High-resolution induced polarization imaging of biogeochemical carbon-turnover hot spots in a peatland

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Abstract

Biogeochemical hot spots are defined as areas where biogeochemical processes occur with anomalously high reaction rates relative to their surroundings. Due to their importance in carbon and nutrient cycling, characterization of hot spots is critical to accurately predict carbon budgets in the context of climate change. However, biogeochemical hot spots are difficult to identify in the environment, as sampling resolutions are often too coarse to find these areas in the subsurface. Here, we present imaging results of a geophysical survey using the non-invasive induced polarization (IP) method to identify biogeochemical hot spots of carbon turnover in a minerotrophic wetland. To interpret the field-scale IP signatures, geochemical analyses were performed on freeze-core samples obtained in areas characterized by anomalously high and low IP responses. Our results reveal large variations in the electrical response, with the highest IP phase values (> 20 mrad) corresponding with high concentrations of phosphates (>4000 μM), an indicator of carbon turnover. Moreover, analysis of the freeze core reveal negligible concentrations of iron sulfides. The extensive geochemical and geophysical data presented in our study demonstrates that IP images can assess changes in the biogeochemical activity in peat, and identify hot spots.

Keywords: biogeochemical carbon turnover; geophysical imaging methods; electrical conductivity; induced polarization; microbiologically active zones
1 Introduction

In terrestrial and aquatic ecosystems ‘patches’ or areas that show disproportionally high biogeochemical reaction rates relative to the surrounding matrix are referred to as biogeochemical ‘hot spots’ (McClain et al., 2003). Hot spots for turnover of redox-sensitive species (e.g. oxygen, nitrate or dissolved organic carbon) are often generated at interfaces between oxic and anoxic environments, where the local presence/absence of oxygen either favours or suppresses biogeochemical reactions such as aerobic respiration, denitrification or oxidation/reduction of iron (McClain et al., 2003). Biogeochemical hot spots are important for nutrient and carbon cycling in various systems such as wetlands (Frei et al., 2010; 2012), lake sediments (Urban, 1994), the vadose zone (Hansen et al., 2014), hyporheic areas (Boano et al., 2014) or aquifers (Gu et al., 1998). Wetlands are distinct elements in the landscape, which are often located where various hydrological flow paths converge, such as at the bottoms of basin shaped catchments, local depressions or around rivers and streams (Cirmo and McDonell 1997). Wetlands are attracting increasing interest because of their important contribution to water supply, water quality, nutrient cycling, and biodiversity (Costanza et al., 1997; 2017). Understanding microbial moderated cycling of nutrients and carbon in wetlands is critical, as these systems store a significant part of the global carbon through the accumulation of decomposed plant material (Kayranli et al., 2010). In wetlands, water table fluctuations as well as plant roots determine the vertical and horizontal distribution of oxic and anoxic areas (Frei et al., 2012; Gutknecht et al., 2006). Small scale subsurface flow processes in wetlands, moderated by micro-topographical structures (hollow and hummocks) (Diamond et al., 2020), can control the spatial presence of redox-sensitive solutes and formation of biogeochemical hot spots (Frei et al., 2010; 2012). Despite their relevance for the carbon and nutrient cycles, the field observations of the basic mechanisms controlling the formation and distribution of biogeochemical hot spots in space is largely missing.

Biogeochemical active areas traditionally have been identified and localized through chemical analyses of point samples from the subsurface and then subsequent interpolation of the data in space (Morse et al., 2014; Capps and Flecker, 2013; Hartley and Schlesinger, 2000). However, such point-based sampling methods may either miss hot spots due to the low spatial resolution of sampling (McClain et al., 2003) or disturb the redox-sensitive conditions in the subsurface by bringing oxygen into anoxic areas during sampling. Non-invasive methods, such as geophysical techniques, have the potential to study biogeochemical hot spots in-situ without interfering with the subsurface environment (e.g., Williams et al., 2005; 2009; Atekwana and Slater 2009; Flores Orozco et al., 2015; 2019; 2020). Geophysical methods permit to map large
areas in 3D and still resolve subsurface physical properties with a high spatial resolution (Binley et al., 2015).

In particular, the induced polarization (IP) technique has recently emerged as a useful tool to delineate biogeochemical processes in the subsurface (e.g., Kemna et al., 2012; Kessouri et al., 2019; Flores Orozco et al., 2020). The IP method provides information about the electrical conductivity and the capacitive properties of the ground, which can be expressed, respectively, in terms of the real and imaginary components of the complex resistivity (Binley and Kemna, 2005). The method is commonly used to explore metallic ores because of the strong polarization response associated to metallic minerals (e.g., Marshall and Madden, 1959; Seigel et al., 2007). Pelton et al. (1978) and Wong (1979) proposed the first models linking the IP response to the size and content of metallic minerals. More recently, the role of chemical and textural properties in the polarization of metallic minerals has been investigated in detail (Revil et al., 2012, 2015a; 2015b, Bücker et al., 2018; 2019). In porous media without a significant metallic content, the IP response can be related to the polarization of the electrical double layer formed at the grain-fluid interface (e.g., Waxman and Smits, 1968; Revil and Florsch, 2010; Revil, 2012). In particular, Revil et al. (2017) carried out complex conductivity measurements on a large set of soil samples, for which they report a linear relation between the magnitude of the polarization response and the cation exchange capacity (CEC), which in turn is related to surface area and surface charge density.

Since the early 2010s, various studies have explored the potential of IP measurement for the investigation of biogeochemical processes (Slater and Attekwana, 2013). Laboratory studies on sediment samples examined the correlation between the spectral induced polarization (SIP) response and X iron sulfid precipitation caused by iron reducing bacteria (Williams et al., 2005; Ntarlagiannis et al., 2005; 2010; Slater et al., 2007; Personna et al., 2008; Zhang et al., 2010; Placencia Gomez et al., 2013; Abdel Aal et al., 2014). Further investigations in the laboratory have also revealed an increase in the polarization response accompanying the accumulation of microbial cells and biofilms (Davis et al., 2006; Abdel Aal et al., 2010a, 2010b; Albrecht et al., 2011; Revil et al., 2012; Zhang et al., 2013; Mellage et al., 2018; Rosier et al., 2019; Kessouri et al., 2019).

Motivated by these successful laboratory measurements (i.e., Abdel Aal and Attekwana, 2014), the IP method has also been used to characterize biogeochemical degradation of contaminants at the field scale (Williams et al., 2009; Slater and Binley 2006; Flores Orozco et al., 2011; 2012b; 2013; 2015; Maurya et al., 2017). Natural accumulations of organic matter in areas with
indigenous iron-reducing bacteria might result in naturally reduced zones and the precipitation of iron sulfide minerals. Wainwright et al. (2016) demonstrated the applicability of IP imaging to identify naturally reduced zones, i.e., hot spots in floodplains. Moreover, in a recent study, Flores Orozco et al. (2020) demonstrated the possibility to delineate biogeochemically active zones in a municipal solid waste landfill even in the absence of iron sulfides. Flores Orozco et al. (2020) argued that the high content of organic matter itself might explain both, the high polarization response and high rates of microbial activity; thus, opening the possibility to delineate biogeochemical hot spots that are not related to iron-reducing bacteria. This conclusion is consistent with previous studies performed in marsh and peat soils, areas with a high organic matter content and high microbial turnover rates (Mansoor and Slater, 2007; McAnallen et al., 2018). Peat soils are characterized by a high surface charge and have been suggested to enhance the IP response (Slater and Reeve, 2002). Mansoor and Slater (2007) concluded that the IP method is a useful tool to map iron cycling and microbial activity in marsh soils. Uhlemann et al. (2016) found differences in the electrical resistivity of peat according to saturation, microbial activity, and pore water conductivity; however, their study was limited to direct-current resistivity and did not investigate variations in the IP response. In contrast to these observations, laboratory studies have shown a low polarization response in samples with a high organic matter content, despite its high CEC (Schwartz and Furman, 2014). Based on field measurements, McAnallen et al. (2018) found that active peat is less polarizable due to variations in groundwater chemistry imposed by sphagnum mosses; while degrading peat resulted in low resistivity values and a high polarization response. Based on measurements with the Fourier Transform Infrared (FTIR) Spectroscopy in water samples, the authors concluded that the carbon-oxygen (C=O) double-bound in degrading peat correlated with the polarization magnitude of the peat material. Based on laboratory investigations, Ponziani et al. (2011) also conclude that decomposition of peat occurs predominantly by aerobic respiration, i.e. using molecular oxygen as the terminal electron acceptor to oxidize organic matter. Thus decomposition rates are expected to be highest at the interface between the oxic and anoxic zones.

Based on these promising previous results, we believe that the IP method is a potentially useful tool for in-situ investigation of biogeochemical processes. However, different lab and field investigations do not offer a clear interpretation scheme of general validity. This study presents an extensive IP imaging dataset collected at a peatland site to investigate the controls on the IP response in biogeochemically active areas. IP monitoring results are compared to geochemical data obtained from freeze cores and pore water sample analysis. Our main objectives are (i) to
assess the applicability of the IP method to spatially delimit highly active biogeochemical areas in the peat soil and (ii) to investigate whether the local IP response is related to the accumulation of iron sulfides or high organic matter turnover.

## 2 Material and methods

### 2.1 Study site

The study site is part of the Lehstenbach catchment located in the Fichtelgebirge mountains (Fig. 1a), a low mountain range in north-eastern Bavaria (Germany) close to the border to the Czech Republic. Various soil types including Dystric Cambisols, Haplic Podsols, and Histosols (i.e. peat soil) cover the catchment area of approximately 4.2 km², situated on-top of variscan granite bedrock (Strohmeier et al., 2013). The catchment is bowl shaped (Fig. 1b), and minerotrophic riparian peatlands have developed around the major streams. The plot scale study site (Fig. 1c) is located in a riparian peatland draining into a nearby stream close to the catchment’s outlet (Fig. 1b).

The groundwater level in this area annually varies within the top 30 cm of the peat soil and was 5-10 cm below the surface during the measurement campaign. The local groundwater flow has a S-SW orientation (Durejka et al., 2019) towards a nearby drainage ditch. Permanently high water saturation of the peat soil favors the development of anoxic biogeochemical processes close to the surface. Frei et al. (2012) demonstrated that hot spots, related to the stimulation of iron reducing bacteria and accumulation of iron sulfides, are generated by small scale subsurface flow processes and the spatial non uniform availability of electron acceptors and donors induced by the peatland’s typical micro topography. Transport limited and spatially non uniform availability of electron acceptors and donors, which to a large extent is controlled by subsurface flow patterns induced by the wetland’s micro-topography (hummocks and hollows), in combination with labile carbon stocks are the primary drivers in generating biogeochemical hot spots in the peatland (Frei et al., 2012; Mishra and Riley, 2015).

### 2.2 Experimental plot and geochemical measurements

The experimental plot for the geophysical measurements covers approximately 160 m² (12.6 x 12.6 m, Fig. 1c) of the riparian peatland. Sphagnum Sp. (peat moss) and Molinia caerulea (purple moor-grass) dominate the vegetation, with the sphagnum and purple moor-grass abundance being higher in the Northern part of the plot (Fig. 2a and 2b). In the Southeastern region, where the sphagnum is less abundant, permanent surface runoff was observed (Fig. 2d). To measure the thickness of the peat, a stainless-steel rod (0.5 cm in diameter) was pushed into...
the ground until it reached the granitic bedrock (similar to Parry et al., 2014). Peat thickness was measured with a 1 m resolution in E-W direction and 0.5 m resolution in N-S direction. During the experiments, the local groundwater level was measured in two piezometers. Groundwater samples were collected at three different locations (S1, S2, and S3 indicated in Fig. 3) using a bailer. Pore water profiles were taken at S1, S2, and S3 at 5 cm intervals to a maximum depth of 50 cm below ground surface (bgs) using stainless steel mini-piezometers. All water samples were filtered through 0.45 µm filters and analyzed for fluoride, chloride, nitrite, bromide, nitrate, phosphate, and sulfate using an ion chromatograph (Compact IC plus 882, Metrohm GMBH). Dissolved organic carbon (DOC) was measured using a Shimadsu TOC analyzer via thermal combustion. Dissolved iron species (Fe$^{2+}$) and total iron (Fe$^{tot}$) concentrations were measured photometrically using the 1,10-phenanthroline method on pore water samples that had been stabilized with 1% vol/vol 1M HCl in the field (Tamura et al., 1974). Two freeze cores (see Fig. 2d) were extracted at locations S1 and S2 (Fig. 3) by pushing an 80-cm long stainless-steel tube into the peat. After the tube was installed, it was filled with a mixture of dry ice and ethanol. After around 20 minutes, the pipe with the frozen peat sample was extracted and stored on dry ice for transportation to the laboratory at the University of Bayreuth. Both freeze cores were cut into 10 cm segments. Each segment was analyzed for reactive iron (1M HCl extraction and measured for Fe$^{tot}$ as described above) (Canfield, 1989), reduced sulfur species using the total reduced inorganic sulfur (TRIS) method (Canfield et al., 1986) and carbon and nitrogen concentrations after combustion using a thermal conductivity detector. Peat samples were also analyzed by FTIR using a Vector 22 FTIR spectrometer (Bruker, Germany) in absorption mode; with subsequent baseline subtraction on KBr pellets (200 mg dried KBr and 2 mg sample). Thirty-two measurements were recorded per sample and averaged from 4.500 to 600 cm$^{-1}$ in a similar manner to Biester et al. (2014).

2.3 Non-invasive techniques: induced polarization measurements

The induced polarization (IP) imaging method, also known as complex conductivity imaging or electrical impedance tomography, is an extension of the electrical resistivity tomography (ERT) method (e.g., Kemna et al., 2012). As such, it is based on four-electrode measurements, where one pair of electrodes is used to inject a current (current dipole) and a second pair of electrodes is used to measure the resulting electrical potential (potential dipole). Modern devices can measure tens of potential dipoles simultaneously for a given current dipole, permitting the collection of dense data sets within a reasonable measuring time. This provides an imaging framework to gain information about lateral and vertical changes in the electrical properties of the subsurface. IP data can be collected in the frequency domain (FD), where an
alternating current is injected into the ground where the polarization of the ground leads to a measurable phase shift between the periodic current and voltage signals. FD measurements can be conducted at different frequencies to gain information about the frequency dependence of the electrical properties of the subsurface. This approach is known as spectral IP (SIP). From the ratio of the magnitudes of the measured voltage and the injected current, as well as the phase shift between the two signals, we can obtain the electrical transfer impedance. The inversion of imaging data sets, i.e. a large set of such four-point transfer-impedance measurements collected at different locations and with different spacing between electrodes along a profile, permits to solve for the spatial distribution of the electrical properties in the subsurface (see deGroot-Hedlin and Constable, 1990; Kemna et al., 2000; Binley and Kemna, 2005).

IP imaging results can be expressed in terms of the complex conductivity \( \sigma^* \), or its inverse the complex resistivity \( \sigma^* = 1/\rho^* \). The complex conductivity can either be denoted in terms of its real \( \sigma' \) and imaginary \( \sigma'' \) components, or in terms of its magnitude \( |\sigma| \) and phase \( \psi \):

\[
\sigma^* = \sigma' + i\sigma'' = |\sigma|e^{i\psi}, \tag{1}
\]

where \( i = \sqrt{-1} \) is the imaginary unit, \( |\sigma| = \sqrt{\sigma'^2 + \sigma''^2} \) and \( \psi = \tan^{-1} \frac{\sigma''}{\sigma'} \).

The real part of the complex conductivity is mainly related to the Ohmic conduction, while the imaginary part is mainly related to the polarization of the subsurface. For a more detailed description of the IP method, the reader is referred to the work of Ward (1988), Binley, and Kemna (2005).

In this study, we conducted FDIP measurements at 1 Hz along 65 lines during a period of four days in July 2019. We used the DAS-1 instrument manufactured by Multi-Phase Technologies (now MTP-IRIS Inc.). We collected 64 N-S oriented lines (referred to as By 1 to By 64) with 20 cm spacing between each line. One additional line (By 68) was collected with a W-E orientation, which intersects the N-S oriented lines at 3 m, as presented in Fig. 3. Each profile consisted of 64 stainless steel electrodes (3 mm diameter) with a separation of 20 cm between each electrode (Fig. 2c). Besides the short electrode spacing, the use of a dipole-dipole configuration with a unit dipole length of 20 cm warranted a high resolution within the upper 50 cm of the peat, where the biogeochemical hot spots were expected. We deployed a dipole-dipole skip-0 (i.e., the dipole length for each measurement is equal to the unit spacing of 20 cm) configuration, voltage measurements were collected across eight adjacent potential dipoles for each current dipole. The dipole-dipole configuration avoids the use of electrodes for potential measurements previously used for current injections to avoid contamination of the data caused by remnant polarization of electrodes. To evaluate data quality, reciprocal readings...
were collected along one profile every day (see e.g., LaBrecque et al., 1996; Flores Orozco et al., 2012a, 2019). Reciprocal readings refer to data collected after interchanging current and potential dipoles. We used coaxial cables to connect the electrodes with the measuring device to minimize the distortion of the data due to electromagnetic coupling and cross talking between the cables (e.g., Zimmermann et al., 2008, 2019; Flores Orozco et al., 2013), with the shields of all coaxial cable running together into one ground electrode (for further details see Flores Orozco et al., 2020).

3 Results
3.1 Data quality and processing

In Fig. 4, we present a modified pseudosection showing both normal (negative pseudodepth) and reciprocal (positive pseudodepth) readings in terms of apparent resistivity (\(\rho_a\)) and apparent phase (\(\phi_a\)) for the data collected along line By 25. Plots in Fig. 4 show consistency between the normal and reciprocal readings of apparent resistivity (4a) and phase (4b). The principle of reciprocity asserts that normal and reciprocal readings should be the same (e.g., Slater et al., 2000). Hence, the misfit between them can be used to detect outliers, and to quantify data errors (LaBrecque et al., 1996; Flores Orozco et al., 2012a; Slater and Binley, 2006; Slater et al., 2000). Figures 4c and 4d show the histograms of the normal-reciprocal misfits along line By 25 for both the resistance and phase (\(\Delta R\) and \(\Delta \phi\) respectively), which exhibit near Gaussian distributions with low standard deviations (as expected for random noise) for both the normalized resistance (\(S_R=0.027\)) and the apparent phase (\(S_\phi=1.1\) mrad). Readings exceeding these standard deviation values were considered as outliers (between 16 and 33% of the data at the different lines) and were removed from the data set prior to the inversion.

For the inversion of the IP imaging data set, we used CRTomo, a smoothness-constrained least-squares algorithm by Kemna (2000) that allows inversion of the data to a level of confidence specified by an error model. We used the resistance and phase error models described by Kemna (2000) and Flores Orozco et al. (2012a). The resistance (\(R\)) error model is expressed as \(s(R)=a+bR\), where \(a\) is the absolute error, which dominates at small resistances (i.e., \(R<0\)), and \(b\) is the relative error, which dominates at high resistance values (LaBrecque et al., 1996; Slater et al., 2000). For the phase, the error model is also expressed as a function of the resistance \(s(\phi_a)=cR^d\), where \(d<0\) in our study due to the relative low range in the measured resistances (see Flores Orozco et al., 2012a for further details). If \(d\to0\), the model reduces to
the constant-phase-error model (Flores Orozco et al., 2012a) with \( s(\phi) = c \) described by Kemna (2000) and Slater and Binley (2006). We quantified the error parameter for each line collected as normal and reciprocal pairs (using the approach outlined by Flores Orozco et al., 2012a) and computed the average value of the error parameters for the inversion of the entire data set. Here, we present inversion results obtained using the error parameters, \( a = 0.001 \, \Omega \), \( b = 2.2 \% \), and \( c = 1 \, \text{mrad} \). For the imaging, we defined a cut-off value of the cumulated sensitivity of \( 10^{2.75} \), with pixels related to a lower cumulated sensitivity blanked in the images. The cumulated sensitivity values are a widely used parameter to assess the depth of investigation (Kemna et al., 2002; Flores Orozco et al., 2013).

### 3.2 Complex conductivity imaging results and their link the peat thickness and land cover

The thickness of the peat in the plot was found to vary between 40 and 160 cm (Fig. 5). The thickness of the peat unit increased in the W-E direction, with much smaller variations in the N-S direction. Variations in the vegetation cover (as indicated by the three vegetation classes, abundant \((av)\), moderate \((mv)\) and sparse \((sv)\)) do not seem to correspond with the variations in the peat thickness. Note that the N-S orientation of the majority of IP lines is approximately aligned with the direction of minimum changes in the peat thickness.

Figure 6 shows the imaging results of the N-S oriented profiles By 25, By 46, and the W-E oriented profile By 68 expressed in terms of the real- \((\sigma')\) and imaginary \((\sigma'')\) part of the complex conductivity. These images reveal three main electrical units: (i) a shallow unit with high \( \sigma' \) (>5 mSm\(^{-1}\)) and high \( \sigma'' \) (>80 μSm\(^{-1}\)) values in the top 10-20 cm bgs, (ii) an intermediate unit with moderate to low \( \sigma' \) (<5 mSm\(^{-1}\)) and moderate \( \sigma'' \) (40–80 μSm\(^{-1}\)) values, and (iii) a deep unit characterized by moderate to low \( \sigma' \) (<5 mSm\(^{-1}\)) values and the lowest \( \sigma'' \) (<40 μSm\(^{-1}\)) values. The low complex conductivity (both \( \sigma' \) and \( \sigma'' \)) in the third unit corresponds to the granite bedrock. The compact structure of the granite, corresponding to low porosity, explains the observed low conductivity values (\( \sigma' <5 \, \text{mSm}^{-1} \)) due to low surface charge and surface area.

The second and third unit boundary corresponds to the contact between peat and granite measured using the metal rod (represented by the dashed line in Fig. 5). This shows that the electrical imaging can accurately resolve the contact with the bedrock - even in areas where it is located deeper than 1 m bgs.

The shallow and intermediate electrical units are related to the relatively heterogeneous peat (Fig. 6). In the first unit, the electrical images reveal lateral changes along the profiles oriented N-S (By 25 and By 46), while we generally observe higher polarization values (\( \sigma'' \)) in the
northern part of the profiles than in the southern part. This lateral change corresponds to the W-E oriented profile By 68, where the polarization values are the highest around sampling point S1.

The phase of the complex conductivity roughly represents the ratio of the polarization \( (\sigma'') \) relative to the Ohmic conduction \( (\sigma') \). Therefore, it has been argued that the phase represents the polarization effect better than the imaginary conductivity itself as it removes effects stemming from changes in fluid conductivity, saturation, and porosity (Kemna et al., 2004; Ulrich and Slater 2004). Similar to the \( \sigma'' \) images, the phase images presented in Fig. 7 resolve the three main units: (i) the shallow peat unit in the top 10-50 cm is characterized by the highest values \( (\phi > 22 \text{ mrad}) \), (ii) the intermediate unit still corresponding to peat, is characterized by moderate \( \phi \) values (between 13 and 22 mrad), and (3) the deep unit, associated to the granitic bedrock, related to the lowest \( \phi \) values \((< 13 \text{ mrad}) \). The polarization images expressed in terms of \( \phi \) solve the thickness of the peat with higher resolution than the \( \sigma' \) (or \( \sigma'' \)) images. Moreover, the shallow unit shows more pronounced lateral variations in the phase than in \( \sigma'' \) and patterns within the peat unit are more clearly defined: Along line By 25, the thickness of the first unit decreases from approx. 0.5 m at 2 m along the profile to 0 m around 10 m at the end of the profile. Along line By 46, the first unit is slightly thicker than 50 cm and shows the highest phase values \((~25 \text{ mrad}) \) between 0 and 6.5 m along the profile. Beyond 6.5 m, the polarizable unit becomes discontinuous with isolated polarizable \((~18 \text{ mrad}) \) zones, which also extend to a depth of 50 cm. The geometry of the polarizable shallow unit is consistent with the corresponding results along line By 68, which crosses By 25 and By 46 at 3 m along these lines (S1 and S3 are located at these intersections). In particular, it is clear that the highest phase values were consistently found in the shallowest 50 cm in the peat unit, at the depth where the presence of biogeochemical hot spots has been reported in previous studies (e.g., Frei et al., 2012).

The histograms presented in the right column of Fig. 7 show the distribution of the phase values in the images, with a different color for model parameters extracted above and below the contact between the peat and the granite. The histograms highlight that the lowest phase values clearly correspond to the granite bedrock \((< 15 \text{ mrad}) \), while higher phase values are characteristic for the peat unit.

Fig. 8 presents maps of the electrical parameters at different depths aiming at the identification of a possible link to the vegetation cover: abundant-\((av)\), moderate- \((mv)\) and sparse vegetation \((sv)\), observed at the field (Fig. 5). Such maps present the interpolation of values inverted in
Along each profile, a value is obtained through the average of model parameters (conductivity magnitude and phase) within the surface and a depth of 20 cm (shallow maps) and between 100 and 120 cm (for deep maps). The western part of the experimental plot (between 0 and 4 m in X-direction and between 2 and 9 m in Y-direction) corresponds to a shallow depth to the bedrock (a peat thickness of ~ 50-70 cm) and is associated with a high electrical response in the shallow maps ($\phi > 17$ mrad, $\sigma' > 7$ mS$^{-1}$ and $\sigma'' > 100$ $\mu$S$^{-1}$). In the Northern part of the experimental plot, in the area with abundant vegetation, we observe a higher polarization response for the top 20 cm ($\phi > 20$ mrad and $\sigma'' > 80$ $\mu$S$^{-1}$) than the one corresponding to the moderate vegetation located at the southern part. In contrast, the lowest polarization values ($\phi < 15$ mrad, and $\sigma'' < 80$ $\mu$S$^{-1}$) values are related to the area with sparse vegetation. Thus, maps in Fig. 8 suggest that the biogeochemical process controlling the polarization response correlate to the vegetation. On the contrary, the electrical conductivity ($\sigma'$) reveals only a weak dependence on the vegetation.

To support our observation regarding the electrical response and the vegetation, we present in Fig. 8j to 8l the histograms of the electrical parameters extracted at each one of the three vegetation features (abundant, moderate and sparse vegetation). These histograms show, in general, that the location with sparse vegetation, where the water can be seen on the surface, presents the lowest phase values (histogram peak at 13 mrad). Moderate vegetation corresponds with moderate phase values (histogram highest peak at 17 mrad), while the abundant vegetation corresponds with the highest phase values (histogram peak at 20 mrad) in the top 20 cm. The histogram of the three vegetation features in terms of $\sigma'$ values overlap each other. However, we observe a strong correlation between $\sigma''$ and the vegetation, with variations in $\sigma''$ permitting to discriminate between areas with a surface cover with abundant and moderate sphagnum and purple moor-grass.

3.3 Comparison of electrical and geochemical data

The pore-fluid conductivity measured in water samples retrieved from the piezometers show little variations with values ranging between 6.7 and 10.4 mS$^{-1}$. The evaluation of the imaging results measured along profiles By 25, By 46, and By 68 were used to guide the selection of locations for the freeze core and sampling of pore water and groundwater. Sampling points S1 and S3 were found in the highly polarizable parts of the uppermost peat unit (high $\sigma'$ and $\sigma''$ values). In contrast, sampling point S2 is located in an area characterized by particularly low polarization values. Figures 9a-e show the chemical parameters measured in the water samples, specifically chloride (Cl$^-$), phosphate (PO$_4^{3-}$), dissolved organic carbon (DOC), total iron (Fe$_{tot}$...
= Fe^{2+} + Fe^{3+}), and pH; whereas Fig. 9f-j show the chemical parameters measured in the peat samples extracted from the freeze cores, namely, cation exchange capacity (CEC), concentrations of iron sulfid (FeS or FeS\(_2\)), total reactive iron (Fe\(_{\text{tot}}\)), potassium (K\(^+\)), and sodium (Na\(^+\)). To facilitate the comparison of electrical parameters and geochemical data, Fig. 9k-m show the complex conductivity values (\(\sigma'\), \(\sigma''\) and \(\phi\)) at the sampling points S1, S2 and S3, which were extracted as vertical 1D profiles from the corresponding imaging results.

As observed in Fig. 6 and 7, the complex conductivity values (\(\sigma'\), \(\sigma''\)) were highest within the uppermost 10-20 cm and rapidly decreased with depth. Furthermore, the values of \(\phi\) and \(\sigma''\) in the top 20 cm at S1 and S3 are significantly higher than those at the location S2. High values of \(\phi\) and \(\sigma''\) at S1 and S3 correspond with high concentrations of DOC, phosphate, Fe\(_{\text{tot}}\) in water samples, as well as with high K\(^+\), and Na\(^+\) contents measured in soil materials extracted from the freeze cores. Fig. 9 reveals consistent patterns between geochemical and geophysical parameters: in the first 10 cm bgs close to the sampling points S1 and S3, we observe complex conductivity values (\(\sigma'\) and \(\sigma''\)) as well as chemical parameters, such as DOC, phosphate, Fe\(_{\text{tot}}\) at least two times higher than those in S2.

Figure 10 shows the actual correlations between complex conductivity and Cl\(^-\), DOC and Fe\(_{\text{tot}}\) concentrations measured in groundwater samples. In Fig. 10, we also provide a simple linear regression analysis to quantify possible correlations. Figure 10 reveals that the phase has a weak to moderate correlation with DOC, Cl\(^-\) and Fe\(_{\text{tot}}\). The conductivity (\(\sigma'\)) shows a slightly stronger correlation with the DOC, the Cl\(^-\) and total iron concentration than the polarization (\(\sigma''\)). The highest \(\sigma''\) values (>100 \(\mu\)Sm\(^{-1}\)) are related to the highest DOC and total iron concentration. Hence, it is possible to interpret the high phase values as those related to biogeochemical hot spots.

Figure 11 shows the FTIR spectroscopy analysis of the freeze core samples. The spectra show the absorbance intensity at different wavenumbers, C-O bond (~1050 cm\(^{-1}\)), C=O double-bound (~1640 cm\(^{-1}\)), carboxyl (~1720 cm\(^{-1}\)) and O-H bonds (~3400 cm\(^{-1}\)). The peaks are also indicated in Fig. 11 with the interpretation based on the typical values reported in peatlands, for instance McAnallen et al. (2018) or Artz et al. (2008).

### 4 DISCUSSION

#### 4.1 Biogeochemical interpretation

The geochemical and geoelectrical parameters presented in Fig. 6-7 and 9 displayed consistent patterns, with the highest values within the uppermost 20 cm around S1 and S3. The high DOC,
K+ and phosphate concentrations in the uppermost peat layers and especially in the areas found to be biogeochemically active, strongly suggest that there is rapid decomposition of dead plant material in these areas (Bragazza et al., 2009). Ions such as K+ and phosphate are essential plant nutrients, and phosphate species especially are often the primary limiting nutrient in peatlands (Hayati and Proctor, 1991). The presence of dissolved phosphate in pore waters suggests that (i) the plant uptake rate of this essential nutrient is exceeded by its production through the decomposition of plant material; and (ii) that organic matter turnover must be rapid indeed to deliver this amount of phosphate to the pore water. This is supported by the DOC concentrations in pore waters exceeding 10 mM. DOC is produced as a decomposition product during microbial hydrolysis and oxidation of solid phase organic carbon via enzymes such as phenol oxidase (Kang et al., 2018). Enzymatic oxidation processes are enhanced by oxygen ingress via diffusion and more importantly by water table fluctuations that work as an ‘oxygen pump’ to the shallow subsurface (Estop-Aragonés et al., 2012). Thus, an increased DOC concentration in the pore water can be used as an indicator for microbial activity (Elifantz et al., 2011; Liu, 2013). The small amount of phosphate measured in the less active area S3 can be explained by adjective transport from the active area S1, which is directly ‘up-stream’ of S2. In this case, adjective water flow through the uppermost peat layers along the hydrological head gradient may have transported a small amount of reaction products from the biogeochemical source areas to the ‘non-active’ area. The high DOC, Fe, K+ and phosphate levels confirm our initial interpretation of the highly conductive and polarizable geophysical units within the first 20 - 50 cm bgs in the surroundings of S1 and S3 as biogeochemically active areas.

The high DOC concentrations are also likely to be directly or indirectly responsible for the Fe maximum in the upper layers. Dissolved Fe was predominantly found as Fe2+ (reducing conditions) suggesting either that high labile DOC levels maintain a low redox potential, or that the dissolved Fe2+ was complexed with the DOC limiting the oxidation kinetics enough so that Fe2+ can accumulate in peat pore waters. The TRIS analysis clearly showed very low levels of sulfide minerals in both freeze cores, especially in the uppermost peat layers. This was unexpected considering the reducing conditions implied by the dominance of pore water Fe2+. We argue that the lack of sulfide minerals is due to insufficient H2S or HS− needed to form FeS or FeS2, or that the redox potential was not low enough to reduce sulfate to H2S or HS−. Both mechanisms are possible, as groundwater in the catchment has generally low sulfate concentrations and sulfate was detected in pore water samples, which would not be expected if redox potentials were low enough to reduce sulfate to sulfide. The chemical analyses do not reveal any significant or systemic vertical gradient in mineral sulfide concentrations, as
expected for the site (Frei et al., 2012). The maximum in extractable (reactive) solid phase Fe was also located in the upper most peat layer at the ‘hot spot’ S1. This Fe was likely in the form of iron oxides or bound to/in the plant organic matter. Such iron rich layers typically form at the redox boundary between oxic and anoxic zones and can be quite dynamic depending on variations in the peatland water levels and oxygen ingress (Wang et al., 2017; Estop-Aragonés, 2013).

Similar to other peatlands, the FTIR spectra show the presence of carbon-oxygen bonds such as C-O, C=O and COOH booth at S1 and S2, furthermore, the peak intensities at S1 tend to decrease with the depth, while the peak intensities at S2 samples tend to increase correlated to the IP response. The phosphate and Fe could potentially form complexes with the O-H groups that show an absorbance peak at 1050 cm$^{-1}$ (Arai and Sparks, 2001; Parikh and Chorover, 2006). Furthermore, the iron can also form complexes with the carboxyl groups (absorbance at ~1720 cm$^{-1}$).

4.2 Correlation between the peat and the electrical signatures

The two electrical units observed within the peat indicate variations in the biogeochemical activity with depth. Thus, it is likely that the uppermost unit characterized by the highest $\sigma'$, $\sigma''$, and $\phi$ values corresponds with an active biogeochemical zone, i.e., a hot spot. Consequently, the second electrical unit associated with moderate $\sigma$, $\sigma''$, and $\phi$ values is probably related to a less biogeochemical active or even inactive zone in the peat. The third unit represents the granitic bedrock. The low metal content and the well crystalized form of the granite leads to low $\sigma''$ values (here, < 40 $\mu$Sm$^{-1}$), as suggested by Marshall and Madden (1959).

The high polarization response of the peat is consistent with the measurements of McAnallen et al. (2018), who performed time-domain IP measurements in different peatlands. They suggest that the active peat is less polarizable due to the presence of the abundant sphagnum cover. They found that in the areas where the peat is actively accumulating, the ratio of the vascular plants and the non-vascular sphagnum is low, and therefore, the oxygen availability is low. However, the sphagnum is expected to exude a small amount of carbon into the peat. Fenner et al. (2004) found that the sphagnum contributes to the DOC leachate to the pore water, which is contradictory to the model of McAnallen et al. (2018). Our observations also show that the high DOC content correlates to an abundant sphagnum cover, in agreement with the observations by Fenner et al. (2004), but we cannot clearly tie this to exudates from roots or decomposition of decaying organic matter. Moreover, at our study site the abundant sphagnum grows in conjunction with abundant purple moor-grass. Figures 8-9 reveal an increase in the polarization
response ($\sigma''$, $\phi$) with both the abundant vegetation as well as with DOC concentrations measured in pore water. Thus, supporting our interpretation that high polarization values ($\sigma'' > 100 \mu\text{Sm}^{-1}$ and $\phi > 20 \text{ mrad}$) correspond to biogeochemical hot spots rather than vegetation type. Recent studies have demonstrated an increase in the polarization response due to the accumulation of biomass and activity in the root system (e.g., Weigand and Kemna 2017; Tsukanov and Schwartz, 2020). However, the sphagnum does not have roots; on the other hand, the vascular purple moor-grass can contribute to the high IP, as the roots transport oxygen into the deeper area which contributes to the increase in the polarity and normalized chargeability of the peat (McAnallen et al., 2018).

4.3 Possible polarization mechanisms

The low-frequency polarization response of subsurface materials is usually attributed to either electrode polarization of highly conductive metallic minerals (e.g., Pelton et al., 1978; Wong, 1979) or the polarization of the electrical double layer (EDL) covering charged surfaces of electrically non-conducting solid matter (e.g., Waxman and Smits, 1968; Revil and Florsch, 2010; Revil, 2012).

Electrode polarization arises from the conductivity contrast between the high electronic (or semi-conductor) conductivity of the metallic mineral and the much lower electrolytic conductivity of the surrounding pore fluid (Wong, 1979). This so-called charge polarization has its maximum at relatively high frequencies ($\gg 1 \text{ kHz}$) unless the conducting minerals are large (Bücker et al., 2018). If charge-transfer reactions take place at the mineral-electrolyte interface, e.g., due to the presence of metallic ions in the electrolyte, the charge polarization is accompanied by a concentration polarization, which potentially shifts the maximum of the electrode-polarization response into the low-frequency range (Bücker et al., 2018). The magnitude of electrode polarization is proportional to the volume content of the conductive minerals (Wong, 1979).

In this study, we have found a strong correlation between the polarization response ($\phi$ and $\sigma''$) and Fe\text{tot} in the solid phase and a less pronounced correlation between the polarization response and the concentration of dissolved iron in the liquid phase (see Fig. 10). If the iron in the solid phase occurred in the form of highly conductive minerals, the two above correlations would clearly point to an electrode-polarization mechanism as a possible explanation for the observed response. Previous studies (e.g., Flores Orozco et al., 2011; 2013) imply that iron sulfides (FeS or FeS$_2$) in sediments polarize as long as sufficient Fe$^{2+}$ cations are available in the pore water. However, in the case of the present study, the lack of sulfide, and the rather high Eh (inferred
from presence of sulfate) in the pore water, do not favor the precipitation of conductive sulfides such as pyrite. Under these conditions, iron would rather precipitate as iron oxide or form iron-organic matter complexes. The electrical conductivity of most iron oxides is orders of magnitude smaller than the conductivity of sulfides (e.g., Cornell and Schwertmann, 1996), and is thus too low to explain an increased electrode polarization. The only highly conductive iron oxide is magnetite, with a conductivity similar to pyrite (Atekwana et al., 2016). Consequently, the presence of magnetite could explain the electrode polarization. However, the low pH (~5) typical for peat systems does not favor the precipitation of magnetite, but rather less conducting, and thus, less polarizable iron (oxy)hydroxides such as ferrihydrite (Andrade et al., 2010; Linke and Gislason, 2018). As indicated by the FTIR analysis, the iron might furthermore have built complexes with the carboxyl (absorbance at ~1720 cm\(^{-1}\)). However, it is not clear if these iron complexes might have a sufficiently high electrical conductivity and - in the fashion of conductive iron minerals - permit electrode polarization mechanism.

In addition to the polarization of conductive minerals, the polarization response can also be caused by the polarization of the electrical double layer (EDL). The EDL consists of an inner Stern layer of cations adsorbed to the charged solid surface and an outer diffuse layer. Together, these two layers screen the (usually) negative charge of the solid surface and represent a highly conductive and, thus, polarizable surface layer. Depending on the specific geometry of solid and liquid phases, EDL polarization can be understood in terms of the polarization of isolated EDL-covered grains (e.g., Schwarz, Schurr, Leroy et al., 2008) or in terms of membrane polarization linked to EDL-covered pore constrictions (e.g., Marshall and Madden, 1959; Bückner and Hördt, 2013; Bückner et al., 2019). Regardless of the specific mechanism, all EDL polarization mechanisms strongly depend on the specific surface area of the material and the charge density at the surface (Revil, 2012; Waxman and Smits, 1968).

The product of both surface charge density and surface area can be quantified by the Cation Exchange Capacity (CEC) of a material. As peat mainly consists of organic matter known to have a high CEC (even compared to most clay minerals; e.g., Schwartz and Furman, 2014; and references therein), EDL polarization of charged organic surfaces may explain the observed IP response. However, in this case, we would expect an increase of the polarization magnitude with the CEC, which we did not observe in our data. The CEC values of the freeze core samples all vary within a narrow range of 5 to 25 meq/kg and we did not observe any correlation between CEC and changes in the polarization magnitude (\(\sigma''\), \(\phi\)). Ponziani et al. (2011), who conducted SIP measurements on a set of peat samples, have reported a similar observation previously. In summary, the measured CEC is high enough to explain a rise in EDL-polarization; however,
the (small) variation in CEC does not explain the observed variation in the polarization magnitude.

The pH of the pore fluid is also known to control the magnitude of EDL polarization; an increase of pH usually corresponds with an increase of the polarization magnitude (e.g., Skold et al., 2011). At low pH values, H\(^+\) ions occupy (negative) surface sites and thus reduce the net surface charge of the EDL (e.g., Hördt et al., 2016; and references therein). Our data seems to show the opposite behavior: we found a lower pH in the highly polarizable anomalies at S1 and S3 compared to site S2 (the inactive and less polarizable location); while the pH increases at depth for decreasing values in the polarization (both \(\sigma''\) and \(\phi\)). At the same time, variations in pH are within the range 4.45 and 5.77 and thus might not be sufficiently large to cause the observed variation in the polarization response.

Besides pH, pore fluid salinity plays a major role in the control of EDL polarization. Laboratory measurements on sand and sandstone samples indicated that an increase in salinity leads to an early increase of the imaginary conductivity, which is eventually followed by a peak and a decrease at very high salinities during later stages of the experiments (e.g., Revil and Skold, 2011; Weller et al., 2015). Hördt et al. (2016) provided a possible theoretical explanation of this behavior: In their membrane-polarization model, salinity controls the thickness of the (diffuse layer) of the EDL and depending on the specific geometry of the pores; there is an optimum thickness, which maximizes the magnitude of the polarization response. In the present study, we observed that an increase in salinity (as indicated by the high Cl\(^-\) concentrations within the uppermost 10 cm at all sampling locations) is associated with an increase in the polarization magnitude response (e.g., Revil and Skold, 2011; Weller et al., 2015; Hördt et al., 2016). However, the highest Cl\(^-\) concentrations were observed for the shallow layers at location S2, where we measured lower polarization magnitudes (in terms of \(\sigma''\), \(\phi\)) compared to S1 and S3.

The strong correlation between the polarization response and the DOC suggest an, as yet not fully understood, causal relationship. A similar observation has recently been reported by Flores Orozco et al. (2020), who found a strong correlation between DOC as a proxy of microbial activity and both \(\sigma'\) and \(\sigma''\) in a municipal waste landfill in Austria. McAnallen et al. (2018) reported a strong correlation between the occurrence of long-chained C=O double bonds and the total chargeability of peat material. The upper peat layers are exposed to oxygen leading to oxidation of the peat and formation of C=O double bonds at solid phase surfaces and in the pore water DOC. Such long-chained organic molecules have an increased polarity and thus more
readily attach (or even form at organic matter surfaces) to the surface of solid organic and mineral particles (Alonso et al., 2009). Based on a membrane-polarization model, Bücker et al. (2017) predict an increase of the polarization magnitude in the presence of wetting (i.e., long-chained) hydrocarbon in the free phase. The long-chained polar DOC attaches to the peat surface, similar to polar hydrocarbon, and so it might provide extra surface charge, thus reducing the pore space and causing membrane polarization (Marshall and Madden, 1959).

5 Conclusions

We investigated the applicability of induced polarization (IP) as a tool to identify and localize biogeochemically active areas or “hot spots” in peatlands. IP imaging results revealed a high polarization response for areas that based on the solid and solution phase chemistry can be interpreted as biogeochemically active. The chemical analysis of the groundwater samples and the freeze core indicate that the top 10-20 cm correspond to the highest geochemical activity, whereas the deeper areas of the peat are less active. Accordingly, only the upper 10-20 cm can be defined as biogeochemical hot spots. Moreover, our geophysical survey shows that IP imaging can delineate the geometry of the hot spot with high spatial resolution. In this regard, indicators for increased biogeochemical activity (DOC, phosphate) correspond to a high IP response (σ’’ and phase values).

Although the exact polarization mechanism is not fully understood, our results reveal that the IP response of the peat changes with the level of biogeochemical activity. Thus, the IP method is capable of distinguishing between biogeochemical active and inactive zones within the peat. The phase values show a contrast between these active and inactive zones and characterize the geometry of the hot spots even if iron sulfides are not present. Previous studies have only investigated the conductive properties of the peat; however, biogeochemical activity within the peat has not been investigated. In this regard, our study shows a new alternative method to characterize biogeochemical changes and their geometry within the peat with high resolution.

For further improvement and understanding of the effect of the organic matter such as DOC or phosphate on the polarization response, further laboratory studies on peat samples with different concentration and mixture of DOC, phosphate and iron in the pore fluid are required.

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**Data availability**

All data are available from the corresponding author upon request.

**Competing interests.**

The authors declare that they have no conflict of interest.

**Author contribution**

AFO and TK designed the experimental set-up, TK conducted the field survey and analysis of
the geophysical data. BG and SF conducted the geochemical measurements and their
interpretation. AFO, MB and TK interpreted the geophysical signatures. TK lead the
preparation of the draft, where SF, BG, MB and AFO contributed equally.
Figure 1: (a) General overview of the experimental plot located in the Fichtel Mountains and (b) structure of the bowl shaped Lehstenbach catchment, and (c) location of the experimental plot.
Figure 2: (a) Panoramic overview of the study site and the measurement setup. Pictures show the experimental setup and differences in the vegetation density between the northern and southern part of the experimental plot. The spectral induced polarization (SIP) lines appear distorted due to the panoramic view. (b) Sphagnum in the northern part of the experimental plot. (c) Coaxial cables and stainless steel electrodes used for SIP measurements. (d) Vegetation and the coaxial cable bundle used for IP measurements at the water covered area in the southeastern part of the experimental plot. (e) The freeze core shows the internal structure of the peat.
Figure 3: Schematic map of the experimental plot. The solid lines represent the measured profiles; the bold lines represent the position of the profiles discussed in this manuscript (By 25, By 46 and By 68). The arrows indicate the ground water flow direction. The points represent the locations of fluid (S1, S2 and S3) and freeze core (S1, S2) samples as well as the position of piezometric tubes, where the water level was measured.
Figure 4: Raw data analysis. Raw-data pseudosections of (a) the apparent resistivity and (b) the apparent phase shift for measurements collected along profile By 25. Histograms of the normal-reciprocal misfits of (c) the measured resistance and (d) the apparent phase shift.
Figure 5: Variations in the thickness of the peat layer, i.e., depth to the granite bedrock. The positions of the three selected IP profiles By 25, By 46, and By 68 are indicated (solid lines) as well as the position of the sampling points and the geometry of the three classes of vegetation cover: abundant vegetation (av), moderate vegetation (mv), and sparse vegetation (sv).
Figure 6: Imaging results for data collected along profiles By 25 (a-b), By 46 (c-d), and By 68 (e-f) expressed as real $\sigma'$ and imaginary $\sigma''$ component of the complex conductivity. The dashed lines represent the contact between the peat and granite; the black dots show the electrode positions at the surface. The vertical lines represent the location of the fluid ($S_1, S_2$ and $S_3$) and freeze core ($S_1, S_2$) samples.
Figure 7: Imaging results for data collected along profiles By 25 (a), By 46 (c), and By 68 (e), expressed as phase values $\phi$ of the complex conductivity. The dashed lines represent the contact between peat and granite; the black dots show the electrode positions at the surface. The vertical lines represent the location of the fluid (S1, S2, S3) and freeze core (S1, S2) samples. The histograms represent the phase values of the granite and peat extracted from the imaging results in (6b, 6d, 6f) according to the geometry of the dashed lines.
Figure 8: Maps of the complex conductivity at different depths. The black lines indicate the profiles By 25, By 46, and By 68. The dots represent the locations of the vertical sampling profiles S1, S2, and S3. The white lines outline areas classified as (av) abundant vegetation, (mv) moderate vegetation, (sv) sparse vegetation, and histograms of the complex-conductivity imaging results of the masked areas, the abundant vegetation (red bins), the moderate vegetation (green bins) and the sparse vegetation (blue bins).
Figure 9: Results of geochemical analyses of water and soil samples. Fluid-sample analysis of the (a) chloride Cl\(^-\), (b) phosphate PO\(_4^{3-}\), (c) dissolved organic carbon, (d) total iron Fe\(_{\text{tot}}\), and (e) pH. Freeze-core sample analysis of the (f) cation exchange capacity CEC, (g) iron sulfide FeS\(_2\), (h) total iron Fe\(_{\text{tot}}\), (i) sodium Na\(^+\), and (j) potassium K\(^+\). Imaging results at the three sampling locations in terms of (k) real component \(\sigma'\), (l) imaginary component \(\sigma''\), and (m) phase \(\phi\) of the complex conductivity.
Figure 10: Correlations between the geophysical and geochemical parameters, phase (ϕ), the real (σ') and imaginary (σ'') component of the complex conductivity (retrieved from the imaging results) and the biogeochemical analysis, expressed in terms of the dissolved organic carbon (DOC), and chloride (Cl\(^{-}\)) content from the pore fluid samples and total iron (Fe\(_{tot}\)) content from pore fluid in μmoll\(^{-1}\) and solid samples in μmolg\(^{-1}\). The correlation coefficients of least square regressions analysis are shown in the top left corners of the subplots.
Figure 11: Fourier transform infrared (FTIR) spectroscopy of the freeze core samples collected at S1 (left panel) and S2 (right panel). Each sample was extracted from the 10 cm segments. The lines represent the depth at every 10 cm between 0 and 80 cm below ground surface. The relevant peaks show the absorbance intensity, the interpretation is based on Artz et al. (2008), Arai and Sparks (2001), Parikh and Chorover (2006).

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