t) = \alpha e^{-\left(\frac{t}{\tau_{KWW}}\right)^{\beta}} \tag{13}$$

The fitting parameters α , β , and τ_{KWW} , further described in Appendix Appendix 455 A, are not used for interpretation in this study. Instead, the stretched exponential fit serves 456 two purposes: (i) calculating a deviation to a stretched exponential to remove outliers 457 polarizability curves where $R^2 < 0.998$ and (ii) smoothing out the polarizability decays 458 that are reasonably deviating to ensure the Debye decomposition is not fitting noise. In 459 order to confirm that this fitting procedure is not introducing a significant bias, the m_{tot} 460 and τ_{mean} resulting from Debye decomposition before and after the fitting procedure are 461 compared in Figures A1 and A2 in Appendix Appendix A. While other types of curves 462 may reflect IP effects, with e.g. negative polarizability values having a physical expla-463 nation in the context of dipole-dipole acquisition with sharp layer boundaries (Fiandaca 464 et al., 2022), the polarizability curves discarded here did not belong to these "hetero-465 dox transients", but rather had a distorted shape, similar to the curve in NN3 at +40466 days at 382-384 m depth, presented in Figure 4. 467

An error is calculated on the resulting values of m_{tot} (equation 3), taking into ac-468 count both the residuals to the stretched exponential fit and the standard deviation within 469 the averaged 2-meter sections, to help discriminate between variations that fall within 470 an uncertainty interval from significant variations over time. First, an error is calculated 471 for each gate of the polarizability curve as the sum of the 2-meter standard deviation 472 (corresponding to the variation within the averaged 2-meter section, see also equation 473 8) and the residuals to the stretched exponential fit (equation 14): if the residual is pos-474 itive, it is added to the positive error bar err_+ , if the residual is negative, it is added to 475 the negative error bar err_{-} . 476

$$Err_{+}(\eta_{i,fit}) = \begin{cases} std(\eta_{i}) + residuals(\eta_{i}, \eta_{i,fit}), & \text{if residuals}(\eta_{i}, \text{fit}) > 0\\ std(\eta_{i}), & \text{otherwise} \end{cases}$$
$$Err_{-}(\eta_{i,fit}) = \begin{cases} std(\eta_{i}) - residuals(\eta_{i}, \eta_{i,fit}), & \text{if residuals}(\eta_{i}, \text{fit}) < 0\\ std(\eta_{i}), & \text{otherwise} \end{cases}$$
(14)

where $\eta_{i,fit}$ is the value of the stretched exponential fit of the polarizability at the i-th gate, and Err_+ and Err_- are the positive and negative error bars on $\eta_{i,fit}$ at each gate.

Similarly to the error on M_{int} , a relative error is calculated for each decay using the root mean square of the error on all the gates divided by the average polarizability $\overline{\eta}$. This relative error is then applied to the total chargeability m_{tot} , then to Δm_{tot} (change from the baseline), and finally to the pyrite volume fraction change Δp_v , following equation 15.

$$Err_{rel} = \frac{1}{\overline{\eta}} \sqrt{\frac{1}{n} \sum_{i=1}^{n} \left(\frac{Err_{+}(\eta_{i,fit}) + Err_{-}(\eta_{i,fit})}{2}\right)^{2}}$$
$$Err(m_{tot}) = Err_{rel} \cdot m_{tot}$$
$$Err(\Delta m_{tot}(d)) = \sqrt{Err(m_{tot}(d))^{2} + Err(m_{tot}(d_{0}))^{2})}$$
$$Err(\Delta p_{v}(d)) = \frac{2}{90} \cdot Err(\Delta m_{tot}(d))$$
(15)

485 4 Results

486

4.1 Baseline resistivity and integral chargeability

In order to evaluate the amplitude and meaning of the changes in the following sec-487 tions, the resistivity and integral chargeability measured in the baseline are first presented 488 in Figure 5. The average resistivity of NN3 and NN4 is about 30 Ωm , with a clear thresh-489 old at 450 m in NN3, separating the well into two sections, above and below 30 Ωm . Both 490 in NN3 and NN4, a few intervals have resistivity values significantly higher than 30 Ωm , 491 up to 200 Ωm . These high-resistivity intervals coincide with low-chargeability intervals 492 (gray rectangles in Figure 5). The average integral chargeability is around 20-30 mV/V, 493 with generally higher values in NN4. 494

Intervals of higher resistivity correspond to more compact, less porous, litholog-495 ical layers, where less hydrothermal fluid flow and thus alteration is expected (Lévy et 496 al., 2018). Therefore, the relatively high chargeability in the baseline, as well as the co-497 incidence of locally high resistivity with locally low chargeability, suggest that hydrother-498 mal alteration had been occurring in the wells prior to the start of H_2S injection, espe-499 cially in intervals of low resistivity. Small amounts of magnetite are expected in unal-500 tered zones, typically 2-4 wt.% (Lévy et al., 2018) and may contribute to the chargeabil-501 ity in high resistivity layers (Lévy, Gibert, et al., 2019; Peshtani et al., 2022). Alteration 502 minerals, such as pyrite and smectite, are especially expected in permeable, fractured 503 layers (Lévy, Gibert, et al., 2019; Gudmundsson et al., 2010; Lévy et al., 2020) and may 504 be responsible for the higher chargeability in low resistivity intervals, particularly be-505 low 450 m in NN3 and below 375 m in NN4. Note that smectite only contributes to po-506 larization when present in small quantities, typically lower than 10% (Lévy, Weller, & 507 Gibert, 2019; Telford et al., 1990; Mendieta et al., 2021), while it tends to decrease the 508 overall polarization response at larger quantities (Lévy, Gibert, et al., 2019), as illustrated in Figure 1. Based on these observations, the interval below 450 m in NN3 is where ex-510 tensive pyrite precipitation is expected following H_2S injection, which is also consistent 511 with the observation that most of the "natural" hydrothermal alteration happens be-512 low 450 m in the area (Helgadóttir, 2021). 513

In both wells, higher chargeability is observed with the 64" spacing compared to the 16". Considering that the fluid is not chargeable in this frequency-range, this illustrates that the 64" images more volume of the chargeable properties of the rock formation, and thus more changes are also expected with the 64" spacing. On the other hand,



Figure 5. Resistivity (blue/red) and integral chargeability (green/black) logs in NN3 (left panels) and NN4 (right panels) using the 16- and 64" spacing (colors). All plots are based on an averaged signal every 2 meters, and the error bars correspond to the standard deviation within these 2-meter intervals. The dashed line shows the resistivity threshold of 30 Ωm . The gray rectangles indicate intervals with locally high resistivity and low integral chargeability.

similar resistivities are observed with both spacings, which reflects the fact that the fluid and the formation have resistivities in the same range. Based on fluid conductivity logging, the borehole fluid resistivity is around 10 Ωm in both wells (Table 1), i.e., only three times lower than the average 30 Ωm measured for the formation.

522

4.2 Changes in resistivity and integral chargeability over time

Logging results at +40 days (NN3 and NN4), +270 days (NN3), +380 days (NN4), 523 and +540 days (NN3 and NN4) are presented as relative differences (in percentage), com-524 pared to the baseline in Figure 6. The average value within a 2-meter thick interval is 525 taken for each round to smooth out rapid vertical variations. Error bars, calculated from 526 the standard deviation of this 2-meter average (equation 9), allow discriminating signif-527 icant changes over time from changes within the local variability. The resistivity after 528 baseline is further corrected to remove the temperature effect, following equation 16 and 529 using temperature logs shown in Figure B1 in order to assess resistivity changes reflect 530 changes in porosity or clay content. 531

$$\frac{\rho(T)}{\rho(T_0)} = 1 + \alpha_T (T_0 - T)$$
(16)

where T and T_0 are the current temperature and the reference temperature (corresponding to the baseline temperature), respectively. And $\alpha_T = 0.02^{\circ}C^{-1}$ is based on Arps (1953).

It can first be observed that changes depend on the electrode spacing for both the resistivity ρ and the integral chargeability M_{int} : relative differences are larger with the 64" spacing, indicating that more changes are happening in the rock formation than near the borehole wall (Figure 6). Resistivity variations are in many places within error bars, and thus, less focus is given here to interpret these variations.

Significant changes in the integral chargeability are observed. In NN4, M_{int} increases 540 at +40 days by 10-20% with the 64" spacing and 5-10% with the 16" spacing (Figure 541 6). At the next monitoring date, i.e., +380 days, M_{int} decreases back to the baseline in 542 most places with both spacings (note that data above 330 m were discarded for this date 543 due to low current injected). At the last monitoring date, +540 days, M_{int} remains at 544 baseline values down to 300 m, but the 64" signal decreases below baseline values be-545 tween 300 m and the bottom of the well (380 m). The 16" signal mostly remains at base-546 line values all along the well. A similar trend is observed in the bottom part of NN3, be-547 low 450 m: first M_{int} increases at +40 days by up to 50% with the 64" and up to 20% 548 with the 16" spacing. Then, M_{int} decreases for both spacings but remains above base-549 line values everywhere, unlike in NN4. At +540 days, below 450 m, slightly higher M_{int} 550 values are observed compared to +270 days for the 64" spacing. The 16" signal follows 551 the same trends everywhere, with lower amplitude. The maximum M_{int} increase is smaller 552 in NN4 than in NN3, but the increase is more uniformly spread along the borehole in 553 NN4. In particular, a significant increase is observed at +40 days in the whole well, i.e., 554 between 200 and 380 m. In NN3, the most important changes are located between 450 555 and 550 m, with only minimal changes down to 450 m. 556

A consistent observation links the 16- and 64" spacing in NN3 and NN4: M_{int} does not follow a monotonous increase as would have been expected if pyrite had been precipitating continuously or even if pyrite had precipitated up to a certain amount and remained in place. To illustrate further the non-monotonous pattern observed in the four panels showing M_{int} variations in Figure 6, the evolution of M_{int} with time in certain depth ranges is presented in Figure 7 for NN3 and NN4, using only the 64" spacing (larger variations) and showing absolute differences in mV/V.

Resistivity changes in NN3 with the 64" spacing show a spatial correlation with 564 M_{int} changes: a decrease of resistivity is observed in the upper part (above 450 m), and 565 an increase is observed in the bottom part. As opposed to the time-evolution observed 566 for M_{int} , the resistivity increase at the bottom of NN3 seems monotonous (Figure 6). 567 In NN4, a resistivity decrease is observed at +380 days above 300 m, and an increase 568 is observed between 300 and 350 m. However, an almost uniform resistivity decrease is 569 observed at +540 days, both with the 64" and 16" spacing, which indicates that the fluid 570 conductivity may have changed due to a change in fluid composition. Since the fluid con-571 ductivity was only measured at the last round (+540 days), further interpretation of the 572 resistivity changes regarding porosity, cementation, or clay content would be uncertain. 573

Overall, this qualitative analysis of the monitoring results indicates that (i) the integral chargeability strongly increases at +40 days in both wells and tends to decrease at the following monitoring rounds, going back or close to the baseline, (ii) the integral chargeability increase at +40 days is rather uniform over the whole depth interval in NN4 (200-400 m), while the increase in NN3 is localized between 450 and 550 m and (iii) the resistivity tends to increase monotonously over time at the bottom (below 450 m) of NN3,



Figure 6. Relative difference of the temperature-corrected resistivity ρ (left) and integral chargeability M_{int} (right) in NN3 (top) and NN4 (bottom), for both 64" and 16" electrode spacing. Colors indicate the monitoring rounds.



Figure 7. Evolution of the integral chargeability (absolute difference in mV/V) over time in NN3 (top) and NN4 (bottom) in different depth intervals, represented by the color scale. The dashed black line indicates the baseline level. Note the different scales for NN3 and NN4. Only data measured with 64" spacing are shown.

which may correspond to a decrease of porosity related to intense pyrite and secondary mineral precipitation, although impossible to quantify further.

In the following section, spectral information is extracted from TDIP monitoring results to allow a first-order quantification of pyrite precipitation in areas with an increase in M_{int} .

585

4.3 Extraction of spectral information through Debye decomposition

Since Debye decomposition tends to amplify the noise in the polarizability curves, decays that could not be fitted by a stretched exponential function (equation 13) with $R^2 \ge 0.998$ were discarded. This also smoothed out the decays that were kept and avoided fitting noise with the Debye decomposition. The theoretical justification and illustration of using a stretched exponential model for TDIP data is further described in Appendix Appendix A (Figures A1 and A2). Figure 4 also illustrates that the stretched exponential fit has minimal difference with the polarizability data in most cases.

⁵⁹³ Debye decompositions, illustrated in Figure 8 for two sets of decays in NN3 and ⁵⁹⁴ NN4 (64" spacing), show a similar trend as observed in Figures 6 and 7. The polariz-⁵⁹⁵ ability decays at +40 days are above the baseline as well as the later monitoring rounds, ⁵⁹⁶ both in NN3 and NN4. This trend is also reflected in the coefficients m_k , especially at ⁵⁹⁷ long τ_k values, and the resulting m_{tot} , larger at +40 days. All Debye decompositions are ⁵⁹⁸ available as figure sets S1 to S4 in Supplementary Information.

As can be seen on Figure A1, only 70% of the decays passed all the processing steps. Most affected datasets were at +380 days in NN4 (above 330 m) and at +40 days in NN3 (above 450 m). The noisy data above 450 m at +40 days is attributed to two factors that can be observed in Figure 3: (i) significantly stronger 50 Hz noise above 450 m than be-



Figure 8. Debye decomposition for two sets of decays measured with the 64" spacing in a) NN3 at 492-494 m and b) NN4 at 362-364 m. Left: polarizability data and fit by the RTD function (equation 10); right: coefficients m_k for all the τ_k at the four measuring rounds.

low at all dates (reason unknown) and (ii) stronger 50 Hz noise at +40 days compared
to baseline, at all depths. As can be observed in Figure 3, the signal level is always better with the 16" (blue) than with the 64" (red) electrode spacing, which is expected considering that the measured voltage decreases for increasing spacing with this configuration. This explains that fewer 16" data were discarded during the Debye decomposition
filtering procedure (A1). However, the 64" data involve a larger depth of penetration around
the borehole and are thus more relevant for conversion to pyrite volume change.

On the one hand, the integral chargeability presents the advantage of smoothing 610 out noise and providing a qualitative overview of the trends for the whole dataset. On 611 the other hand, Debye decomposition allows extracting spectral information and further 612 quantifying pyrite volume fraction change. Ideally, a unique relation would exist between 613 m_{tot} values obtained by Debye decomposition and the integral chargeability M_{int} , which 614 would allow to convert M_{int} into pyrite volume fraction. However, we find that differ-615 ent relations exist between m_{tot} and M_{int} , depending on the mean relaxation time. In-616 vestigating further these relations is beyond the scope of this study. Thus, pyrite esti-617 mations in the next section are calculated on a reduced portion of the dataset, where De-618 by decomposition was carried out. 619

4.4 Quantification of pyrite volume fraction change

The absolute difference in total chargeability, Δm_{tot} , further converted into Δp_v 621 are presented in Figure 9 for the three monitoring rounds. While the same trends are 622 observed as in Figure 6, this conversion allows estimating the extent of pyrite precipi-623 tation or dissolution. Error bars also differentiate between significant differences in pyrite 624 volume fraction versus differences too uncertain to be interpreted further. Overall, pre-625 cipitation of up to 2% and 1% are observed with the 64" and 16" electrode spacing, re-626 spectively. Figure 9 illustrates that more volume fraction changes are observed when tak-627 ing into consideration a larger volume around the boreholes, which emphasizes once more 628 that changes are not limited to the very near vicinity of the injection boreholes. 629

In NN4, an increase in pyrite volume fraction at +40 days (+0.5-1% compared to 630 baseline) is observed with both the 16" and 64" electrode spacings at all depths, followed 631 by a decrease at +380 and +540 days. In addition, the 64" data indicate that the de-632 crease at +540 days extends below baseline values (-0.2-0.5% compared to baseline). In 633 NN3, changes are more depth-dependent than in NN4. Between 200 and 450 m, no sig-634 nificant changes are observed at any rounds. However, it is important to note that, at 635 +40 days, most of the 64" data in this depth-range were discarded during the Debye de-636 composition filtering procedure. Below 450 m, all the way down to 550 m, +1-2% in the 637 pyrite volume fraction is suggested by the 64" data and +0.5-1% by the 16" data. At 638 +380 and +540 days, the 64" and 16" data, where available, suggest a decrease towards 639 baseline values, yet never extending below baseline values, as opposed to NN4. 640

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4.5 Analysis of the mean relaxation time

Overall, relaxation times are shorter in NN4 than in NN3, with a maximum of 300 642 msec in NN4 and above 1000 msec in NN3 (Figure 10). It can also be noted that the vari-643 ations of the average relaxation times across the different monitoring rounds follow the 644 variations of m_{tot} . In NN3 below 450 m, τ_{mean} increases at +40 days and decreases back 645 at +270 days. In NN4 across the whole well, τ_{mean} increases at +40 days and decreases 646 back at +270 days. In NN3, significantly longer relaxation times are observed below 450647 m, which reflects the different shape of the polarizability decay curves, with most of the 648 decay happening at late times, as opposed to depths above 450 m. Similarly, in NN4, 649 decays at 346-348 m and at 380-382 m have a different shape and correspond to min-650 imum and maximum values of τ_{mean} over the whole well, 40 and 400 msec, respectively. 651



Figure 9. Estimations of pyrite fraction change (absolute difference of pyrite vol-%) in NN3 (top) and NN4 (bottom), based on TDIP logging with 64" and 16" electrode spacing, using Debye decomposition of the decays.



Figure 10. Left: mean relaxation times, τ_{mean} , from Debye decomposition in NN3 (top) and NN4 (bottom), for both the 64" and 16" spacings, at different monitoring rounds. Polarizability curves for four selected depths for each well, using the 64" spacing. The corresponding depths are shown on the depth- τ_{mean} plots with gray rectangles.

As was observed in Figure 1, longer relaxation times, corresponding to lower fre-652 quencies, can be associated with either larger metallic grain sizes or connected metal-653 lic grains, e.g., forming veins. In this case, relatively small pyrite grain sizes are expected 654 due to the short time available, especially after +40 days (around 10 μm according to 655 laboratory experiments presented by (Prikryl et al., 2018)). Therefore, long relaxation 656 times in NN3, below 450 m, may indicate connected pyrite particles in fractures. On the 657 other hand, shorter relaxation times in NN4 may be associated with small, disseminated 658 pyrite particles. 659

It is important to note here that TDIP data are only capturing polarization processes associated with relaxation times larger than 3 msec, which is above the typical range of relaxation times investigated by FDIP laboratory studies, such as Martin and Weller (2023) and (Revil, Abdel Aal, et al., 2015). Therefore, converting the relaxation time distributions into grain size distribution would be uncertain.

5 5 Discussion

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5.1 Interpreting subsurface processes behind the monitored resistivity and TDIP responses

First, the justification for converting total chargeability into pyrite volume frac-668 tion changes is discussed here. This conversion assumes that pyrite precipitation is the 669 process governing the increase in chargeability. Based on reaction path modeling car-670 ried out in similar conditions, it is likely that some amount of smectite precipitates (Galeczka 671 et al., 2022). Still, smectite precipitation is expected to have little influence on the IP 672 signal in the presence of pyrite and could not alone explain the observed chargeability 673 increases (Lévy, Gibert, et al., 2019; Lévy, Weller, & Gibert, 2019; Revil et al., 2017). 674 Precipitation of iron-(oxyhydro)oxides following basalt dissolution is also possible, but 675 only magnetic iron-oxides, such as magnetite or ilmenite, yield a strong polarization re-676 sponse (Peshtani et al., 2022). In these conditions, only non-magnetic Fe(III)-bearing 677 phases are expected, such as goethite or hematite, based on laboratory experiments (Menefee 678 et al., 2018; Andreani et al., 2009), field-scale modeling (Galeczka et al., 2022), and the 679 general low-temperature alteration sequence of Icelandic basalt (Crovisier et al., 1992; 680 Arnórsson et al., 1983; Gunnlaugsson & Arnòrsson, 1982; Stefánsson & Gíslason, 2001). 681 In addition, amorphous silica and other silica-based minerals, such as quartz and zeo-682 lites, are expected (Daval et al., 2011; Galeczka et al., 2022). Amorphous silica, zeolites, 683 and Fe(III)-bearing phases further have a strong tendency to clog the pore space (Menefee 684 et al., 2018; Andreani et al., 2009), without creating a distinctive polarization signal (Revil 685 et al., 2002). Finally, the influence of microbial activity on the IP response is still poorly 686 understood but seems to yield polarization of much smaller amplitude than the observed 687 changes (Mellage et al., 2018) and will therefore not be considered here. 688

With these considerations in mind, TDIP monitoring brings three types of insights: spatial variability, long-term evolution, and quantity of pyrite volume changes.

The spatial variability of pyrite precipitation is assessed as a function of depth, vari-691 ability between NN3 and NN4, and distance from injection well. First, it is clear that 692 more pyrite precipitation occurs in the depth range 450-550 m in NN3 than in the rest 693 of the well, while in NN4, rather uniform pyrite precipitation is inferred (Figures 6 and 694 9). This can be understood in light of the baseline resistivity (Figure 5), showing resis-695 tivity lower than 30 Ωm at levels deeper than 450 m in NN3. In this 450-550 m inter-696 val, more clay content was found at the time of drilling (Helgadóttir, 2021), indicating 697 preferential hydrothermal flow. While the injection rate is ten times larger in NN4 than 698 in NN3 (Table 1), more precipitation seems to occur at the bottom of NN3 compared 699 to NN4. This may be due to fracture flow transporting the H_2S -rich fluid rapidly away 700 in NN4, not allowing time for pyrite precipitation. This would be consistent with the over-701

⁷⁰² all lower resistivity in NN4 in the common depth interval 200-380 m, compared to NN3 ⁷⁰³ (Figure 5), and would mean that there exists an optimal permeability that maximizes ⁷⁰⁴ the volumetric change in pyrite, ensuring at the same time ample fluid supply (H_2S sup-⁷⁰⁵ ply) and sufficient fluid residence time (basalt dissolution).

Second, it can be inferred that more precipitation occurs away from the well than on the borehole walls, based on the comparison between the 64" and 16" datasets in both wells (Figure 6).

The time-evolution of resistivity, integral chargeability, total chargeability and mean 709 relaxation time show that subsurface processes may be more complex than the expected 710 continuous pyrite precipitation suggested by previous studies (Clark et al., 2020; Gun-711 narsson et al., 2018). Integral chargeability time-series presented in Figure 7 show that 712 in both wells, at most depths, the initial increase at +40 days is followed by a decrease, 713 extending below baseline values in NN4 at depths below 330 m. This trend, which is com-714 mon to the 64" and 16" spacings (Figure 6), is also observed with the total chargeabil-715 ity and relaxation time (Figures 10, 9 and A1). Here, three different interpretations are 716 explored: (i) pyrite is still precipitating after +40 days, but the IP signal decreases any-717 way, (ii) previously formed pyrite grains are "passivated" by a coating layer of secondary 718 minerals preventing polarization while no more pyrite is forming and (iii) pyrite is be-719 ing re-dissolved. 720

Interpretation (i) would be consistent with the observation by Martin and Weller (2023) that the same amount of pyrite, with increasing grain size, yields a lower IP signal. However, the relaxation time, which is the main indicator for particle size (Figure 1) also starts decreasing after +40 days (Figure 10), which indicates that pyrite particles, if they are still there, are not growing in size. This interpretation is, therefore, ruled out.

Interpretation (ii) is based on the fact that strong polarization effects are caused 727 by metallic particles only when they have a clean surface. If secondary minerals isolate 728 the pyrite surface from the fluid, the particles are passivated, and the polarization de-729 creases (Gurin et al., 2019). Amorphous silica and Fe(III)-bearing minerals are common 730 coating layers, which may coat pyrite grain surfaces, as well as reduce basalt dissolution 731 rates by coating fresh basalt and clogging the pore space (Menefee et al., 2018; Andreani 732 et al., 2009; Daval et al., 2011). Corrosion of pyrite surface in the presence of little amounts 733 of oxygen may also lead to a similar passivation effect, without significantly dissolving 734 the pyrite particle (Placencia-Gómez et al., 2013). 735

Interpretation (iii) contradicts the common idea that pyrite mineralization is sta-736 ble in this context once formed. However, practical field conditions differ from typical 737 hypotheses in reactive transport models, which are constrained by the availability of suit-738 able frameworks and databases. Firstly, a high disequilibrium near the borehole, where 739 H_2S -rich fluids are constantly injected, makes conventional "steady-state" assumptions 740 irrelevant. Secondly, previous experiments on CO_2 injection in basalt at Hellisheiði (sim-741 ilar set-up as the Nesjavellir field case presented here) triggered a bloom of iron-oxidizing 742 bacteria, such as *Gallionellaceae*, and more generally sulfur- and iron- oxidation mark-743 ers (Trias et al., 2017). At Nesjavellir, provided that oxygen was present in the system 744 at some point (which could be the case due to e.g., leakage in the pipe system happen-745 ing at the beginning of the H_2S injection), specific microbial communities may have de-746 veloped and triggered pyrite oxidation (Mielke et al., 2003; Percak-Dennett et al., 2017). 747 The fracture networks that are ubiquitous in basaltic rocks could also be a pathway for 748 intermittent oxygen delivery, which would extend the depth of the habitable zone for iron-749 oxidizing bacteria (Bochet et al., 2020). Overall, biologically-driven mechanisms, which 750 could explain an unexpected pyrite dissolution over time, are not taken into account in 751 previous studies claiming that pyrite mineralization is stable due to the difficulty of rep-752 resenting them correctly in reactive transport simulations. While high-temperature geother-753
mal systems may prevent the development of most microbial communities, the shallow
system studied here has a relatively low reservoir temperature, although the injected fluid
temperature is higher (Table 1).

The pyrite volume fraction change estimated in Figure 9 assumes, to simplify, that 757 all the decrease of chargeability is explained by pyrite dissolution. In practice, there may 758 be a mix of passivation and re-dissolution and, thus, a more conservative interpretation 759 of the vol.% presented in Figure 9 would be the change in pyrite volume fraction con-760 tributing to the IP signal. While it is impossible from geophysical data alone to deter-761 mine whether passivation (interpretation ii) or re-dissolution (interpretation iii) are re-762 sponsible for the observed signals, both could be further investigated by laboratory stud-763 ies, as well as by installing a more continuous geophysical monitoring infrastructure. 764

In addition to the uncertainty on the source of polarization decrease, the quantifi-765 cation of pyrite volume presents additional uncertainty. It should first be noted that the 766 influence of background polarization not related to sulfide minerals in the baseline (Fig-767 ure 5) is managed by only converting the difference of total chargeability between dif-768 ferent monitoring rounds into the difference in pyrite volume fraction. However, equa-769 tion 5 proposed by Revil, Florsch, and Mao (2015) and used here presents other limits, 770 as Martin and Weller (2023) found that the exact relation between m_{tot} and p_v depend 771 on the grain size of pyrite particles. Furthermore, the use of an equation calibrated with 772 FDIP measurements to convert TDIP measurements is made possible by "long" current 773 injections (2 seconds), where the TDIP and FDIP responses are supposed to overlap (Martin 774 et al., 2021). However, 2-second current injections remain short, compared to 64"second 775 used by Martin et al. (2021) to show the equivalence with laboratory measurements. There-776 fore, the pyrite difference calculated here from 2-second TDIP data is certainly under-777 estimated. It was also assumed here that, at a given depth, precipitation occurs uniformly 778 over a certain (undetermined) lateral extent, considered representative of near-borehole 779 conditions. Further modeling of the contribution from the different cylindrical layers around 780 the borehole to the measured signal, and thus of the pyrite volume in each of these lay-781 ers, could be achieved with inversion. 782

Finally, it is interesting to note that the mean relaxation time is not only time-dependent but also depth-dependent and that an overall consistency of the decay shapes at a given depth is seen over time (Figure 10). This suggests that the local lithology and fracture network influence the relaxation time, most likely by influencing the patterns for pyrite precipitation. Although this is purely speculative, it may have implications for a better understanding of fracture networks in subsurface storage systems.

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5.2 Requirements and added-value of IP-logging compared to current practices

As mentioned in the Introduction, current monitoring of field-scale H_2S reinjec-791 tion, and similarly of CO_2 reinjection, have major pitfalls: (i) tracer tests capture less 792 than 5% of the gas migration flow (Matter et al., 2016), (ii) mass balance calculations 793 assume that the amount of sulfur not captured by downstream measurements, compared 794 to expectations based on tracer tests, has mineralized into sulfide minerals (Gunnarsson 795 et al., 2018) and (iii) long-term predictions by reactive transport models are not calibrated 796 by field monitoring and thus present significant uncertainty (Ratouis et al., 2022; Aradót-797 tir et al., 2012; White et al., 2020). Little has been researched on the long-term stabil-798 ity of the newly formed minerals in a context where the continuous injection of H_2S (and similarly, CO_2) creates a constant disequilibrium in the system, in terms of acidity and 800 redox conditions in particular. Overall, current monitoring approaches are still indirect 801 and cannot physically confirm the presence, location, and quantity of expected miner-802 als in the rock matrix, nor their long-term stability. The results presented here show the 803 feasibility of monitoring pyrite precipitation with logging TDIP chargeability measure-804

ments within injection wells. However, several conditions/constraints/obstacles need to be overcome.

To start with, an appropriate tool is required. Compared to more conventional ap-807 plications of TDIP, typically in mining exploration, the expected polarization signal is 808 weaker (smaller volumes of sulfide minerals). At the same time, the presence of power 809 lines due to the proximity to the power plant creates a significant 50 Hz background noise. 810 Therefore, large current injection is essential to ensure a sufficient signal-to-noise ratio, 811 and the acquisition of voltage with a high sampling rate is essential to separate the con-812 813 tribution from 50 Hz noise and from the rock to the voltage waveform. In addition, the need for quantification of pyrite precipitation, and thus comparison to frequency-domain 814 IP calibrations, requires that current injection and voltage acquisition happen over a suf-815 ficiently long time to capture the polarization information needed for the interpretation. 816 All these specifications required the development of new hardware in the existing QL40-817 IP instrument. In particular, the need for both longer injection/acquisition time and a 818 higher sampling rate means that more data needs to be collected and recorded in the in-819 strument's memory. The instrument went from recording 100 samples per cycle (depth) 820 to 980. Nevertheless, a trade-off was adopted, and the current waveform, deemed less 821 critical for the analysis, was not recorded; only the average current at each depth was 822 recorded. In addition, the maximum injection/acquisition times were 2 seconds. Ideally, 823 recording the current full waveform, injecting during longer times, and combining dif-824 ferent injection times would allow a more accurate comparison to FDIP calibrations. 825

Field conditions are also important to consider to carry out successful TDIP log-826 ging. The instrument has a temperature limit of $70^{\circ}C$, and its exposure to corrosive flu-827 ids shall be limited. This can be challenging in the context of CO_2 and H_2S injection 828 wells at geothermal sites. On the other hand, injection wells are the location with strong 829 changes in mineralogy. At the Nesjavellir site, the need to stop the injection and cool 830 down the wells a few days before the measurements represented the main obstacle to re-831 peating the measurements more regularly. In addition, the method requires an opera-832 tor on-site to operate the instrument. In this context, where data redundancy is limited 833 while TDIP measurements are easily subject to noise, data loss is a critical aspect to man-834 age, especially for quantitative conversion. 835

Despite field limitations and remaining uncertainties, IP-logging monitoring brings 836 novel insights into the processes at play, as emphasized in the previous subsection. A pos-837 sible way forward would be to carry out the logging in dedicated monitoring wells rel-838 atively close to the injection wells to avoid the financial and environmental cost of stop-839 ping the injection during monitoring. Another possibility that would also fulfill the need 840 for more continuous data acquisition to overcome episodic data losses would be the in-841 stallation of permanent electrodes in dedicated wells. The electrodes would connect to 842 an instrument remaining at the surface during the whole monitoring period, and only 843 batteries would need to be changed periodically. Furthermore, this approach could al-844 low 2D or 3D monitoring if several wells are used together in a cross-borehole electrical 845 tomography approach. External data acquisition would also be helpful to further elu-846 cidate the physical processes behind the observed IP response. Monitoring the fluid con-847 ductivity over time appears essential to assess changes in fluid composition and further 848 interpret resistivity changes. Complementary logging methods may include sonic and den-849 sity logging for assessment of porosity changes, acoustic televiewer, and spinner logs for 850 mapping fractures and flow paths, and color optical camera for direct observations of the 851 precipitation of pyrite and biofilms at the borehole wall. 852

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5.3 Can the IP-logging method be applied for monitoring other reactive processes?

Due to the strong signal caused by sulfides and other metallic minerals (Pelton et 855 al., 1978), mapping ore deposits is still the main industrial application of IP methods 856 today. Here, we present the first field example where reactive processes involving the pre-857 cipitation and/or dissolution of metallic minerals are monitored with IP-logging. Other 858 field examples for monitoring, with IP methods, reactive processes involving metallic min-859 erals include the immobilization of uranium contamination by stimulating iron and sulfate-860 reducing microorganisms (Flores Orozco et al., 2011) as well as the injection of zero-valent-861 iron amendment to aid bioremediation of chlorinated solvents (Flores Orozco et al., 2015). 862 In these examples, surface TDIP is used, and the limited spatial and temporal resolu-863 tion complicates the interpretation. There, IP-logging could help improve the understand-864 ing of physical processes at play. 865

Studies on the link between IP and bacterial activity often find that microbially-866 mediated metallic minerals cause an IP response (Williams et al., 2005; Flores Orozco 867 et al., 2011; Slater et al., 2007). However, recent studies suggest that polarization of bac-868 terial cells themselves may control the IP response in certain cases (Mellage et al., 2018; 869 Strobel et al., 2023). Therefore, in addition to monitoring the bioremediation of contam-870 inants, IP-logging could also be applied to the study of microbial life in a dynamic nat-871 ural environment, such as intermittent oxic-anoxic fluid mixing in fractured rocks (Bochet 872 et al., 2020), provided that upscaling issues can be overcome (Mellage et al., 2018). 873

Finally, we evaluate to what extent the method developed here could be applied 874 to monitoring CO_2 injection in basalts. Laboratory studies suggest that calcite precip-875 itation causes significant polarization increase (Wu et al., 2010). However, others observe 876 limited polarization increase, or even a decrease, in response to calcite precipitation (Wu et al., 2011; Zhang et al., 2012; Saneiyan et al., 2018) as well as no decrease associated 878 with calcite dissolution (Halisch et al., 2018). Recent studies attribute these contradic-879 tory findings to the dominating influence of pore water chemical changes or dissolution 880 changes on the IP response (Rembert, 2021; Izumoto et al., 2020). Therefore, the cur-881 rent petrophysical knowledge on the electrical response of calcite is insufficient to allow 882 direct mapping of calcite in the field with IP methods. Nevertheless, H_2S is often as-883 sociated with CO_2 in emissions from geothermal or coal power plants (Wang et al., 2011; 884 Fridriksson et al., 2016). H_2S is often considered an impurity and is only injected with 885 CO_2 to minimize the cost of separation. Still, the formation of pyrite could be used as 886 a marker of mineralization processes, given that pyrite and calcite precipitations are pri-887 marily controlled by basalt dissolution rates (Prikryl et al., 2018; Stefánsson et al., 2011) 888 and also share similar kinetics (Plummer et al., 1979; Williamson & Rimstidt, 1994). 889

⁸⁹⁰ 6 Conclusions

To date, monitoring efforts of an implemented field-scale H_2S reinjection system have only occurred via concentration measurements, mass balance calculations, and analysis of precipitates on submersible pumps in injection and downgradient boreholes. These studies, along with geochemical numerical models and laboratory simulations, concluded effective H_2S sequestration through pyrite formation. Borehole sampling of fluids, although effective at identifying the occurrence of sulfide mineralization, provides limited quantitative, spatial, and temporal information on in-situ sulfide mineralization.

In order to bring insights into the subsurface physical processes controlling mineralization in the context of H_2S injection in basalts, as well as quantitative information on the sulfide mineralization in a specific case, a "shallow" (200-550 m) injection of H_2S at Nesjavellir geothermal field was monitored by means of resistivity and time-domain induced polarization wireline logging. Pyrite precipitation was expected to affect primarily the induced polarization signal, while other changes (e.g., fluid conductivity, temperature, clay minerals precipitation) were expected to affect the resistivity.

The QL40-IP logging tool, developed by the company Advanced Logic Technology, was modified to allow longer current injection and a larger sampling rate than previously. This was needed to filter a strong 50 Hz background signal due to the proximity of a buried power line and convert the TDIP data into sulfide volumes using a laboratory-based relationship.

The first logging measurements were carried out in the summer of 2019, and the baseline with the modified tool was measured in September 2020. H_2S injection began on January 29th, 2021. After the start of injection, five logging monitoring rounds (+40, +270, +380, +540, +630 days) were carried out in wells NN3 and NN4. Out of these five rounds, only three remained for each well due to too low current injected in the other rounds for unknown reasons.

Monitoring results show that polarization strongly increases at +40 days in both 916 wells and tends to decrease at the following monitoring rounds, going back close to the 917 baseline or below. The polarization increase is rather uniform over the whole depth in-918 terval in NN4 (200-400 m), while the increase is clearly localized between 450 and 550 919 m in NN3, where more hydrothermal alteration was also found prior to the injection of 920 any wastewater. Conversion of chargeability absolute difference into pyrite volume frac-921 tion change indicates precipitation of up to 1% in NN4 and 2% in NN3 at +40 days. In 922 both wells, changes are more pronounced with the larger electrode spacing (64"), indi-923 cating that pyrite precipitation takes place away from the well. 924

The decrease of chargeability observed in both wells at +270 and +380 days sug-925 gests that pyrite is either passivated or re-dissolved after precipitating. The presence of 926 specific microbial communities that can produce oxygen at these depths should be fur-927 ther investigated. Recommendations for future monitoring include: (i) installation of per-928 manent monitoring systems to better understand the trends, (ii) considering the possi-929 bility of pyrite re-dissolution in reactive transport modeling of H_2S sequestration, and 930 (iii) validation of geophysical results by additional in-situ measurements and observa-931 tion, such as color camera and multi-level water sampling in wells. 932

933 7 Open Research

The TDIP data used for monitoring H_2S fate in the study, together with the resistivity, fluid conductivity and temperature data, will be available on the following Zenodo repository at the time of publication (Lévy, 2023): https://doi.org/10.5281/zenodo .10145643 with Creative Commons Attribution 4.0 International license. For review purposes, data are shared as supplementary information.

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Appendix A Added-value and justification of the stretched exponential fitting procedure

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The stretched exponential function $f(t) = \alpha e^{-\left(\frac{t}{\tau_{KWW}}\right)^{\beta}}$, also named Kohlrausch-1373 Williams-Watts (KWW), and the Havriliak-Negami (HN) relaxation function $g(\omega) = \frac{1}{(1+(i\omega\tau_{HN})^{\alpha}_{HN})^{\gamma}_{HN}}$ have been widely used to describe the relaxation behavior of glass-forming liquids and complex systems, in time- and frequency-domain respectively (Alvarez et al., 1374 1375 1376 1991). The HN relaxation function is a generalization of the Pelton model with one more 1377 parameter (Havriliak & Negami, 1966). It also corresponds to the "Generalized Cole-1378 Cole" model (Pelton et al., 1983). A relationship among the parameters of the KWW 1379 and HN models is suggested by the fact that both models yield an accurate description 1380 of real data. An empirical and mathematical relation between the coefficients of the KWW 1381 and HN functions is found by Alvarez et al. (1991). This relationship is not analytical 1382 since the HN and the KWW relaxation functions are not exactly Fourier transforms of 1383

each other. In particular, the relation between the relaxation times in both functions depends on the exponent beta in the KWW function, β . For example, for β close to 1, $\tau_{HN} \simeq \tau_{KWW}$, and for $\beta = 0.1$, $\tau_{HN} = 300\tau_{KWW}$. The KWW has one less parameter, so it can always be transformed into an HN model, but the contrary is not always true. The product $\alpha_{HN}\gamma_{HN}$ is close to β . In addition, even though the HN model is a generalized version of the Pelton model, the fitted chargeability parameter cannot be used to quantify pyrite as it is not strictly comparable to the chargeability in the Pelton model.

That's why the Debye decomposition is used here to relate the measured TDIP data with petrophysical relationships calibrated on FDIP data.

Nevertheless, based on the theory described by Alvarez et al. (1991), the stretched 1393 exponential function is used to automatically discriminate noisy discharge curves that 1394 do not follow the stretched exponential shape. Debye decomposition is only applied to 1395 discharge curves that can be reasonably fitted by a stretched exponential. Moreover, the 1396 stretched exponential fit allows smoothing out the "accepted" discharge curves that may 1397 still contain some noise, which in turn results in more meaningful outcomes from the De-1398 by decomposition. This is illustrated for the total chargeability, M_{tot} , and the mean re-1399 laxation time, τ_{mean} in Figures A1 and A2. In these figures, Debye decomposition car-1400 ried out on the stretched exponential fitted function is compared to Debye decomposi-1401 tion carried out on the polarizability data themselves (before the fitting step). The com-1402 parison clearly indicates less noise in the curves of SE fitted spectra. 1403

¹⁴⁰⁴ Appendix B Temperature and borehole diameter in NN3 and NN4



♦ +380 days ♦ +540 days Baseline +40 days ÷ ÷

Figure A1. Total chargeability from Debye decomposition in NN3 (top) and NN4 (bottom), with 64" (left) and 16" (right) spacing. The results are shown for Debye decompositions carried out on stretched exponential fit ("SE fit") and on original data. Note that the Debye decomposition on original data includes more discharge curves since it happens before the filtering procedure related to the SE fit.



Figure A2. Weighted-average relaxation time from Debye decomposition in NN3 (top) and NN4 (bottom), with 64" (left) and 16" (right) spacing. The results are shown for Debye decompositions carried out on stretched exponential fit ("SE fit") and on original data. Note that the Debye decomposition on original data includes more discharge curves since it happens before the filtering procedure related to the SE fit.



Figure B1. Temperature measured in NN3 and NN4 at the four monitoring rounds.



Figure B2. Borehole diameter (caliper) below the casing, measured in 2001 after drilling, in NN3 and NN4.

Supporting Information for "Understanding the fate of H_2S injected in basalts by means of time-domain induced polarization geophysical logging"

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Supporting Information (Files uploaded separately)

- 1. Captions for Datasets S1 to S5
- 2. Captions for Figure sets S1 to S4

Introduction

Two types of supporting information are provided to the article: (i) data sets S1 to S5 corresponding to the geophysical monitoring and (ii) sets of figures S1 to S4 showing the Debye decompositions for all the decays (all depths, all dates, both 16- and 64-inch electrode spacing).

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Data Set S1. Resistivity monitoring data for NN3, at all dates, with four different electrode spacings (including 16- and 64-inch, as well as 8- and 32-inch). Naming convention: ds01

Data Set S2. Resistivity monitoring data for NN4, at all dates, with four different electrode spacings (including 16- and 64-inch, as well as 8- and 32-inch). Naming convention: ds02

Data Set S3. Temperature and fluid conductivity measured in july 2022 in both NN3 and NN4 (+540 days monitoring round). Naming convention: ds03

Data Set S4. Full waveform time-domain induced polarization data in NN3, at all dates. Each spreadsheet corresponds to a given date (refer to Table 2 in the main manuscript for the correspondance between dates and time after injection start). The first two worksheets have the voltage waveforms for 16- and 64-inches. The third worksheet includes measurements of the injected electric current. Additional worksheet contains additional information on the acquisition settings. Naming convention: ds04

Data Set S5. Full waveform time-domain induced polarization data in NN4, at all dates. Each spreadsheet corresponds to a given date (refer to Table 2 in the main manuscript for the correspondance between dates and time after injection start). The first two worksheets have the voltage waveforms for 16- and 64-inches. The third worksheet includes measurements of the injected electric current. Additional worksheet contains additional information on the acquisition settings. Naming convention: ds05

Figure set S1.

Debye decomposition of 2m-averaged polarizability decays, after the step of stretched exponential fit, for data in NN3, with the 16-inch spacing, at all monitoring dates. Naming convention: fs01

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Figure set S2.

Debye decomposition of 2m-averaged polarizability decays, after the step of stretched exponential fit, for data in NN3, with the 64-inch spacing, at all monitoring dates. Naming convention: fs02

Figure set S3.

Debye decomposition of 2m-averaged polarizability decays, after the step of stretched exponential fit, for data in NN4, with the 16-inch spacing, at all monitoring dates. Naming convention: fs03

Figure set S4.

Debye decomposition of 2m-averaged polarizability decays, after the step of stretched exponential fit, for data in NN4, with the 64-inch spacing, at all monitoring dates. Naming convention: fs04