Geochemical Variability of Surface Sediment in Post-Mining Lakes Located in the Muskau Arch (Poland) and Its Relation to Water Chemistry

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Abstract Water and sediment chemical compositions were investigated in 75 post-mining lakes in the eastern part of the Muskau Arch region (Western Poland, Central Europe). The lakes, originating from lignite and clay surfaces and underground mining activity, exhibited a wide gradient of physical and chemical water parameters. The most pronounced variability in water parameters was observed in pH values, conductivity, and oxygen concentration in the mixolimnion and monimolimnion. In addition, the lakes varied in terms of their trophic status, indicated by their dissolved organic carbon (DOC) and phosphorous concentrations. In general, lakes with higher pH values had higher trophic states, and some even exhibited hypertrophic conditions. The Monte Carlo test indicated that the water pH, phosphorus, DOC, and silicon concentrations could significantly explain the chemical characteristics of sediments.

Keywords Lake sediments · Coal mining · AMD · Aluminium · Iron

1 Introduction

Coal mining impacts the natural environment in many ways, including landscape changes, waste production, water, soil and air contamination, water table changes, and noise pollution (Chadwick et al. 1987; Tiwary 2001; Younger 2004; Bian et al. 2010). The negative impacts of mining were generally neglected until the 1980s. Since the 1980s, multiple mines that strongly affected the environment have been closed. Unfortunately, this has failed to stop the negative effects on the ecosystem, and new hazards have arisen. One of the most significant hazards is the weathering of coal and its host rocks, contributing to the acidity of the ground and surface waters (acid mine drainage (AMD)) (Blowes et al. 2005). AMD is a major water contaminant in numerous post-mining areas in Europe and globally (Williamson et al. 2006). This process can intensify in the near future, leading to several ecological challenges and water unavailability for municipal and industrial purposes (Younger 2002). In addition, AMD has a major impact on lake water and surrounding soils. AMD-affected lakes are commonly known for their high conductivity and increased concentrations of metals such as iron and aluminium (Geller et al. 2013). Acidification of surface waters results in a decline in fish diversity and population. In water bodies impacted by AMD, fish are unable to survive if pH drops below 5. Moreover, AMD negatively impacts the diversity and development of aquatic plants and animals (Katz 1969; Sienkiewicz and Gąsiorowski 2016, 2019).

Lake water at sites affected by mining reacts with minerals in the surrounding rocks and soils. Some of these minerals act as a pH buffer. This leads to natural neutralisation of AMD-affected pit lakes. In general, minerals that play a role in natural neutralisation of
acidic lakes include calcite, siderite, aluminosilicate, clay, aluminium hydroxysulphate, aluminium and iron hydroxide, and ferrihydrite (Jurjovec et al. 2002; Gunsinger et al. 2006). Although different minerals buffer the solution in different pH ranges (Blowes and Pateck 1994), the carbonate system greatly impacts the buffer intensity and the neutralising capacity of water (Langmuir 1997). In addition to the buffering capacity of minerals in bedrock, the neutralisation process of pit lakes depends on many other factors. One of these factors is the age of such lakes and the exposure of sulphides, which are present in the surrounding rocks, to water and aerobic conditions. Although natural neutralisation often begins several decades after the creation of a lake, it can progress relatively quickly (Sienkiewicz and Gąsiorowski 2016).

One of the biggest regions in Central Europe affected by AMD is the Muskau Arch located on the Polish–German border. Over 100 lakes can be found in post-mining pits that developed in former lignite and clay mines. The lignite deposits in the area are of the Oligocene age and were originally deposited at depths of 100–200 m (Lutyńska and Labus 2015). Later, they integrated with glacial deposits of large marginal moraine from the Elsterian glaciation (Urbański 2007). The oldest lignite mines in the region began operating in the 1840s, with some mines closing as early as the end of the nineteenth century (Solski et al. 1988). The closed mines were usually immediately flooded with ground and meteoric waters. Lignite contains large amounts of iron sulphides, for example, pyrite, which easily oxidises in contact with oxygen dissolved in infiltrating water as well as atmospheric oxygen. This process led to the production of sulphuric acid, which resulted in the strong acidification of many anthropogenic lakes in the region (Williamson et al. 2006). The exploitation of lignite in the Polish part of the region ended in 1973 when the ‘Babina’ mine located in the east of Leknica stopped operating (Koźma 2005). Since then, multiple lakes in the Muskau Arch still show signs of AMD. This is especially apparent in younger lakes, formed 44–49 years ago, exhibiting higher mineralisation and acidity. In addition, the chemical composition of the water from these lakes is characterised by the presence of dissolved compounds of S, Si, and Fe, which are oxidation products of minerals (e.g. pyrite) from the surrounding rocks (Pukacz et al. 2018). Hence, the oxidation of sulphides has a strong impact on water chemistry, which is greater than that of the dissolution of rock-forming minerals (Lutyńska and Labus 2015). Furthermore, research has shown that the parameters and concentrations of ion such as Eh, SO₄²⁻, Ca²⁺, Mg²⁺, Na⁺, K⁺, Fe₃⁺, Mn⁵⁺, and Al⁺⁺ increase with the declining pH levels of lake water (Lutyńska and Labus 2015). Lakes with higher acidity (pH < 4) are characterised by higher concentrations of dissolved Fe, Ca, and Mg. High concentrations of Ca and Mg are due to the dissolution of Ca²⁺- and Mg²⁺-containing minerals (e.g. carbonates silicates) by acidic water (Solski et al. 1988).

This study analysed the chemical composition of water and surface sediments in the 75 lakes situated at the Muskau Arch, with the aim of providing an update on water chemistry studies conducted in the 1980s (Solski et al. 1988) and to further explain the chemical diversity of these lakes. Employing a dataset on water and sediment chemistry, we attempted to find relationships between specific parameters. We hypothesised that some water parameters in post-mining lakes, such as pH, trophic state, or presence of summer stratification, impacted the composition of bottom sediments. We tested this hypothesis using multivariate analyses and determined the relationship between specific water parameters or elemental content in the water and evaluated water parameters that could explain the chemical composition of the sediment. Understanding the relationship between water and sediment chemistry can help in devising policies for the remediation/protection of post-mining lakes.

2 Material and Methods

2.1 Study Area

The studied lakes are located in the Muskau Arch in the western part of Poland (Central Europe), close to the border with Germany (Fig. 1). The Muskau Arch is an arch-shaped glaciotectonic moraine formed during the Last Glacial Maximum. This structure is composed of a thick bed of glacial till and some clay and lignite from the Tertiary Period. Lignite occurs as strongly faulted horizons or lenses (Urbański 2007; Lutyńska and Labus 2015). The glacial origin of the structure resulted in a characteristic geomorphology, which was extensively modified by mining activity. Lignite was extracted by the subsurface or surface mining method. Several lakes in the Muskau Arch formed as a result of lignite and clay mining (Solski et al. 1988).
Post-mining lakes in the region can be divided into two types: depressions created over underground passages and chambers after their collapse and water bodies filling open-mine pits. Lakes of the first type are usually narrow and long, with very steep shores. Lakes of the second type are usually larger and have an irregular shape. In addition, natural lakes related to lignite deposits are also present in the region. These natural lakes formed owing to the weathering of lignite beds and the subsequent lowering of the terrain surface. These basins are relatively small and shallow (Koźma 2005).

2.2 Fieldwork

Samples were collected in June and September 2013 and June 2014. The surface sediments were collected from 75 lakes (Fig. 1) at their deepest sites using a Kayak-type gravity corer with a polycarbonate pipe that had a diameter of 55 mm. Only the top 2-cm-thick layer, representing the last 1–4 years (Sienkiewicz and Gąsiorowski 2016, 2017, 2018, 2019), was sampled for geochemical analysis. Sediment samples were stored in plastic bags under dry and cold (4 °C) conditions. Water parameters (temperature, conductivity, pH, dissolved oxygen (DO), and dissolved organic carbon (DOC)) were measured in situ for each sampled lake, using multiparameter MultiLineP4 SET manufactured by WTW, both at the surface and near the bottom, 0.5 m above the lake bed. In selected lakes (e.g. site 61), the water parameters were measured every 1 m to investigate thermal and chemical stratification. Water samples for laboratory analyses were collected from the top water layer at a depth of 0.5 m using a 5-L vertical water sampler. Further, the water samples were filtered through a membrane filter with pore size of 0.45 μm, stored in 50 mL polyethylene bottles, and transported to the laboratory in a portable fridge under cold (4 °C) conditions.

2.3 Surface Sediment and Water Analysis

Sediment samples were weighed and dried at 65 °C for 48 h to determine the bulk density and water content. Dry, grounded, and homogenised sediment samples were analysed for selected elements (Ca, Al, Fe, Mn, P, total nitrogen (TN), total organic carbon (TOC), total hydrogen (TH), and total sulphur (TS)). For elemental analysis, the sediment samples were pre-treated with 10% hydrochloric acid to dissolve carbonates. Subsequently, TOC, TN, TH, and TS contents of the pre-treated samples were determined using the elemental analyser Vario MicroCUBE at the Institute of Geological Sciences PAS in Warsaw. Sulphanilic acid was used as a standard for analysis, and the typical analytical error was smaller than 0.1%.

For metal analysis, sediment samples were dissolved with a mixture of 65% nitric acid and 35–38% hydrochloric acid (3:1) in the microwave mineraliser Magnum V2. Filtered water samples were additionally acidified using 1 mL of 65% ultra-pure nitric acid. The TOC in the water samples was measured using the Shimadzu carbon analyser TOC-5000 1997, SS14-5000. The nitrogen concentration in water samples was determined using the Kjeldahl method. The contents of Na and K in the sediment and water samples were investigated using flame atomic emission spectroscopy; Ca and Mg were investigated using flame atomic absorption spectroscopy (Qz 989 SOLAAR by Unicam), and Fe, Si, Mn, Al, and P were measured using inductively coupled plasma–optical emission spectrometer (Varian Vista-PRO). All measurements were performed at the Institute of Environmental Protection-National Research Institute in Warsaw.

2.4 Numerical Analysis

For data analysis, we used the principal component analysis (PCA) and redundancy analysis (RDA). Prior to the multivariate analysis, 15 water variables (water temperature, DO, pH, conductivity, DOC, Ca, Mg, Na, K, Fe, Mn, Al, P, Si, and TN) were standardised using logarithms. This precluded effective visualisation in the dendrogram and scatter biplot. PCA was performed to examine the pattern of variation in the surface and deep water chemistry. RDA was employed to explain the pattern of variation in lake sediment chemistry using water parameters as explanatory variables. The relative importance of explanatory variables was determined using forward selection of the RDA. A Monte Carlo test with 999 unrestricted permutations under a reduced model was run to verify the environmental variables that could significantly explain the variation of the dependent variables. Multivariate analyses were performed using Canoco software (version 5; ter Braak and Šmilauer 2012).

3 Results

3.1 Water Chemistry

Field measurements of water parameters highlighted the strong variability of water pH (Suppl. 1). The pH of the
lake water varied from 2.52 (lake no. 58) to 8.80 (lake no. 68). The lowest pH was found in lakes formed as a result of lignite mining (2.5–6.0) and the highest was found in post-clay-pit lakes (6.7–8.8) and some natural shallow lakes (> 7.3). The collected water samples varied in terms of conductivity, which ranged from 81 μS cm⁻¹ (site 29) in lakes with circumneutral waters to 2430 μS cm⁻¹ (site 59) in lakes with acidic water. Water samples from acid lakes were enriched in Fe, Al, Si, Ca, and Mg (e.g. sites 57, 58), while in samples from alkaline lakes, the concentrations of these elements were significantly lower (e.g. sites 31, 32, 33, 34).

In lake 61, where clear stratification was observed, pH, DO, conductivity, and temperature were measured in the water column at 1-m intervals (Fig. 3). Based on the measurements, we determined the presence of the chemocline between 6 and 7 m. At this depth, only temperature changed significantly, but the pH, conductivity, and DO remained almost the same. Below 6 m, the DO began to decline from 7 mg L⁻¹, and at 7 m, it
was below the detection limit. At 6 m, the pH and conductivity started to increase, with pH increasing from 3.04 to 6.52 and conductivity increasing from 1855 to 3070 \( \mu \text{S cm}^{-1} \).

PCA was used to determine the factors that contributed the most to the variability in the chemical composition of surface waters (Fig. 2). All environmental variables (water depth, temperature, DO, pH, conductivity, DOC, Ca, Mg, Na, K, Fe, Mn, Al, P, Si, and N) were included in the analysis. We present results only from the first (PC1) and second (PC2) components (Fig. 2a), which explained 45.9% and 12.7% of the variation in the dataset, respectively. PC1 can be primarily explained by the changes in water mineralisation, expressed as conductivity, both in surface and near-to-bottom water (\( r = 0.98 \) and 0.90, respectively), and the concentration of the main cations. Moreover, changes in water pH can be observed along the first PC axis (\( r = -0.85 \)). The site distribution along PC2 can be interpreted as the variation in nutrient content (i.e. DOC and P; \( r = 0.64 \) and 0.69, respectively) and DO concentration (\( r = -0.60 \)).

The sites on the PCA plots were divided into two groups with respect to water pH (acidic and neutral/alkaline lakes) with a limit value of 5 (Fig. 2b). In addition to grouping according to water pH, three subgroups can be distinguished: (1) lakes with high total mineralisation (expressed as high conductivity, above 1700 \( \mu \text{S cm}^{-1} \), and high cation content); (2) relatively deep lakes with high oxygen content in near-to-bottom water; and (3) eutrophic lakes with dissolved carbon content (> 12 mg L\(^{-1}\)) and phosphorus (> 35 \( \mu \text{g L}^{-1} \)).

### 3.2 Sediment Chemistry

The collected sediments were classified as sand, mud, sapropel, and detritus gyttja with varying amounts of organic matter. The major and trace element contents, namely, Al, Ca, Fe, Mn, P, TOC, TH, TN, and TS, were determined in all samples (Suppl. 2). The most prominent feature of the dataset is the high variability of Al and Fe concentrations in sediments. The concentrations of these elements were negatively correlated at many sites, that is, lakes with a high concentration of Al in sediments (e.g. sites 41, 50, 71) had low Fe concentrations, and conversely, sites with high concentrations of Fe (3.5, 23, 51, 60) had low Al concentrations (< 10 mg kg\(^{-1}\)). This relationship is unclear for samples with intermediate concentrations of both elements, resulting in a low overall correlation (\( r = 0.43 \)) between these elements for all datasets (Fig. 4). In the studied sediments, Mn and Ca concentrations were positively correlated, as were the concentrations of P, C, and N. Moreover, the concentration of elements indicating the organic matter content strongly varied between lakes. The highest concentrations of TOC (10–20 wt.% of dry mass) and TN (up to 3% of dry mass) were recorded in lakes from the middle part of the region. The absolute maximum TOC (approximately 50 wt.% of dry mass) was recorded at site 55, comprising sandy sediments full of lignite clasts. The C/N ratio for most samples varied between 10 and 20 (Fig. 5). Several samples had C/N ratios in the range of 20–35, and one had a high C/N ratio of 78 (site 55).

### 3.3 Explanation of Variations in Lake Sediment by Water Parameters

The relationship between the sediment and the top-layer water chemistry was tested using ordination techniques. Environmental variables with a large variance inflation factor (VIF > 10), which were conductivity and Ca concentration, were excluded from the analysis. Because of the relatively short gradients (0.2 SD) in the response dataset, RDA with forward selection was conducted on the sediment data using water parameters as explanatory variables (Fig. 6). The test indicated that only four variables, pH, P, Si, and DOC content, were statistically significant (\( p < 0.05 \)). Together, these variables explained 23.3% of the variance in the sediment dataset. According to the RDA results, the concentrations of organic matter, Ca, and Mn in sediments increased with P content in water. The contents of TOC, TN, and TS in sediments were higher in lakes with a high pH and DOC content in water. Sedimentary Fe and Al varied along RDA axis 2, and sedimentary Fe increased with increasing Si concentration in water.

### 4 Discussion

#### 4.1 Factors Determining Water Chemistry

The chemistry of water and sediments in the anthropogenic lakes located in the Muskau Arch region was influenced by multiple factors, including the origin of the lake (i.e. coal versus clay mining), the age of the lake (Pukacz et al. 2018), hydrogeological and geological...
conditions, presence of lignite, pyrite content (Labus and Skoczyliska 2013), and ore geometry. This results in high variability in lake water and sediment chemistry, even when comparing closely located sites (Sienkiewicz and Gąsiorowski 2019). Moreover, the complex geological structure of glaciogenic sediments containing lignite makes the changes in the chemistry of the studied lakes difficult to model and predict (Sienkiewicz and Gąsiorowski 2016).

The parameters that significantly differentiate the studied lakes were water conductivity and pH. Moreover, few factors determine the pH of the studied lakes. Lakes that originate as a result of lignite exploitation usually have a lower pH than post-clay-pit and natural lakes. This is caused by the oxidation of sulphide compounds present in lignite deposits (Blowes et al. 2005).

An important factor influencing the state of water bodies created by lignite mining is their age (Sienkiewicz and Gąsiorowski 2016; Pukacz et al. 2018). It is known that the AMD-affected lakes have the ability to neutralise water due to the properties of the clay minerals present in the bedrock (Sienkiewicz and Gąsiorowski 2018), the slow weathering of glacial till containing Ca, and the inflow of water with high mineralisation (Stottrmeister et al. 1999). However, this is a long process, and in the Muskau Arch, this process is controlled by the time of closure of the lignite mines. The oldest lakes, formed at the beginning of the twentieth century, are located in the northern part of the area near Tuplice (Dawczyk and Maciantowicz 2014). The pH of these lakes was significantly higher (> 6) than that of the lakes in the central and southern parts of the region (Fig. 2b). The mine closure progressed from north to south. The youngest lakes are on the southern side of the studied area near Leknica. These lakes (sites 38–65, except sites 41 and 52), formed in the second half of the twentieth century, are characterised by low pH (2.5–4), as reported by Lutyńska and Labus (2015). This is owing to the lack of time required for the natural neutralisation of younger water bodies. However, the increased concentration of Ca in acidic lakes proves the occurrence of the neutralisation process. CaCO₃ present in the glacial tills (CaO content 1.8–3.1%; Maziarz 1980) dissolves in the presence of H⁺ ions and together with Mg ions (r = 0.76) is transported to the lakes. Artificial introduction of calcium hydroxide to a lake is another way of accelerating lake neutralisation (Feng et al. 2019), which was done in the case of lakes 14, 16, 17, 27, 28, and 31 (Oszkinis-Golon et al. 2020). In some other lakes (e.g. lake 10), wastewater was introduced to ensure better conditions for fish breeding.

The high variability in the geological structure of the Muskau Arch resulted in varied water chemistry parameters, even in lakes in close proximity, for example, sites 40 and 42 versus 41. Site 41 with slightly alkaline water (pH = 7.44) is close to a nearby area with a thick layer of clay, the exploitation of which led to the formation of this lake. It is located only 120 m from lakes 40 and 42,
which had water pH levels of 3.16 and 3.13, respectively. Moreover, even in a single lake, stratification in the water column can result in two chemically different water layers. This process occurs mainly in relatively deep lakes with wind action limited by the direction of their longer axis perpendicular to the dominant wind direction (which in the studied region is west–east). These lakes were stratified based on water density. This is caused by a pronounced temperature gradient caused by changes in weather conditions at the surface of the lakes. This leads to the formation of a thermocline and allows deeper layers that are isolated from the atmosphere to develop chemical conditions different from those in the mixolimnion. Thermal stratification can be strengthened by an inflow of highly mineralised groundwater of a high density directly to the hypolimnion or monimolimnion (Schultze et al. 2016). Examples of such stratification were found in sites 38 (site 24 in Lutyńska and Labus 2015), 54, and 61 (Fig. 3). In these acidic lakes, the hypolimnion water, depleted in DO, had relatively high pH and conductivity compared to the surface water. This is due to the consumption of oxygen in the near-to-bottom zone by the decomposition of organic matter and reduction of Fe, leading to its dissolution (Boehrer and Schultze 2009). In the mixolimnion and chemocline zones, DO in water results in Fe and S oxidation and, consequently, a decrease in pH. A similar pattern of high pH in the monimolimnion has been observed in stratified acidic pit lakes in other regions (Moser and Weisse 2011; Hrdinka et al. 2013).

In general, acidified water has higher total mineralisation and electroconductivity. Hence, the water conductivity of lignite mining pit lakes is much higher than that of post-clay-pit lakes or natural lakes in the studied area and other regions of Poland (Kostka and Leśniak 2020). This is caused by the hydrolysis of ferric iron during which H\(^+\) ions are released, affecting the carbonate buffering system, lowering pH, and increasing conductivity (Schultz 2013). Furthermore, the concentration of specific metals is usually higher in acidified water, for example, Fe and Al have higher concentrations at low pH (España et al. 2008). Despite this general trend, the concentration of some elements was not significantly correlated with water pH. The Al concentration is high in water with a pH below 4. This pH is the point below which Al concentration significantly increased. The Fe concentration is also related to pH and is the highest in low pH waters, but decreases gradually rather than abruptly with an increase in pH (González et al. 2020). The Mn concentration is also higher in waters below pH 4, which is due to the release of Mn ions from the reductive dissolution of Fe and Mn hydroxides (Torres and Auleda 2013; Ählgren et al. 2020). Both ferric iron and manganese can still be dissolved in circumneutral waters due to the high solubility of Fe\(^{2+}\) and Mn\(^{2+}\) under anoxic conditions that occur in the hypolimnion in some lakes (e.g. site 38 or 61) or even near-to-surface water of extremely shallow lakes (e.g. site 14.5).

The concentrations of P and DOC in water are good approximators of the trophic state of lakes. The highest concentrations of P and DOC were recorded in shallow natural and post-mining lakes (e.g. site 68, 4). These lakes are characterised by strong green algae blooms during the summer despite their shallow depth and oxygen depletion. In addition, zooplankton assemblages indicate eutrophic or hypertrophic conditions (Sienkiewicz and Gąsiorowski 2017, 2019).
4.2 Sediment Chemistry in Relation to Water Parameters

The chemical composition of sediments relies on various reactions occurring in lake ecosystems. It is primarily controlled by reactions occurring in water and detrital material transported to the lake with runoff, groundwater, mass movements, or wind action. The Monte Carlo permutation test indicated that pH, Si, P, and DOC could explain the chemical composition of the sediment samples ($p < 0.05$).

The chemical reactions in the studied lakes, including sediment precipitation/dissolution processes, depend highly on water pH. This relationship is evident in the distribution of Fe and Al in surface sediments. In general, although the studied sediments displayed high concentrations of Fe and Al, lake sediments exhibiting high concentrations of Fe often had low concentrations of Al and vice versa. The RDA results showed that Fe concentrations in sediments increased with low pH (e.g. at sites 22, 51, 57, and 58). This is possibly related to the precipitation of schwertmannite, goethite, jarosite, and gypsum, which were recognised in lake sediments in this region (Rzepa and Bożęcki 2007). Sites with a high concentration of Fe are more common in the studied area, especially in the southern part (e.g. sites 22, 51, 57), whereas lakes with sediments enriched in Al are found mainly in the northern and central parts of the Muskau Arch (e.g. sites 35, 69, 71). Al in lake sediments possibly results from the precipitation of kaolinite and illite (Bożęcki 2013). These lakes are mostly older or were created close to former clay pits and are characterised by circumneutral (pH = 6.5–7.5) waters. This negative correlation results from the characteristic geochemical nature of these elements. Ionic species distribution in water for both Fe and Al depends on pH. These metals can exist in-lake primary production (C/N < 20) and orange bars indicate lakes with dominant allochthonous organic matter originating mainly from the lake catchment. Number of lakes are shown in Fig. 1.
in water in many different forms such as sulphates, hydroxides, or hydroxysulphates. When the pH of the water increases above 2, Fe compounds begin to precipitate due to hydrolysis, unlike Al compounds that tend to be hydrolysed and precipitated at pH 4.5–5.0 (España et al. 2008). Lower concentrations of Fe in sediments also occur in lakes with relatively high Fe content in the water, as well as with thermal stratification and oxygen depletion in the hypolimnion (e.g. sites 14.5, 54, and 57). The water in the bottom layer of these lakes exhibits low concentrations of DO, which indicates anaerobic conditions and the reduction of Fe$^{3+}$ to Fe$^{2+}$ ions. As Fe$^{2+}$ ions are more soluble than ferric Fe, the reduced Fe dissolves, enriching the water in this zone and preventing most of the Fe from precipitating into the sediments (Hem and Cropper 1962). This process results in no correlation between the Fe concentration in water and sediment ($r = 0.03$).

Although Si was identified in the RDA as one of the drivers controlling sediment chemistry, its relation to sediment parameters is unclear. The Si in freshwater originates from the weathering of silicate minerals, mainly feldspars. This process occurs mainly in the soil layer under aerobic conditions and is catalysed at low (< 4) and high (> 8) pH (Sverdrup et al. 2019). The Si is transported to the lake as an ion or in a colloid form, and it stays in a liquid form under acidic conditions. Consequently, its concentration is positively correlated with conductivity ($r = 0.52$) and negatively correlated with pH ($r = -0.70$). The negative correlation with pH and weak negative correlation with DOC ($r = -0.37$) suggest that in neutral and alkaline lakes, which are more productive (e.g. sites 31 or 68), at least part of the Si budget is consumed by diatoms (Sienkiewicz and Gąsiorowski 2019) and subsequently deposited into sediments. This process results in a higher concentration of Si in lakes with low pH and a simple food web, characterised by sediments rich in iron hydroxides (Fig. 6).

The groups of sites clearly separated on the RDA diagram were eutrophic and hypertrophic lakes (e.g. sites 2, 4, 68). These sites were shallow lakes with sediments enriched in P, Mg, C, and N. Furthermore, the P content in water of these lakes was relatively high,
exceeding 100 μg L⁻¹ in the hypertrophic lake group. Although these lakes are very shallow (maximum depth of 1.5–2 m), oxygen stratification might occur because of the intensive mineralisation of organic matter deposited at their bottom. The organic matter of these lakes, characterised with a low C/N ratio (~ 10), originates almost exclusively from in-lake primary production (Meyers and Teranes 2001). In contrast, some lakes (34, 55, 71) have a significantly higher C/N ratio (30–90), suggesting that organic matter in these water bodies mainly originates from their catchment areas. We can assume that lignite deposits in the bedrock of these lakes, or even pieces of lignite leached from the surrounding rocks, may be the cause of the increased C/N ratios in the sediments (Blodau et al. 2000).

Analysis of the relationship between water and sediment chemical composition indicated that even small changes in lake characteristics (e.g. changes in oxygen depletion in the hypolimnion or breaking the water stratification) can lead to substantial changes in chemistry. An increase in oxygen concentration can lead to oxidation of Fe²⁺ cations to less soluble Fe³⁺ forms or, particularly in eutrophic lakes, to the intensive oxidation of organic matter deposited at the bottom and further eutrophication. These changes might be driven by natural (e.g. weathering of the catchment and carbonate supply) or artificial (e.g. neutralisation with calcium carbonate or sediment dragging) factors. To plan the remediation of these lakes, it is important to identify the chemical characteristics of water and sediments in each lake individually. Because of the high geochemical and biological variability, a universal practice to revitalise these lakes would be ineffective. These lakes are unique ecosystems that need to be protected, at least partially. Hence, a recovery programme should be developed for specific lakes following detailed investigation and monitoring.

5 Conclusions

The water and sediment chemical composition of post-mining lakes in the Muskau Arch region indicates elevated concentration of some elements, for example, Al and Fe, compared to lakes in other areas of Central Europe. On the other hand, the concentration of these elements is approximately three orders of magnitude lower compared to other regions impacted by AMD, such as the Iberian Pyrite Belt.

This study has shown that, similar to water, surface sediments of investigated lakes can vary, even in lakes in close proximity. This is especially common in the middle part of the region. It is not surprising that water pH, varying along the gradient in the area, was one of the most important factors explaining the variability in sediment chemical composition. In addition, P and DOC concentrations indicated the trophic state of the lake and determined its productivity and impact on sediment chemistry. Surprisingly, Fe and Al concentrations or water conductivity (reflecting water total mineralisation) were not statistically significant factors to explain the sediment composition. Future studies should analyse Fe and Al speciation or vertical changes in lake water chemistry to identify their relationship with sediment composition.

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