Geochemical focusing and sequestration of manganese during eutrophication of Lake Stechlin (NE Germany)

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Abstract Significant sedimentation of manganese (Mn) in form of manganese oxides (MnOₓ) and the subsequent formation of authigenic calcium-rich rhodochrosite (Mn(Ca)CO₃) were observed in the seasonally stratified hard water Lake Stechlin in northeastern Germany. This manganese enrichment was assumed to be associated with recent eutrophication of the formerly oligotrophic lake. The mechanisms and processes involved were examined by analysing: (i) short sediment cores obtained from seven locations along a depth transect ranging from 69.5 m (the deepest point) to 38 m; (ii) sediment traps located at 20 m and 60 m water depths; (iii) water column profiles; and (iv) porewater profiles at 69.5 m and 58 m depths. Sedimentary Mn enrichment was observed at water depths below 56 m and increased to more than 25 wt% at the deepest site. Between 2010 and 2017, Mn accumulation at the deepest site was 815 g Mn m⁻². Transfer of Mn from the shallower towards the deepest parts of the lake was initiated by reductive dissolution of MnOₓ and diffusion of dissolved Mn from the sediment to the overlying water column. Manganese was then dissipated via turbulent mixing and subsequently oxidised to MnOₓ before being transported towards the deepest zone. Transformation of the redeposited MnOₓ to Mn(Ca)CO₃ favoured the final burial of Mn. We show that eutrophication and the areal spreading of anoxic conditions may intensify diagenetic processes and cause the spatial redistribution of Mn as well as its effective burial. Contrary to many previous findings, we show that increases of Mn and Mn/Fe can also be used as indicators for increasing anoxic conditions in previously oligotrophic lakes.

Keywords Lacustrine sediments · Geochemical focusing · Eutrophication · Diagenesis · Rhodochrosite · Varves
Introduction

Manganese (Mn) is a redox-sensitive element and an important terminal electron acceptor in the anaerobic mineralisation of organic matter in aquatic ecosystems. At the oxic–anoxic transition zone, i.e., at the sediment-water interface (SWI) or in the water column of stratified water bodies, it typically circulates between the dissolved phase (Mn$^{2+}$) in the underlying reducing environment and the solid phase in the overlying oxic environment (e.g. Calvert and Pedersen 1993; Davison 1993; Tribovillard et al. 2006). The predominant reduced form of this metal found in nature is Mn$^{2+}$, whereas Mn$^{3+}$ is found as an important intermediate form in various suboxic and oxic waters (Trouwborst et al. 2006; Dellwig et al. 2012; Madison et al. 2013; Oldham et al. 2017). In its particulate phase, Mn forms oxide (oxidation state (IV), carbonate (II), or sulphide (II) mineral phases (Suess 1979; Davison 1993; Dellwig et al. 2010; Häusler et al. 2018).

Depending on the morphological and hydrophysical characteristics of lakes and marine basins, redox changes may cause a spatial redistribution of particulate Mn in lakes (Delfino et al. 1969; Ostendorp and Frevert 1979; Schaller and Wehrli 1996; Dellwig et al. 2018) and marine basins (Rajendran et al. 1992; Lenz et al. 2015a, b; Häusler et al. 2018; Schaller and Wehrli 1996) introduced the term "geochemical focusing" to address the mobilisation of Mn from shallow sediments, its transportation within the water column, and sedimentation in the deep zones of lakes. In marine geology and sedimentology, the term “shelf-to-basin shuttle” is more common for this transport process (Lyons and Severmann 2006; Scholz et al. 2013).

While geochemical focusing describes the phenomenon of spatial Mn redistribution, it does not imply a long-term burial of Mn. Effective burial may occur if, for example, Mn is rapidly buried due to high Mn sedimentation rates in combination with low Mn oxide (MnO$_x$) reduction rates (Granina et al. 2004; Och et al. 2012). Alternatively, a mineral phase that is resistant to reductive dissolution within sediments may form (Hamilton-Taylor and Price 1983; Stauffer 1987; Giblin 2009). Such mineral may be calcium (Ca)-rich rhodochrosite (here referred to as Mn(Ca)CO$_3$) that has been reported in lacustrine (Mayer et al. 1982; Nuhfer et al. 1993; Friedl et al. 1997; Roeser et al. 2016; Dräger et al. 2017) and marine sediments (Suess 1979; Calvert and Pedersen 1996; Neumann et al. 2002; Johnson et al. 2016; Dellwig et al. 2018; Häusler et al. 2018).

The formation of Mn(Ca)CO$_3$ is fairly well-studied in anoxic basins of the Baltic Sea and has been associated with inflow events from the North Sea that oxidise formerly sulphidic waters highly enriched in Mn$^{2+}$, which ultimately results in significant deposition of MnO$_x$ on the seafloor (e.g. Huckriede and Meischner 1996). When reducing conditions are re-established and cause substantial MnO$_x$ dissolution, Mn(Ca)CO$_3$ formation may occur close to the surface of sediments where Mn$^{2+}$ concentrations reach supersaturation (Middelburg et al. 1987; Calvert and Pedersen 1993) although involvement of bacteria has also been suggested (Aller and Rude 1988; Burdige 1993; Yao and Millero 1993; Neumann et al. 2002; Lee et al. 2011; Henkel et al. 2019). The long-term oxygenation of bottom waters in contact with anoxic sediment was found to facilitate the formation of Mn(Ca)CO$_3$ in the Baltic Sea (Dellwig et al. 2018; Häusler et al. 2018). Alternative models of Mn(Ca)CO$_3$ formation were presented by Herndon et al. (2018) and Wittkop et al. (2020), where the formation of Mn-rich calcite and rhodochrosite was suggested to occur also within redox-stratified water bodies of Fayetteville Green Lake and Brownie Lake supported by calcite presence.

The concept of geochemical focusing is still not well understood and suffers from a substantial lack of in-situ documentation. Many studies of the phenomenon focussed on the marine environment (e.g. Lyons and Severmann 2006; Scholz et al. 2013; Lenz et al. 2015a, b; Häusler et al. 2018), while lacustrine studies are notably rare (Schaller and Wehrli 1996; Naehler et al. 2013). The diagenesis of Mn and the resulting role of Mn as a proxy of changing redox conditions is similarly poorly understood.

Exceptionally high Mn contents were found in the surface sediments of the seasonally stratified Lake Stechlin in Germany (Kleeberg 2014). However, this could not be simply explained by marine models developed for temporarily stratified basins because the waters of Lake Stechlin are mostly oxic-suboxic due to the annual mixing of the waterbody. Furthermore, the suggestion by Mackereth (1966) and Wersin et al. (1991) that Mn should be increasingly depleted from
sediments upon exposure to more reducing conditions could not explain the Mn enrichment of Lake Stechlin.

To address these knowledge gaps, we aimed to explain the recent accumulation of Mn in the deepest areas of Lake Stechlin and describe the pathways of geochemical focusing. We hypothesised that recent eutrophication and lowered hypolimnetic O$_2$ concentrations have influenced the lake’s internal Mn cycle. Based on the quantitative analysis of Mn pools and fluxes using sediment core sampling from various water depths, highly resolved measurements at the SWI, sediment trap sampling at two different water depths, and monitoring of Mn$_{diss}$ in the water column, we develop a new conceptual model to describe the observed phenomenon.

**Study site**

Lake Stechlin is located in north-eastern Germany (53°09'N, 13°02'E, Fig. 1) in a sandy glacial landscape with a catchment area covered in 80% by forest. The dimictic lake is characterised by an area of 4.23 km$^2$, a mean depth of 23.3 m, a maximum depth of 69.5 m (Koschel and Adams 2003), and is thermally stratified from about April until the end of December. Except for two narrow and shallow channels that connect to Lake Nehmitz in the southwest, Lake Stechlin lacks substantial surface in- or outflows. The water level is nearly constant since 1957 as the lake outlet is regulated (Pöschke et al. 2018). The major sources of water are precipitation and groundwater (Richter and Koschel 1985). During the operation of a nuclear power plant between 1966 and 1990, 300,000 m$^3$ day$^{-1}$ water from the neighbouring Lake Nehmitz was used for cooling and was discharged into Lake Stechlin heated by 10 °C. The water residence time during that time was 335 days and since 1990 it increased to > 40 years (Koschel and Adams 2003). Thermal pollution from the nuclear power plant, and more recently climate change, have increased the duration of the summer stratification period in the lake (Kirillin et al. 2013). Long-term monitoring data of the Leibniz Institute of Freshwater Ecology and Inland Fisheries (IGB) showed that the lake annual average total phosphorus concentration has increased from 14 ± 3 (2000–2010) to 33 ± 3 μg L$^{-1}$ (2015–2017). The annual mean chlorophyll-$a$ concentration in the surface waters for the period from April to October also consistently increased from 2.6 ± 0.6 (2000–2010) to 3.3 ± 0.4 μg L$^{-1}$ (2015–2017). Minimal O$_2$ concentrations were observed at a water depth below 60 m, which decreased from 4.8 ± 0.7 (2000–2010) to 0.7 ± 0.4 mg L$^{-1}$ (2015–2017) with major drops in 2003 and 2011. The higher O$_2$ consumption rate has resulted in a steadily increasing area of the lake bottom waters with less than 2 mg O$_2$ L$^{-1}$ at the end of the stratification period. Consequently, Lake Stechlin has lost its oligotrophic status and seems on a trajectory towards more eutrophic conditions. These recent changes in water quality have been described in detail by Selmeczy et al. (2019).

**Material and methods**

**Sampling and preparation of sediment cores**

Eight sediment cores (60 cm length, 6 cm diameter) were obtained with a Kajak gravity corer (UWITEC, Mondsee, Austria) during two field campaigns: one core was obtained from the deepest part of the lake at a water depth of 69.5 m on 24 August 2017 and the other six cores (at depths of 65 m, 64 m, 61 m, 56 m, 51 m, and 38 m) were obtained along a depth-transect towards the south-eastern slope on 3 May 2018 (Fig. 1). After sampling, the taped cores were placed horizontally and split with a stainless-steel plate into two halves. The sediment was stabilised by a floral foam from above so that the sediment remained undisturbed when placed horizontally. One half was used for imaging and X-ray fluorescence (XRF) core scanning, and the second half was sub-sampled into 1 cm sections. The sediments were freeze-dried, ground with an agate mortar, and used for further geochemical analyses. On 15 November 2017, 12 additional cores were taken at depths of 69.5 m and 58 m, respectively (Fig. 1). The surface sediments (0–1 cm and 1–2 cm) at each depth were pooled and used for the analysis of the Mn fraction by sequential extraction. The total mass accumulation of Mn in the sediment cores was calculated based on the total Mn content of their Mn-rich upper layers (defined as samples with > 1.2 wt% Mn) multiplied by dry mass per area of the respective layers. The cores from the deepest part of the lake, which were used for Scanning Electron Microscopy-Energy-Dispersive X-ray
Spectroscopy (SEM-EDX) analysis, were obtained on 19 November 2018 and were freeze-dried but not ground.

Sampling of settling seston

Sediment traps were positioned at depths of 20 m and 60 m in the water column above the deepest point of the lake to determine Mn sedimentation rates (Fig. 1). The sediment traps comprised four cylinders 9 cm in diameter and 100 cm in height. The traps were treated with 0.13% formaldehyde to avoid microbial degradation (Lee et al. 1992) and sampled at 1–2.5-month since autumn 2009; a significant break in the measurements occurred between 2013 and the middle of 2016 as well as the following shorter periods: 24 September to 22 October 2009, 31 August to 13 October 2010, 19 December to 24 March 2012, 13 June to 4 July 2016, and 19 December to 31 March 2016. The sedimentation rate of Mn (mg m\(^{-2}\) year\(^{-1}\)) was calculated by multiplying Mn content by the dry mass sedimentation rate. To calculate the total sedimentation of Mn in the traps during the period 2010–2017, the shorter gaps in the sampling record...
were filled using linearly interpolated values, but the longest gap in the record (between 2013 and mid-2016) was not interpolated. To illustrate the formation of authigentic Mn minerals (e.g., MnO\textsubscript{x} formed at the redoxcline), the Mn contents in the sediment traps were related to aluminium (Al) as representative of the lithogenic background sedimentation.

Water and porewater sampling

Water samples were collected every month from August 2017 until January 2019, except February and March 2018, at several water depths depending on the mixing regime: a surface water sample and every 5 m below 30 m during the stagnation period, and a surface water sample, 40 m, 60 m, 65 m and 68 m during the mixing period. All water samples were taken at the deepest site of the lake using a water sampler (Limnos, Turku, Finland). Dissolved manganese (Mn\textsubscript{diss}) concentrations were determined by ICP-OES (iCAP 6300, Thermo Fischer Scientific, Waltham, USA) after membrane filtration of water samples using a 0.45-μm cellulose acetate syringe filter (Whatman GmbH, Dassel, Germany) and fixing with 10 μL of 2 M HCl per 1 mL of sample. The mass of Mn\textsubscript{diss} in the water column at the deepest point (g m\textsuperscript{-2}) was calculated by the depth-weighted mean value multiplied by a water depth of 69 m (see also supplementary dataset). Oxygen concentrations were determined by O\textsubscript{2} logger (YSI 6600 V2-4 Xylem Analytics; Rye Brook, USA).

Porewater profiles were obtained using dialysis chambers (30 × 3.2 × 1.5 cm) consisting of acrylic glass covered with a HT-Tuffryn 200 membrane (GELMAN) following the method of Hesslein (1976). A frame was used to keep the dialysis chamber in the vertical position within the sediments. The dialysis chambers were filled with deionised water, deoxygenated using N\textsubscript{2}, and placed in the sediments at water depths of 69.5 m and 58 m (Fig. 1). After 14 days of exposure, the dialysis chambers were retrieved on 29 November 2017 and 26 June 2018, and the collected porewater was acidified with 10 μL of 2 M HCl per 1 mL of sample. Porewater profiles for sulphate and sulphide were obtained using an analogous dialysis chamber at the deepest site (69.5 m) on 19 November 2018. Dissolved sulphide was fixed with a 0.2 mL ZnAc solution (about 50 mM L\textsuperscript{-1} Zn) per 2 mL sample.

Geochemical analyses

XRF core scanning

The sediment cores were scanned for Mn, Ca, titanium (Ti), and silicon (Si) using an ITRAX core scanner (Cox, Mölnadal, Sweden; Croudace et al. 2006; Jarvis et al. 2015) at the Leibniz Institute of Baltic Sea Research Warnemünde (IOW). Core halves were scanned using a Cr-tube set to 30 kV, 30 mA, and with an exposure time of 15 s per step. The measurement resolution in the core from 69.5 to 0.05 mm for the uppermost 7 cm (varved/laminated section) and 0.2 mm for the lower homogenous section; the other sections of the cores were measured using a 0.1 mm resolution. The range of the resulting XRF signal was 3500–170,000 cps for Mn; 2000–80,000 cps for Ca; 130–1800 cps for Ti; and 75–1200 cps for Si. X-ray images were also obtained with the ITRAX core scanner at a 0.2 mm downcore resolution.

Dissolved and solid elemental and compound analysis

The Mn, Ca, Al, and Fe content of the 69.5 m depth core was determined using inductively coupled plasma-optical emission spectrometry (ICP-OES; iCAP 7400 Duo, Thermo Fisher Scientific, Waltham, USA) after acid digestion (HNO\textsubscript{3}, HClO\textsubscript{4}, and HF) following the procedure after Dellwig et al. (2019). Precision and trueness were checked by using the international reference material SGR-1b (USGS) and were better than 6% and −5%, respectively. The relative content of Mn and Ca, as determined by the XRF, was linearly calibrated with the absolute content determined by ICP-OES as successfully applied by Moller et al. (2012). Prior calibration using an offset-correction of depth based on five prominent peaks in Ca and Mn was performed between the halves of the cores used for the XRF and ICP-OES analyses. Then the average XRF values of less resolved ICP-OES intervals were plotted against the ICP-OES measurements, giving linear regressions coefficients of R\textsuperscript{2} = 0.87 for Mn and R\textsuperscript{2} = 0.96 for Ca. The sediment trap material was digested by a mixture of 36% HCl and 65% HNO\textsubscript{3} at a volumetric ratio of 1:3 (aqua regia) in a high-pressure microwave (MLS GmbH, Leutkirch, Germany). The Mn and Al contents in the digested trap material, in the core from 38 m water depth, and dissolved Mn and Ca in the water samples

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(lake and porewater) were measured using ICP-OES (iCAP 6300 Duo, Thermo Fisher Scientific). The corresponding precision and accuracy were < 9% and < 5%, respectively. The pH of the porewaters was determined using a SenTix® 940 electrode (WTW 3430, Weilheim, Germany). Dissolved sulphide concentrations were measured using the method after Cline (1969) and the precision of this method is usually around 3% (Dellwig et al. 2019). Sulphate concentrations were determined via ion chromatography (Shimadzu, Kyoto, Japan) and the relative standard error was < 1%. The contents of total carbon (TC) and total organic carbon (TOC), the latter after treatment with 1 M HCl in order to remove carbonates, were determined using a CNS element analyser (Vario EL, Elementar Analysensysteme GmbH, Langenselbold, Germany). Total inorganic carbon (TIC) was calculated as the difference between TC and TOC. Each sample of the CNS analysis was performed as duplicate and the relative standard error was < 1% for TC and < 4% for TOC. Loss on ignition (LOI) for the flux calculations was determined as the loss of weight after the combustion of dried sediment at 450 °C for 3 h, and the relative standard error (n = 2) was < 1%.

Manganese fractionation

To distinguish reducible MnOx from acid-soluble Mn carbonates from the sediment trap material from 60 m (exposure time: 19 September–29 October 2018, and 29 October–18 December 2018) and surface sediments (0–1 and 1–2 cm, 69.5 m and 58 m water depth), we deployed a sequential wet-chemical extraction protocol that follows the procedure of Psenner et al. (1984) originally developed for phosphorus fractions. The extraction scheme comprised four steps: (1) 1 M ammonium chloride (NH4Cl), representing loosely adsorbed and porewater Mn, (2) 0.11 M bicarbonate-dithionite (BD) solution, for the reductive-soluble fraction representing MnOx mineral phases, (3) 1 M sodium hydroxide (NaOH), and (4) 0.5 M hydrochloric acid (HCl) targeting (acid-soluble) carbonate-bound Mn. For testing the plausibility, we compared the total Mn content determined by ICP-OES with the sum of the individual fractions. Each sample was performed in duplicate, the results were presented as averages, and the relative standard error was < 5%. The Mn concentrations in extracts 1, 2, and 4 were routinely measured by atomic absorption spectroscopy (AAS; Perkin Elmer 3300, Rodgau-Juegesheim, Germany). Since Mn was not measured in extract 3, NaOH soluble Mn is a part of residual Mn that was calculated as the difference between the total Mn and the sum of Mn in the NH4Cl, BD, and HCl fractions.

SEM-EDX

Scanning electron microscopy with energy dispersive X-ray fluorescence (SEM-EDX, FESEM Zeiss Merlin VP Compact) served for identification of authigenic mineral phases in the sediment trap material and sediment samples. The solid material was suspended in an ultrasonic bath and filtrated with 0.4-μm polycarbonate filters. After drying at 60 °C for 24 h, the filters were fixed on Al stubs and carbon-coated. The working distance in the SEM-EDX was 8.5 mm and spot analysis was done at an accelerating voltage of 15 kV using AZtec Energy software (Oxford Instruments). The identification and determination of relative abundances of the mineral phases were done by automated particle analyses. This procedure includes recognition of particles by grey level thresholds with a minimum ECD of 0.67 μm at a magnification of 2000. Identified particles (~ 1500 particles per sample) were measured for their average elemental composition by a secondary electron detector comprising several spot analyses per particle followed by image processing and particle classification using the AZtec Feature software (Oxford Instruments). Peak overlapping (e.g. Mn kβ and Fe kα lines at 6.4–6.5 keV) was solved by peak deconvolution. Depending on the chosen instrument parameters, detection limits of the EDX measurements of our environmental samples typically range from 0.1 to 1 wt% and values < 1 wt% should be considered as informative only. Based on the chemical composition of the minerals and previous experiences with suspended particulate matter and sediment samples from the Baltic Sea (Dellwig et al. 2010, 2018; Häusler et al. 2018), the threshold values for the identification of the mineral phases were as follows: Mn > 30%, Ca < 6%, Fe < 5%, and P < 5% for Mn oxide; Mn > 30%, Ca > 6%, and Fe < 5% for Ca-rich Mn carbonate; and Mn > 10%, Fe > 5%, Ca < 6%, and P > 1% for Mn-Fe-P phases, which. In addition to Ca, Fe, Mn, and P, the contents of Mg and Si were also determined by the EDX analysis.
Fluxes of dissolved manganese

Diffusive fluxes of $M_n_{\text{diss}}$ were calculated from the respective concentration gradients across the SWI according to Fick’s first law of diffusion (following the formulas presented by Lewandowski et al. 2002):

$$ J_{M_n} = \frac{\varphi}{\theta^2} \times D_{M_n} \times \frac{\Delta C_{M_n}}{\Delta z} $$

with $J$ is the diffusion rate of Mn (g m$^{-2}$ day$^{-1}$), $\varphi$ is the porosity (m$^3$ porewater/m$^3$ wet sediment), $\theta$ is the tortuosity (unitless), $D_{M_n}$ is the molar diffusion coefficient of Mn in the sediment ($6.88 \times 10^{-6}$ cm$^2$ s$^{-1}$; Li and Gregory 1974), $\Delta C/\Delta z$ is the concentration gradient at the SWI (g m$^{-4}$). Porosity was calculated as:

$$ \phi = \frac{w}{w_{\text{water}}} + \frac{1-w}{\rho_{\text{sediment}}} $$

where $w$ is the water content (wt%/100), $\rho_{\text{water}}$ density of water at 4 °C. The density of sediment ($\rho_{\text{sediment}}$), was calculated as:

$$ \rho_{\text{sediment}} = \frac{1}{\rho_{\text{org}} + \frac{1-loi}{\rho_{\text{min}}}} $$

where LOI is the loss on ignition (wt%/100), $\rho_{\text{org}}$ density of organic matter (1.4 g cm$^{-3}$) and $\rho_{\text{min}}$ density mineral matter (2.65 g cm$^{-3}$). Tortuosity was calculated following the formula by Boudreau (1996):

$$ \theta = \sqrt{1 - \ln \phi^2} $$

To calculate the diffusion coefficient at 4 °C, the following formula as presented by Lewandowski et al. (2002) was used:

$$ D_{M_n} = D_{M_n,25\,^\circC} \times \frac{\mu_{25\,^\circC} \times T}{\mu_T \times 298.15\,^\circK} $$

where $\mu$ is the dynamic viscosity of water at 25 °C (0.890 g m$^{-1}$ s$^{-1}$) and $T$ is the temperature at the SWI considered as 4 °C. The diffusion coefficient of $D_{M_n}$ at 4 °C was calculated as $3.12 \times 10^{-5}$ m$^2$ day$^{-1}$. Porosity and tortuosity were determined using the 69.5 m and 56 m depth sediment cores for the porewater fluxes at 69.5 m and 58 m, respectively. The hypolimnetic Mn accumulation was calculated based on the $M_n_{\text{diss}}$ concentration in different water layers multiplied by the volume of the respective depth interval.

**Results**

**Sediment investigations**

**Sediment properties and varve formation**

The upper < 5 cm of the three deepest cores (between 69.5 and 65 m) contained macroscopical varves, represented by variations of Si, Ca and Mn (Fig. 2). In the sediment core obtained at 69.5 m in August 2017, eight varves were counted within the uppermost 4 cm, so the beginning of continuous varve formation coincides with the major acceleration of eutrophication in 2010. By comparing the profiles of Ca, Mn and Fe of a core that was sampled at the same site in 2010 (Kleeberg 2014), we can infer that the newly deposited layers amount to 4–5 cm. In contrast, varves were disturbed or absent at the shallower sites. From 12 cm upwards in the 69.5 m core, the Al and TOC contents decreased from 2 to 0.15 wt% and from 25 to 10 wt%, respectively (Fig. 3). A distinct increase in TIC was observed with a peak of 4.5 wt% at 9 cm and 6 wt% in the upper part of the varved sections. The Ca content increased almost parallel to TIC and reached a maximum of 16 wt% at approximately 9 cm. The top part of all of the cores along the transect contained a dark grey, Ca-rich section above 15 cm depth (Figs. 2 and 4).

**Manganese and calcium in sediment cores along a water depth transect**

A distinct surface Mn enrichment was only present in deepest cores below 56 m, and the thickness of the Mn-rich layer increased from 3 cm in the 56 m core to 8 cm in the 69.5 m core (Fig. 4; see Mn vs. Ti in Figs. S3 and S4 in supplementary material). Furthermore, the Mn content at the SWI increased towards the deepest point, reaching 18 wt% in 1 cm intervals (Fig. 3) and up to approximately 45 wt% in single laminations in the uppermost parts of the cores (Fig. 4). In the 0–5 cm interval of the core from 38 m water depth, the ICP-OES derived Mn content was 0.04–0.06 wt% (see supplementary dataset).
contrast to other cores, additional three Mn-rich layers (Figs. 3 and 4) were present in the deepest core (69.5 m), at sediment depths of 38 cm, 31 cm, and 25 cm, and less distinct in the second-deepest core (65 m). The Mn content in these layers was distinctly lower (max. 6.0 wt%) than the Mn enrichment at the core top. The lowermost Mn-rich layer was located in an Al-rich layer, accompanied by increased TIC and Ca contents (Fig. 3). The Mn-rich layers in the middle of the core (at 31 cm and 25 cm) were also accompanied by a weakly increased TIC content, but not by increased Ca or Al contents. Whilst the Fe content decreased towards the sediment surface from 7 to 0.7 wt%, the Mn/Fe ratio increased in the Mn-rich upper portions (Fig. 3). The total Mn accumulation in the Mn-rich layer (0–8 cm) of the 69.5 m core (sampled on 24 August 2017) was 940 g m$^{-2}$. In comparison, the Mn-rich layer at the top of a core sampled at the same site on 30 October 2010 (Kleeberg 2014) was 4-cm thick and had a Mn content equivalent to approximately 3 wt%. The calculated Mn accumulation based on this core was 125 g m$^{-2}$, and the
difference between 2017 and 2010 was 815 g m\(^{-2}\), giving a mean accumulation rate of 116 g m\(^{-2}\) year\(^{-1}\).

Dissolved manganese in the water column

Concentrations of Mn\(_{\text{diss}}\) in the water column showed a clear dependence on the seasonal patterns of stratification and O\(_2\) concentrations (Fig. 5). During the summer stratification period on 8 August 2017, Mn\(_{\text{diss}}\) concentrations increased towards the O\(_2\)-poor bottom waters to a maximum of 0.3 mg L\(^{-1}\) at still oxic conditions of > 5 mg L\(^{-1}\). Manganese concentrations increased up to 0.6 mg L\(^{-1}\) in the bottom waters in September 2017, but unexpectedly decreased to \(~\) 0.005 mg L\(^{-1}\) during most O\(_2\)-depleted bottom waters at the end of the summer stratification period (< 2 mg L\(^{-1}\) in the bottom water column on 12 December 2017; Fig. 5). The Mn\(_{\text{diss}}\) concentration remained low (< 0.005 mg L\(^{-1}\)) during the following winter mixing period (9 January 2018), until the beginning of the next stratification period in spring 2018 when bottom water Mn\(_{\text{diss}}\) rose again to \(~\) 0.2 mg L\(^{-1}\) (26 April 2018). During the stratification period in 2018, Mn\(_{\text{diss}}\) concentrations increased up to 5 mg L\(^{-1}\) in the few lowermost metres, while the bottom water became oxygen-free.

Porewater concentrations and diffusive fluxes

The Mn\(_{\text{diss}}\) concentrations in the sediment porewaters were up to several times higher than in the water column (Fig. 6). In the core obtained at 58 m water depth in June, the Mn\(_{\text{diss}}\) profile was almost constant, at approximately 6 mg L\(^{-1}\), with only a slight decrease to 5 mg L\(^{-1}\) at 3 cm depth and a peak at the SWI that reached 7.6 mg L\(^{-1}\) in June and 11 mg L\(^{-1}\) in November. At 69.5 m, Mn\(_{\text{diss}}\) concentrations were very similar during November 2017 and June 2018, and were higher than at 58 m; Mn\(_{\text{diss}}\) concentrations peaked with 25 mg L\(^{-1}\) just below the SWI. Notable Mn\(_{\text{diss}}\) concentrations at 69.5 m were observed in the overlying water column, reaching more than 1.8 mg L\(^{-1}\) in June and 0.2 mg L\(^{-1}\) in November.

At 69.5 m, the diffusive flux of Mn out of the sediment to the water column was 16.8 g m\(^{-2}\) year\(^{-1}\) in November and 7.5 g m\(^{-2}\) year\(^{-1}\) in June. At 58 m, the diffusive flux was 3.2 g m\(^{-2}\) year\(^{-1}\) in November 2017 and 8.2 g m\(^{-2}\) year\(^{-1}\) in June 2018. Mean annual fluxes were calculated as the average of the June and November values, i.e. 12.1 g m\(^{-2}\) year\(^{-1}\) at 69.5 m and 5.7 g m\(^{-2}\) year\(^{-1}\) at 58 m.

The dissolved Ca concentrations (Ca\(_{\text{diss}}\)) in the porewater and bottom water column in November and June amounted to approximately 40–50 mg L\(^{-1}\) at 58 m and 69.5 m, with the highest concentrations occurring in June (Fig. 6). In the top 10 cm, Ca\(_{\text{diss}}\) concentrations dropped by approximately 5 mg L\(^{-1}\) relative to the deeper layers and in the overlying water both in June and November at 69.5 m and in June at 58 m. Sulphate concentrations decreased abruptly from 26 mg L\(^{-1}\) in the water column to approximately

![Fig. 4](https://www.springer.com)
3 mg L\(^{-1}\) in the porewater at both depths in June and November (Fig. 6). The pH was very similar at both depths and was almost constant in June at approximately 7.6 in the water column and 7.4 in the sediment. In November, pH increased to 7.8, reaching a peak just below the SWI relative to the background level in the sediment (7.4) and the water column (7.2). Total dissolved sulphide was absent below 25 cm, increased towards the SWI to 4.4 mg L\(^{-1}\), and dropped to below 0.1 mg L\(^{-1}\) 10 cm above the SWI (Fig. 7).

Manganese in sediment traps

Manganese sedimentation rates showed seasonal variations and were much higher at 60 m in the water column than at 20 m (Fig. 8). At the 20 m water depth, the Mn content in the trap material varied between < 0.1 and 1.7 wt% (Fig. 9) with an average of 0.3 wt% between 2010 and 2018. The Mn sedimentation rate at 20 m varied between < 0.1 and 4.8 mg m\(^{-2}\) day\(^{-1}\) but did not show any discernible seasonal pattern (Fig. 8). In the trap at 60 m water depth, the Mn content varied between 0.2 and 26 wt%, with an average of 8.7 wt% (Fig. 9). The Mn sedimentation rates at 60 m varied between 3 mg m\(^{-2}\) day\(^{-1}\) and 404 mg m\(^{-2}\) day\(^{-1}\) with the minima occurring during the mixing period in winter and the maxima occurring during the late stagnation period. Between 2010 and 2017, the total Mn sedimentation recorded by the traps was 4.3 g m\(^{-2}\) at 20 m and 297 g m\(^{-2}\) at 60 m (supplementary dataset). In 2017, the Mn sedimentation rate was 0.7 g m\(^{-2}\) year\(^{-1}\) at 20 m and 50.9 g m\(^{-2}\) year\(^{-1}\) at 60 m. The annual maximum Mn
sedimentation rate at 60 m gradually increased between 2010 and 2012 and has remained constant until at least 2016 (Fig. 8). Minimum O\(_2\) concentrations decreased every year, reaching approximately 0.3 mg L\(^{-1}\) by the end of the stagnation period. The dry mass sedimentation rate ranged between 0.05 and 1.3 g m\(^{-2}\) day\(^{-1}\) (average 0.49 g m\(^{-2}\) day\(^{-1}\)) in the 20 m trap and between 0.07 and 8.1 g m\(^{-2}\) day\(^{-1}\) (average 1.1 g m\(^{-2}\) day\(^{-1}\)) in the 60 m trap (see Fig. S1 in supplementary material). The highest dry mass sedimentation occurred during the summer period.

The Mn content in the sediment traps deployed during the mixing period at both 20 m and 60 m depths in the water column showed a relatively close relation to external sedimentary input represented by Al (Fig. 9). While during the stagnation period, the Mn vs. Al linear relationship remained fairly unchanged at 20 m, the samples at 60 m water depth were enriched in Mn relative to Al having up to five times higher Mn contents compared to the trap material obtained during the mixing period.

Manganese fractionation

The Mn fractions differed substantially between the sediment trap material and the surface sediment sampled in the early summer during a stratification period (26 June 2018) and before the end of the stratification period in late autumn (29 November 2017).
Sediment trap samples (60 m) from the end of the stagnation period had an overall high Mn content of 20 wt% and 26 wt% in trap a and b, respectively. The reductive soluble fraction (BD-Mn), representing MnOx mineral phases, was most abundant and amounted to ca. 95% of the total Mn content. The other fractions represented only a minor portion of the total Mn: 0.1% was HCl-Mn, ca. 3% was NH4Cl-Mn, and ca. 2% was residual Mn. Although dissolution of MnOx particles by HCl is conceivable, the BD-Mn fraction is assumed to have extracted all MnOx in the previous extraction step.

While being almost completely absent in the sediment trap material, the HCl-Mn fraction, representing Mn carbonates, was most abundant in the surface sediment and amounted to approximately 3.1 wt% at 58 m and 5–7 wt% at 69.5 m (equivalent to 54–61% and 32–52% of total Mn, respectively). The BD-Mn fraction amounted to <1 wt% at 58 m and 1–3 wt% at 69.5 m and, in contrast to the sediment trap material, constituted a minor part of the total Mn (12–17% and 10–13%, respectively). It is notable that the NH4Cl-Mn fraction yielded higher values at 69.5 m (2.7–8 wt%) than in 58 m (approximately 1 wt%), and that the content of the BD-Mn fraction decreased slightly while the HCl-Mn fraction increased from 0 to 1 cm towards 1–2 cm depths (Fig. 10).

SEM-EDX analysis

Of the sediment trap particles, 67% were identified as Mn oxides and 2.7% as Mn carbonates (supplementary dataset). Further 8.2% of the particles were referred to as Mn-Fe-P-rich particles, as described in more detail by Dellwig et al. (2010). The Mn oxides from the trap material had variable sizes ranging from a few to hundreds of micrometres (Fig. 11), were poorly crystalline and branching vermicular in shape, resembling Mn oxides described in previous studies (Dellwig et al. 2012, 2018; Schnetger and Dellwig 2012). The Mn oxides from the 60 m trap typically contained only a small amount of Ca (2.4 wt%), resulting in a high molar Mn/Ca ratio of approximately 14. As further accessory elements Fe (3.1 wt%), P (1.2 wt%), Si (2.8 wt%) as well as trace amounts of S (0.4 wt%) and Mg (0.1 wt%) were determined, with the last two element contents within the range of the EDX detection limit.

In the 0–1 cm sediment horizon at a depth of 69.5 m, 8% of the particles were identified as Mn oxides, 67% as Mn carbonates, and 1% as Mn-Fe-P-rich particles. In the 1–2 cm horizon of the same core, the proportion of Mn oxides dropped to 1.3%, whereas 42% were Mn carbonates and 1.2% were classified as Mn-Fe-P-rich particles. The Mn carbonate particles had a size of approximately 2 μm, were spherical smooth-edge rhombohedral or “almond-like” in shape, had an average stoichiometry of (Mn0.79-Ca0.21)CO3, and are grouped into larger aggregates (Fig. 11; supplementary dataset). Unlike in Mn carbonates from the Big Watab Lake in the USA (Stevens et al. 2000) or in the Landsort Deep (Häusler et al. 2018), the deepest basin in the Baltic Proper, no Mg could be determined in the Mn carbonate phase, but significant amounts of Si (2.9 wt%), Fe (0.5 wt%), P (1.2 wt%), and S (1 wt%) were present.
Discussion

Manganese enrichment in sediments of Lake Stechlin

Sediments of Lake Stechlin in this study were found to be enriched in Mn only below 56 m water depth (Fig. 4), accounting for a relatively small area of approximately 3.9% of the total lake. During a field campaign in 1991, Mn content was determined in 51 surface sediments from various depths between > 1 and 69.5 m in the Lake Stechlin (Casper et al. 1995). It was observed that the deepest site contained the most Mn, although the contents of Mn found in the surface sediments were much lower (a maximum of 1.3 wt%) than determined in this study.

This Mn-rich upper layer postdates the onset of calcite (CaCO₃) precipitation (Figs. 3 and 4) that was dated to approximately 1900–1930 (Casper 1994; Gonsiorczyk et al. 1995). The calcite precipitation in Lake Stechlin was explained by increased CO₂-assimilation by primary production and increased water temperature (Koschel et al. 1985). The appearance of the Mn-rich layer slightly pre-dates the appearance of varves, which occurs in the uppermost section of the deepest core and can be counted back from 2017 to 2010 (Fig. 2). As such, the Mn-rich deposits coincide with the major drop in O₂ in the hypolimnion in 2010. Such a trend of increasing Mn accumulation alongside decreasing O₂ concentrations was also documented in the sediment trap record since 2010 (Fig. 8). Varve formation resulted from the seasonal and sequential enhanced sedimentation of organic matter, calcite, opal, and MnOₓ as a direct or indirect result of increased primary productivity and a lack of bioturbating organisms.

Manganese mineral phases

Manganese minerals in the water column collected by the sediment traps at 60 m were MnOₓ particles, according to SEM-EDX (Fig. 11). These particles were the dominant phase of the settling seston, as shown by particle counting under the SEM-EDX and a very high (95%) proportion of (reductive-soluble) BD-Mn fraction in the total Mn (Fig. 10). The relationship between Mn and Al for the particles mostly deposited during the oxygenated mixing phase (Fig. 9) could be considered as a background Mn sedimentation during...
periods where the redoxcline is near or within the sediment. High Mn contents and Mn/Al ratios in the 60 m traps (Fig. 9) indicate that MnO\textsubscript{x} particles precipitated from the water column during stratification.

In contrast to the trap material, SEM-EDX indicated that the dominant phase in the sediment was Ca-rich rhodochrosite (Mn(Ca)CO\textsubscript{3}; Fig. 11) when compared with studies on the structural and chemical composition of Ca-rich rhodochrosite in the hypoxic/euxinic basins of the Baltic Sea (Suess 1979; Dellwig et al. 2018; Häusler et al. 2018). The presence of this mineral phase was also suggested by the high amounts of HCl-soluble Mn (Fig. 10) and TIC (Fig. 3) that indicated an existence of both Mn and Ca carbonates. Similar enrichments in Mn(Ca)CO\textsubscript{3} must have occurred in the past, as evidenced by the Mn layers in lower sections of the deepest core (Figs. 3 and 4). The Ca-rich mixed phases are not expected in the two Mn-rich middle sections (31 cm and 25 cm) of the deepest core (69.5 m) as the Ca signal remained low in these layers (Fig. 3). The increased Mn, Ca, and TIC contents within the detritus-rich interval (at 38 cm depth) suggested the presence of Mn(Ca)CO\textsubscript{3}.

**Fig. 10** Left: contents of the four Mn fractions from the sequential extraction in the sediment trap material in 60 m (above) obtained on **a** 19 September–29 October 2018 and **b** 29 October–18 December 2018 and of the sediments at 58 m and 69.5 m water depth (below). Right: contents of the Mn fractions normalised to the total Mn content.

**Fig. 11** Exemplary SEM-photograph showing a MnO\textsubscript{x} particle (ID 6997) from the 60 m sediment trap (left) and Mn(Ca)CO\textsubscript{3} particles (ID 1466) in the surface sediment (right) at the deepest site of Lake Stechlin. Crosses mark the spots of the EDX elemental analyses. Additional SEM-EDX data are provided in the supplementary dataset.

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Previously, Röper and Schwarz (2003) detected Ca-rich rhodochrosite, using XRD, in a core collected from the deepest part of Lake Stechlin in 1993, in a layer below 25 cm, that should correspond with the Mn-rich interval observed in our study at 38 cm depth.

Formation of Mn(Ca)CO$_3$

The distinct difference between the dominance of MnO$_x$ in the water column and Mn(Ca)CO$_3$ at the sediment surface indicated that Mn(Ca)CO$_3$ formation has taken place at the SWI. We suppose that a mineral phase transition from Mn oxide to Mn carbonate required reductive dissolution of deposited MnO$_x$ and re-precipitation as Mn(Ca)CO$_3$. An active diagenetic transformation of MnO$_x$ to Mn(Ca)CO$_3$ was confirmed by the downward trend of BD-Mn loss in favour of Mn-HCl (Fig. 10). The confined stratigraphic appearance of Mn peaks (Fig. 2) further points to a quick conversion of MnO$_x$ into Mn(Ca)CO$_3$ and successful burial close to the SWI rather than dispersed post-depositional re-precipitation as also suggested for the Landsort Deep by Häusler et al. (2018). Sedimentation of MnO$_x$ was not abrupt but continuous and occurred mainly along the stagnation period from about April until December, as evidenced by the sediment traps (Fig. 8). Precipitation of calcite in Lake Stechlin occurs in June–July, and the particulate Mn nearly co-appears with calcite, as shown by the parallel occurrence of Ca and Mn peaks in the varved section (Fig. 2). An abrupt sedimentation or formation of particulate Mn, e.g. due to mixing with oxygenated surface waters, would have rather resulted in single thick Mn-laminations. Several factors may favour the authigenesis of Mn(Ca)CO$_3$, as for instance high porewater Mn$_{diss}$ concentrations (Fig. 6), which result from the reductive dissolution of abundant MnO$_x$ at the SWI (Figs. 4 and 10). Formation and burial of Mn as Mn(Ca)CO$_3$ did not occur and apparently was not possible in sediments at shallower water depths. We attribute this to insufficient amounts of reducible MnO$_x$, a higher water column O$_2$ levels inhibiting MnO$_x$ reduction, and an insufficient input of reactive organic matter to induce persistently reducing conditions at the SWI.

The first step for Mn(Ca)CO$_3$ authigenesis, i.e. the reduction of MnO$_x$, may be mediated directly by porewater sulphide (e.g. Burdige 1993; Yao and Millero 1993) or microbes (e.g. Aller and Rude 1988; Neumann et al. 2002; Tebo et al. 2004; Lee et al. 2011; Henkel et al. 2019). Active bacterial sulphate reduction and the involvement of sulphide in MnO$_x$ reduction is supported by the rapid drop in sulphate concentrations and the peak in sulphide concentrations at the SWI (Figs. 6 and 7). In addition, bacterial sulphate reduction may also result in significant alkalinity, thus favouring Mn(Ca)CO$_3$ precipitation (Berner et al. 1970; Huckriede and Meischner 1996).

The calcite-rich top portion of the sediment (Fig. 2) suggested that dissolution of calcite at the SWI is a potential source of carbonate ions for the Mn(Ca)CO$_3$ authigenesis; however, the Mn intervals at 31 cm and 25 cm within the deepest core contained no Ca-enrichments, which suggest that the presence of calcite was not obligatory for past Mn(Ca)CO$_3$ formation (Fig. 4). The slightly basic pH of the porewater (Fig. 6) does not support calcite dissolution as a source of Ca and carbonate ions. More likely, carbonate ions may be the product of oxidised organic matter. Indeed, based on low $\delta^{13}$C isotope ratios (Suess 1979; Huckriede and Meischner 1996; Lepland and Stevens 1998; Stevens et al. 2000; Neumann et al. 2002), a significant portion of the carbonate involved in Mn(Ca)CO$_3$ formation must come from organic matter degradation. In contrast, Pedersen and Price (1982) reported $\delta^{13}$C data that indicated a non-organic source of carbonate in Mn(Ca)CO$_3$. Nevertheless, the degradation of organic matter—the sedimentation of which has increased since the onset of eutrophication of Lake Stechlin—likely has provided the carbonate ions for the Mn(Ca)CO$_3$ authigenesis. The addition of Ca-ions to the mixed Mn-Ca phase does not necessarily require the dissolution of CaCO$_3$ to Ca$^{2+}$ as the water of Lake Stechlin is already rich in Ca$^{2+}$ (Fig. 6). The decrease of Ca$^{2+}$ in the porewater during the summer stagnation period also does not support the scenario of CaCO$_3$ dissolution. Instead, it points to the precipitation of Ca$^{2+}$ as Mn(Ca)CO$_3$. During the oligotrophic period, conditions necessary for Mn carbonate formation were probably not met, and Mn was cycling between the water column and the sediment without being buried.

Fluxes and redistribution of manganese

Calculations of internal Mn fluxes were made to explain the source and sedimentation patterns of Mn
and were summarised in Fig. 12. The flux of porewater Mn\textsubscript{diss} below 58 m water depths cannot explain the Mn accumulation into the 60 m trap alone; the porewater fluxes measured at 58 m and 69.5 m did not exceed 16.8 g Mn m\textsuperscript{-2} year\textsuperscript{-1}, so were much lower than the Mn accumulation rate of 50.9 g m\textsuperscript{-2} year\textsuperscript{-1} in the trap at 60 m. Therefore, porewater fluxes from shallower areas must contribute as well. Although distinctly lower compared to the central lake, a Mn flux into the overlying water also occurs in the porewaters of the shallower areas. Thus, Mn release rates vary between 0.3 and 36 mg Mn m\textsuperscript{-2} day\textsuperscript{-1} on a transect carried out from 17 to 68 m water depth in May–July 2017 (Lungfiel 2018; Fig. S2 and Table S1 in supplementary material). Relating this data to the large source area between 0 and 60 m water depth (4.15 km\textsuperscript{2}), a total Mn mass of 3078 kg year\textsuperscript{-1} (0.74 g m\textsuperscript{-2} year\textsuperscript{-1}) could be theoretically released into the water, and 37.2 g m\textsuperscript{-2} year\textsuperscript{-1} could be deposited in the small sink area (0.08 km\textsuperscript{2}) below 60 m depth. Despite existent inaccuracies, this is in the order of magnitude of the mean additional Mn accumulation in the sediments (50.9 g m\textsuperscript{-2} year\textsuperscript{-1}; Fig. 12). Lake-external Mn inputs are excluded as significant sources for Mn enrichment in deeper parts because there were no major environmental changes in the watershed area in the last decades that would explain an intensified Mn input to the lake and its geochemical focusing. Groundwater as the presumed main source of Mn could not have caused such Mn enrichment at the deepest sites directly, as it enters Lake Stechlin at the shoreline (Pöschke et al. 2018).

Manganese enrichment at the sediment surface was present below the 56 m water depth (Fig. 12), although a certain portion was not permanently fixed as Mn(Ca)CO\textsubscript{3} but was still present as MnO\textsubscript{x} (Fig. 10) and will be subject to ongoing redistribution. The difference between total Mn accumulated at the SWI at the deepest site in 2010 (125 g m\textsuperscript{-2}) and 2017 (940 g m\textsuperscript{-2}) is 815 g m\textsuperscript{-2}. However, the sediment trap at 60 m in the water column above the deepest site recorded a Mn accumulation of just 297 g m\textsuperscript{-2} during this period. The difference between the amount of Mn accumulated in the sediment and the sediment trap is explained by the lateral, near-sediment transport of an additional 518 g m\textsuperscript{-2} year\textsuperscript{-1} beneath the sediment trap at 60 m. Furthermore, the highest mass per square meter of Mn\textsubscript{diss} in the water column above the 69.5 m site was obtained on 8 August 2017, and equalled 5.5 g m\textsuperscript{-2} (Fig. 12). This amount is approximately 10-times smaller than the accumulation in the trap at 60 m (50.9 g year\textsuperscript{-1}) and is much lower than the average known amount of deposited Mn in the period 2010–2017 (116 g year\textsuperscript{-1}). Therefore, the Mn measured in the water column cannot explain the overall Mn accumulation rates, but instead, MnO\textsubscript{x} must be repeatedly dissolved and precipitated.

Geochemical focusing

Based on the observations and mass balance calculations, a conceptual model of the geochemical focusing and diagenesis of Mn in Lake Stechlin was developed (numbers 1 through 8 in Fig. 13). (1) The intensification of all redox-related processes at the SWI can be attributed to the supply of labile organic matter due to increased primary production during a recent period of eutrophication. (2) In addition, pH seasonally increases due to CO\textsubscript{2} uptake by phototrophic organisms leading to calcite precipitation from the epilimnion in summer. (3) The degradation of organic matter (OM) consumes O\textsubscript{2} leading to anoxic conditions at the SWI resulting in a reduction of particulate MnO\textsubscript{x} to Mn\textsubscript{diss} in the porewater forming a peak at the SWI (Fig. 6). Because Mn\textsubscript{diss} has already accumulated in the water column during the beginning of the stagnation period (26 April 2018 in Fig. 5), the mobilisation of Mn can be attributed to the reductive dissolution of MnO\textsubscript{x} by organic matter degradation coming from algae blooms in spring, rather than to stagnation duration. A similar increase in Mn\textsubscript{diss} in porewater and in water column during spring/summer has been reported in Lake Mendota (Wisconsin, USA) by Delfino and Lee (1971), Oneida Lake (New York, USA) by Aguilar and Nealson (1998), the seasonally anoxic Lake Rostherne Mere, UK (Davison and Woof 1984), and Esthwaite Water, UK (Hamilton-Taylor et al. 1996). The long water residence time of Lake Stechlin of > 40 years (Koschel and Adams 2003), resulting from a lack of major watercourses, is probably a major factor enabling Mn cycling and prevents the outflow of Mn\textsubscript{diss}. (4) After the reductive dissolution of MnO\textsubscript{x} and diffusion of Mn\textsubscript{diss} into the water column (Fig. 5), Mn\textsubscript{diss} is oxidised to MnO\textsubscript{x} (Figs. 10 and 11). (5) The transportation and focusing of the particulate MnO\textsubscript{x} towards greater water depths can be explained by diffusion of Mn\textsubscript{diss} within the water column and, as proposed by Schaller and Wehrli (1996), by the
Fig. 12 Balance of Mn fluxes in Lake Stechlin. The maximum encountered total Mn$_{diss}$ mass in the water column was measured on 8 August 2017 (WD: water depth); dark-blue text = total accumulation of Mn at the sediment surface (maximum depth = 8 cm) sampled in 2017 (deepest core) and 2018 (other cores); purple area = Mn-rich layer with an increasing thickness with the water depth; green text = diffusion of Mn$_{diss}$ from porewaters at 58 m and 69.5 m; purple text = net accumulation rates into the sediment traps at 20 m and 60 m for 2017 and 2010–2017 (right, black); bottom right = net difference between Mn accumulation at the top of the core sampled in 2010 (125 g m$^{-2}$) and the total Mn accumulation in the core sampled in 2017 (940 g m$^{-2}$). The gap between the sedimentation in the trap and the net sedimentation at the lake bottom is explained by an additional lateral flux of 518 g m$^{-2}$ below the trap. (Color figure online)

Fig. 13 Left: Conceptual model of geochemical focusing of Mn and the formation of Mn(Ca)CO$_3$ in Lake Stechlin. Numbers 1–8 represent the key processes involved and are explained in the main text. Purple layers represent buried Mn. Right: The porewater Mn flux during the oligotrophic phase was likely essentially lower due to a deeper O$_2$ penetration into the sediment and less organic matter deposition limiting dissimilatory MnO$_x$ reduction as well as bacterial sulphate reduction finally causing lower Mn$_{diss}$ concentrations.
turbulent mixing of dissolved or colloidal Mn throughout the basin. Mass balance calculations showed that MnOx is transported through the water column from shallow depths towards the deepest site close to the sediment surface. (6) MnOx then accumulates at the deepest point in the lake where, owing to renewed reductive dissolution, porewater Mn_diss concentrations increase. (7) Simultaneously, carbonate-ions are supplied either by organic matter degradation or calcite dissolution. (8) Finally, Mn(Ca)CO3 precipitation is facilitated by high Mn_diss concentrations potentially supported by bacterial mediation.

In this model, shallow-water sediments act as the source of Mn_diss, whereas sediments in the deepest part of the lake act as a net sink of particulate Mn. Consequently, geochemical focusing is not expected to occur if a geochemically resistant mineral phase is formed (e.g. Mn(Ca)CO3) at shallower water depths or, more generally, in the source area. As described by Schaller and Wehrli (1996), such geochemical focusing occurs when an anoxic sediment environment is in contact with an oxic water column. Accordingly, the reason why geochemical focusing of Mn in Lake Stechlin was not active during the pre-eutrophication phase was probably because of dissolved oxygen that could penetrate sufficiently deep into the sediment to prevent substantial Mn_diss diffusion out of the sediment (Fig. 13). The role of organic matter should be emphasised, as the TOC-rich sediments of Lake Stechlin provide a sharp redox gradient in contact with the (sub-)oxic water column, as evidenced by the sharp gradient of dissolved sulphide concentration just below the SWI (Fig. 7). Quite similar to our findings in Lake Stechlin, Mn(Ca)CO3 is found occasionally in organic-rich marine sediments (e.g. Calvert and Pedersen 1996; Häusler et al. 2018).

Previous studies on Mn as a redox-proxy (e.g. Mackereth 1966; Wersin et al. 1991; Naehler et al. 2013) have stated that increasing Mn/Fe ratios are indicative of oxic conditions. Our findings show the opposite (Fig. 3), whereby eutrophication and more reducing conditions caused the geochemical focusing and burial of Mn as Mn(Ca)CO3, showing that geochemical focusing may occur during a transition from oligotrophic to mesotrophic conditions. In Lake Stechlin (this study) and Lake Zurich (Naehler et al. 2013), geochemical focusing caused the enrichment of Mn towards the deepest sites, reflected by high Mn/Fe ratios in comparison to other sites. However, Mn geochemical focusing began in Lake Zurich during oligotrophic conditions from eutrophic towards mesotrophic conditions (Naehler et al. 2013), whereas in Lake Stechlin, the process was active during eutrophication from oligotrophic towards mesotrophic conditions. During periods with O2-free bottom waters in Lake Zurich between 1940 and 1950 (Naehler et al. 2013), insufficient Mn availability in the sediments due to reductive dissolution of MnOx and coupled Mn_diss escape into the water column probably inhibited the formation of stable Mn minerals including rhodochrosite. Therefore, we conclude that similar to the Baltic Sea (Häusler et al. 2018) geochemical focusing and Mn(Ca)CO3 formation may occur only when bottom waters are replenished with O2 containing waters (e.g. by annual mixing or water inflows), such that MnOx can accumulate at the sediment surface and a stable Mn mineral phase forms during renewed reductive dissolution.

Our measurements of O2 concentrations in the water column suggest that oxic or suboxic—but not sulphidic—conditions occurred throughout the year until approximately 2010. Since then, anoxic (or even sulphidic) bottom waters appeared at the end of the summer stagnation period. However, if climatic warming prolongs the stratification period of Lake Stechlin and, additionally, the ongoing eutrophication results in increasingly O2-depleted bottom waters, the lake may become meromictic so that geochemical focusing and coupled Mn(Ca)CO3 formation may become inhibited.

Conclusions

We have shown that Mn enrichment of Lake Stechlin is linked to a recent period of eutrophication in this once oligotrophic lake. Dissolved Mn originates from shallow and deep sediments and diffuses out of the sediment into the oxic-suboxic water column. Dissolved Mn is oxidised to MnOx particles, transported through the water column close to the sediment surface, and is deposited in the deepest areas of the lake as a part of a varved section. The burial of Mn was enabled by the partial conversion of MnOx into Ca-rich rhodochrosites (Mn(Ca)CO3) shortly after sedimentation. The formation of Mn(Ca)CO3 does not occur at the shallower sites, most likely due to the low Mn_diss concentrations resulting from the low MnOx.

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content of the sediments. Increased primary production, concomitant with the eutrophication, provides organic matter that intensifies various biogeochemical diagenetic processes upon degradation. Further Mn peaks buried deeper in the sediments at the deepest areas of the lake indicate that the geochemical focusing process has also been active in the past. As such, Mn may be used as a proxy of paleo-redox dynamics. In accordance with the concept of the geochemical focusing and studies from the deep Baltic basins, the redox-driven re-sedimentation of Mn seems possible only if anoxic sediment is in contact with an O₂-containing water column. Our study of Lake Stechlin also shows that oligotrophic lakes that experience eutrophication may be subject to geochemical focusing and parameters including lake morphology (e.g. the relatively small area of the deepest site), organic-rich sediment, and a long water residence time may favour this phenomenon. In comparison to previous research, Mn/Fe values were found to function oppositely in our study, and as such, the applicability of this ratio as a redox-proxy requires further assessment.

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