Using small-scale measurements to estimate hypolimnetic oxygen depletion in a deep lake

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Abstract

Low oxygen concentrations in lakes and reservoirs are an ongoing environmental concern, particularly in light of increasing anthropogenic activity and climate change. Oxygen depletion processes in lakes are still not completely understood and a variety of models have been proposed based on limited field observations. Here, we present field measurements of oxygen depletion processes in a deep lake, Lake Geneva (Switzerland). The aim of this study was to quantify three basic processes controlling hypolimnetic oxygen depletion and their relative contribution to the total oxygen depletion (TOD) rate. Sediment oxygen uptake (SOU) and the flux of reduced substances were estimated based on oxygen microprofile measurements and sediment core data of reduced substances. Acoustic Doppler current profiler measurements and hydrodynamic modeling were used to ensure that SOU was measured under typical hydrodynamic conditions. Comparison with long-term monitoring data allowed for an estimate of the relative importance of SOU and water column mineralization (WCM). Results show a decrease in both SOU and WCM down to mid-depth which could not be explained by changes in hydrodynamic conditions or temperature. Below mid-depth, TOD increased due to an enhanced sediment area to water volume ratio \((a)\). This vertical pattern of oxygen depletion is driven by (1) lake morphometry paired with increasing \(a\), and (2) decreasing organic matter mineralization in the water column with depth. The findings are explained by a model which separates the oxygen depletion into an exponentially decreasing component, representing the fast-decaying fraction of the organic matter, and a constant background component.

Efficient lake management is based on accurately describing and modeling physical and biogeochemical processes. Dissolved oxygen (DO) demand is one of the most important parameters for characterizing lake ecosystem health and, consequently, DO concentrations in lakes have been intensely studied since the end of the 19th century (e.g., Thienemann 1928; Hutchinson 1957). Since the middle of the last century, anthropogenic pressure has caused eutrophication and temperature increases that negatively affect the DO budget of lakes and oceans. Thus, DO falls often below biologically acceptable concentrations (Diaz 2001; Friedrich et al. 2014; North et al. 2014).

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Additional Supporting Information may be found in the online version of this article.
Oxygen depletion is caused by mineralization of organic matter in the water column or within the upper sediments. Aerobic respiration is the dominant process for organic matter mineralization under oxic conditions. However, under anoxic conditions (e.g., in the deep water of meromictic lakes or in anoxic sediments), other electron acceptors come into play leading to the formation of reduced compounds including CH$_4$, NH$_4^+$, Mn(II), and Fe(II), as in the deep water of meromictic lakes or in anoxic sediments. These substances diffuse toward the oxic zones of the sediment or into the open water and are then oxidized, thereby contributing to further DO uptake. These fluxes of reduced substances can contribute significantly to DO depletion in eutrophic lakes (Gelda et al. 1995; Matzinger et al. 2010). Fluxes of reduced substances and oxic respiration depend on the efficiency of organic matter mineralization. The mineralization rate is in turn dependent on several parameters such as temperature (Gudasz et al. 2010) or organic matter quality (Arndt et al. 2013) which may show extensive variation within a lake. Current lake DO models use either simplified empirically derived expressions that do not quantitatively describe the complex processes upon which they are built (Fang and Stefan 2009; Rucinski et al. 2010) or they rely on a large number of free parameters that require calibration based on empirical field data (Rucinski et al. 2014). For both approaches, good knowledge of the sinks of DO and its spatial variability is crucial to improving the reliability and predictive power of the models. Hence, detailed measurements of the SOU at different lake depth and the relative importance of the flux of reduced substances for the total oxygen depletion (TOD) are essential to understand and reproduce oxygen depletion processes.

In this study, we investigate DO depletion in Lake Geneva, which is currently recovering from anthropogenic eutrophication but shows still high levels of DO depletion and periodic hypoxia in the deep hypolimnion (Schwefel et al. 2016). Sediment-water microprofile measurements were performed at different depths of the lake to determine the spatial variability of SOU (i.e., the flux of DO from the water toward the sediment surface). Simultaneous pore-water analyses of sediment cores allowed for the quantification of the fluxes of reduced substances out of the sediment. Combined with a set of long-term monitoring data from over more than 40 yr, collected from the water column at the deepest location of the lake, these observations allow for a distinction between the various pathways of DO consumption as a function of depth. The aims of this study were to
characterize the spatial distribution and variability of DO depletion, (2) estimate the relative importance of the two sediment molecular fluxes: the SOU and the reduced substances flux from the sediment, and (3) explain these findings within a conceptual basin-scale oxygen budget.

Methods

Study site

Lake Geneva is a deep perialpine lake situated on the border of France and Switzerland with a maximum and mean depth of 309 m and 153 m, respectively, a surface area of 582 km$^2$, and a volume of 89 km$^3$. Once considered oligotrophic, Lake Geneva has experienced high nutrient inputs since the 1950s. Phosphorus concentrations increased from 15 mg m$^{-2}$ around 1960 to up to 90 mg m$^{-2}$ in the 1980s. Consequently, water transparency decreased, the phytoplankton communities shifted to species more typical for eutrophic environments, and phyto- and zooplankton biomass increased (Anneville and Pelletier 2000; Loizeau and Dominik 2005). While DO has been monitored in the deep hypolimnion since the late 1890s (e.g., Forel 1895; Delebecque 1898; Vivier 1944), low DO concentrations had never been observed prior to 1957, either via direct field measurements (Delebecque 1898; Vivier 1944) or sediment core sampling (Jenny et al. 2014). Since then, periods of hypoxia have been observed regularly (Savoye et al. 2015; Fig. 3).

Recent governmental measures to limit the phosphorus input into the lake reduced total phosphorus to the present concentrations of ~20 mg m$^{-3}$. Despite the high reduction in nutrient loads, total biomass has remained high and DO
depletion rates and concentrations have not shown any significant trend over the last 40 yr (Anneville and Pelletier 2000; Savoye et al. 2015; Schwefel et al. 2016; Figs. 1, 3).

### Microprofile measurements

In total, 97 DO profiles were measured using a MP4/8 Microprofiler (Unisense A/S) equipped with oxygen microsensors during eight different measurement campaigns between autumn 2013 and summer 2017 (Table 1). Measurements were performed in the northwestern part of the deep basin of Lake Geneva in depth between 43 m and 224 m (Table 1). The microprofiler was deployed for between 1 d and 3 d for each individual campaign; measurement locations and dates are shown in Fig. 4 and summarized in Supporting Information Table S1. The Clark-type oxygen microsensors had a tip size of 100 \( \mu \)m with a detection limit of 0.01 mg L\(^{-1}\) and a response time of <8 s. The vertical spatial resolution of the measurements is limited by the diameter of the sensor tip. An RBR concerto data logger (RBR, Ottawa) equipped with an oxygen optode (Aanderaa Data Instruments) as well as a temperature and pressure sensor was attached to the microprofiler frame and positioned outside of the diffusive boundary layer (DBL) at \( \sim \) 25 cm above the sediment to provide background time series of DO concentrations in the BBL. To calibrate the oxygen microsensors in the field, we performed a two-point, linear calibration with the in situ data from the optode as the upper reference point and the anoxic sediment as the lower reference point.

The detection of the sediment-water interface (SWI) is critical when investigating SOU. In Lake Geneva, sediment and water are distinctly separated by a defined SWI as observed during MIR submersible measurements in 2014.

### Table 1. Summary of the microprofile measurement results. The sediment oxic zone is defined as the layer, in which the oxygen concentration stays above 0.1 mg L\(^{-1}\).

<table>
<thead>
<tr>
<th>Number</th>
<th>Depth (m)</th>
<th># of profiles</th>
<th>SOU (gDO m(^{-2}) d(^{-1}))</th>
<th>DBL thickness (mm)</th>
<th>Sediment oxic zone (mm)</th>
<th>BBL temperature (°C)</th>
<th>BBL DO concentration (g m(^{-3}))</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>MP1</td>
<td>43</td>
<td>4</td>
<td>1.08 ± 0.51</td>
<td>1.1 ± 0.8</td>
<td>2.1 ± 0.1</td>
<td>6.6</td>
<td>8.8</td>
<td>Oct 2013</td>
</tr>
<tr>
<td>MP2</td>
<td>45</td>
<td>12</td>
<td>1.08 ± 0.33</td>
<td>0.8 ± 0.6</td>
<td>2.4 ± 0.5</td>
<td>6.6</td>
<td>10.3</td>
<td>Mar 2014</td>
</tr>
<tr>
<td>MP3</td>
<td>72</td>
<td>7</td>
<td>0.50 ± 0.11</td>
<td>1.5 ± 0.8</td>
<td>2.9 ± 0.7</td>
<td>6.6</td>
<td>10.1</td>
<td>May 2014</td>
</tr>
<tr>
<td>MP4</td>
<td>76</td>
<td>13</td>
<td>0.46 ± 0.16</td>
<td>1.4 ± 0.6</td>
<td>3.8 ± 1.0</td>
<td>6.4</td>
<td>9.4</td>
<td>Jul 2014</td>
</tr>
<tr>
<td>MP5</td>
<td>93</td>
<td>30</td>
<td>0.32 ± 0.16</td>
<td>2.0 ± 1.3</td>
<td>4.5 ± 0.7</td>
<td>6.3</td>
<td>8.5</td>
<td>Aug 2015</td>
</tr>
<tr>
<td>MP6</td>
<td>109</td>
<td>13</td>
<td>0.39 ± 0.07</td>
<td>1.7 ± 0.4</td>
<td>4.3 ± 0.3</td>
<td>5.9</td>
<td>8.7</td>
<td>Jul 2014</td>
</tr>
<tr>
<td>MP7</td>
<td>133</td>
<td>13</td>
<td>0.21 ± 0.06</td>
<td>2.1 ± 0.8</td>
<td>5.0 ± 0.8</td>
<td>6.0</td>
<td>7.8</td>
<td>Jul 2015</td>
</tr>
<tr>
<td>MP8</td>
<td>224</td>
<td>5</td>
<td>0.14 ± 0.14</td>
<td>1.9 ± 1.2</td>
<td>4.0 ± 0.5</td>
<td>5.8</td>
<td>4.5</td>
<td>May 2017</td>
</tr>
</tbody>
</table>

Fig. 4. Left panel: Bathymetry of Lake Geneva. Right panel: Measurement locations, red diamonds: microprofiles (Table 1, MP1 and MP2 as well as MP3 and MP4 are displayed as one point as they are too closely located to be resolved), white circles: sediment cores (Table 2), red cross: long-term monitoring point SHL2.
(Wüst et al. 2014). As our instrument was very slowly but continuously sinking into the sediment throughout the experiments (by less than 1 mm h\(^{-1}\)), we had to perform accurate SWI detection measurements prior to each profile. We programmed the microprofiler so that before each profile, the microsensor was first lowered continuously until the DO signal was reduced to 60% of the bulk BBL DO concentration. This drop was defined as a good proxy for identifying the close proximity of the DBL. Afterward, the sensors were raised by \(\frac{C24}{24}\) mm and the high-resolution DO microprofile measurements were started in typically 150–300 vertical steps of 100 \(\mu\)m fully covering the water-side DBL and the sediment oxic zone beneath. Per profile, sediment detection and profiling generally lasted about 70 min and varied depending on the number of steps and the time needed for sediment detection. Final determination of the SWI was based on visual identification of the gradient change in the DO profiles across the SWI that results from differing diffusivities between water and sediment (Røy et al. 2004; Bryant et al. 2010). Analysis of the DO variance provided additional evidence for the correct location of the SWI (Schwefel et al. 2017).

The SOU was estimated from the water-side DO gradient based on Fick’s law:

\[
\text{SOU} = D \frac{\partial O_2}{\partial z},
\]

where \(D\) is the temperature-dependent molecular diffusion coefficient of oxygen in water. \(\frac{\partial O_2}{\partial z}\) is the DO gradient in the linear region above the SWI where \(z\) is defined positive downward. Hence, positive values indicate fluxes into the sediment. The lower boundary of the sediment oxic zone was defined by the depth where DO concentrations fell below 0.1 mg L\(^{-1}\) (\(\sim 3 \mu\text{mol L}^{-1}\) as in Bryant et al. 2010). Finally, the thickness of the DBL was calculated by extrapolating \(\frac{\partial O_2}{\partial z}\) up to the point where DO(\(z\)) reaches the averaged bulk DO concentrations of the BBL. In Jørgensen and Revsbech (1985), this term is called the “effective DBL.”

**Current measurements and modeling**

Flow measurements were performed with an upward-looking 2 MHz acoustic Doppler current profiler (ADCP; Nortek Aquadopp). The ADCP was deployed close to the microprofiler (\(\sim 100\) m away) to quantify the BBL flow at the same time as the SOU estimate. During all deployments, the vertical resolution was 5 cm. Due to the blanking distance of the ADCP, the first measurement cell was \(\sim 20\) cm above the sediment. Measurements were performed in burst mode in 20 min intervals with 256–1024 measurements per burst. Due to instrument and/or deployment issues, velocity data were not available for all deployments (see Supporting Information Table S1). The red dashed line gives the mean observed velocities 1 m above bottom measured by an ADCP during the microprofile measurements. For measurements MP1, MP3, MP4, and MP8, no ADCP data were available.

The hydrodynamic conditions of the four relatively short (< 1 week) measurements periods were compared to typical conditions in Lake Geneva by using the three-dimensional hydrodynamic model Delft3D (Razmi et al. 2013). The model provided a map of the average speed of the BBL currents at 1 m above the sediment during the year 2012. Over a 1 yr period, we observed a clear decrease in the bottom currents with depth, as shown in the contour map in Fig. 5a and in the frequency distribution at different measuring locations in Fig. 5b. The modeled bottom current distribution indicates that at the 43 m deep site, measurements were collected during comparatively quiet conditions compared to long-term BBL currents at this location. During the measurements at the deeper sites, the flows were close to the modeled mean bottom currents (Fig. 5b).
Sediment cores

A total of 16 sediment cores were collected with a UWI-Tec corer equipped with a PVC tube (6.5 cm inner diameter, 60 cm length) at eight different sites from 40 m to 310 m depth (Fig. 4; Table 2). At each site, one core was sampled for the analysis of CH₄ via gas chromatography and a second core was sampled for the measurement of pore-water concentrations of cations and anions using capillary electrophoresis (Kubán et al. 2007; Torres et al. 2013).

For pore-water retrieval, PVC tubes with pre-drilled holes of 2 mm diameter at 5 mm vertical intervals were used. The holes were sealed with adhesive tape prior to sampling. Resolution of the measurements was 0.5 cm for the first 10 cm and 1 cm for the next 10 cm of the sediment. Immediately upon retrieval, the cores were brought to the lab and 10–50 L of sediment pore water were sampled by insertion of a MicroRhizon filter tube (1 mm diameter, 0.20 μm pore size; Rhizosphere Research Products) through the pre-drilled holes. Additionally, one sample was taken from the overlying water. Measurements were performed with two portable capillary electrophoresis systems equipped with capacitive coupled contactless conductivity detectors. Cations were analyzed in all eight cores. The analysis of anions was omitted for logistical reasons in the 2014 campaign. Full separation of cations of interest (NH₄⁺, Mn(II), and Fe(II)) was achieved within 6 min. Standard deviations of all measurements were < 5%. The setup is described in detail by Torres et al. (2013) and Steinsberger et al. (2017).

For the CH₄ measurements, holes of 1.2 cm diameter were drilled staggered at 1-cm vertical intervals in the sediment sampling tube and covered with adhesive tape. Immediately after retrieval, cores were sampled from top to bottom by cutting the tape and inserting a plastic syringe which had the tip cut off. Two milliliters of sediment were transferred into 125 mL serum flasks prefilled with 2 mL of 7 M NaOH and sealed with butyl rubber stoppers. Methane was analyzed by headspace gas chromatography (Agilent) using a 1010 Supelco Carboxene column.

The fluxes of reduced substances (F₈CH₄, F₈NH₄⁺, F₈Mn, and F₈Fe for CH₄, NH₄⁺, Mn(II), and Fe(II), respectively) were calculated based on the concentration gradients measured in the topmost layer of the sediment cores. A one-dimensional balance model originally developed by Epping and Helder (1997) for oxygen and adapted to other dissolved substances was used to estimate species fluxes. The model, a reaction-diffusion model based on the assumption of steady-state conditions in the sediment, is described in detail in Müller et al. (2003) and Steinsberger et al. (2017). The modeled fluxes of reduced substances were then converted to oxygen equivalents based on redox stoichiometry (Matzinger et al. 2010) by:

\[
F_{\text{red}} = 2F_{\text{CH}_4} + 2F_{\text{NH}_4} + 0.5F_{\text{Mn}} + 0.25F_{\text{Fe}} \tag{2}
\]

where \( F_{\text{red}} \) represents the sum of all reduced substances (estimated in [mol m⁻² d⁻¹]) and expressed in oxygen equivalents [g eq O₂ m⁻² d⁻¹] by using Eq. 2.

Monitoring data

Since 1957, DO measurements were performed on a regular basis by the Commission International pour la Protection des Eaux du Léman (CIPEL). DO was measured along with other parameters (e.g., total phosphorus, transparency, and temperature) at the deepest point of the lake (SHL2, see Fig. 4) throughout the entire water column for 8–12 times per year between 1957 and 1970 and at least once per month thereafter to present. Comparisons between the measured DO concentrations in the water column of SHL2 and four other offshore measurement stations at six different depths show only minor variations of 6% on average with higher variability in spring as compared to summer and autumn (Supporting Information Fig. S1). Therefore, data from SHL2 was considered to be representative for the entire lake during stratified summer conditions.

Results

Microprofile measurements

Two typical microprofiles from 45 m and 133 m depth (Table 1) are shown as representative examples in Fig. 6. The BBL temperatures were measured simultaneously with the DO microprofiles during each deployment. While BBL temperatures showed only minor differences between individual measurements, BBL DO concentrations decreased with depth and were, as expected, lower in autumn compared to spring and summer (Table 1). The mean SOU, sediment oxic zone depth (i.e., oxygen penetration depth), and BBL temperature

### Table 2. Flux of reduced substances in units of oxygen equivalents (g eq O₂ m⁻² d⁻¹) using Eq. 2.

<table>
<thead>
<tr>
<th>Depth (m)</th>
<th>LG1</th>
<th>LG2</th>
<th>LG3</th>
<th>LG4</th>
<th>LG5</th>
<th>LG6</th>
<th>LG7</th>
<th>LG8</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>40</td>
<td>0.029</td>
<td>&lt;0.001</td>
<td>0.003</td>
<td>0.035</td>
<td>0.051</td>
<td>0.039</td>
<td>0.005</td>
<td>0.056</td>
<td>0.027 (55%)</td>
</tr>
<tr>
<td>Ammonium</td>
<td>0.017</td>
<td>0.027</td>
<td>0.010</td>
<td>0.019</td>
<td>0.033</td>
<td>0.013</td>
<td>0.008</td>
<td>0.010</td>
<td>0.017 (35%)</td>
</tr>
<tr>
<td>Manganese(II)</td>
<td>&lt;0.001</td>
<td>0.003</td>
<td>0.002</td>
<td>0.004</td>
<td>0.001</td>
<td>0.002</td>
<td>0.006</td>
<td>0.005</td>
<td>0.003 (6%)</td>
</tr>
<tr>
<td>Iron(II)</td>
<td>0.003</td>
<td>0.002</td>
<td>0.003</td>
<td>0.006</td>
<td>0.002</td>
<td>&lt;0.001</td>
<td>0.002</td>
<td>&lt;0.001</td>
<td>0.002 (4%)</td>
</tr>
<tr>
<td>Sum</td>
<td>0.049</td>
<td>0.032</td>
<td>0.018</td>
<td>0.064</td>
<td>0.086</td>
<td>0.055</td>
<td>0.021</td>
<td>0.071</td>
<td>0.049</td>
</tr>
</tbody>
</table>
and DO concentration at the different measurement sites are summarized in Table 1. Figure 7 shows the variation of SOU with depth. The individually estimated SOU rates, based on averages from 4 to 30 profiles per site, varied strongly during each measurement session (Table 1). These variations can be partially explained by BBL turbulence dynamics that was also observed in other studies (Lorke et al. 2003; Bryant et al. 2010a). In addition to temporal variations in the individual profiles at each site, SOU also showed strong spatial variability among the different measurement sites with a distinct decrease with depth ranging from $1.08 \pm 0.51 \text{ g DO m}^{-2} \text{d}^{-1}$ at 45 m depth to only $0.14 \pm 0.14 \text{ g DO m}^{-2} \text{d}^{-1}$ at 224 m depth.

In parallel, the sediment oxic zone increased with depth from $\sim 2 \text{ mm}$ at 43 m depth to $\sim 5 \text{ mm}$ at 133 m depth (Table 1). At 224 m, the sediment oxic zone was slightly shallower (4 mm), which was most likely due to the generally lower oxygen concentrations present during measurements at this depth.

**Sediment cores**

In the top few millimeters of the sediment, DO is depleted by two processes: (1) aerobic respiration in the sediment and (2) reduced substances fluxes from the deeper sediment (Fig. 2). The latter leads to a DO consumption if the reduced substances diffuse into the oxic zone of the sediment. An unknown fraction of the reduced substances is oxidized in the sediment, thereby contributing to the SOU (Frenzel et al. 1990; Bryant et al. 2011). The remaining fraction diffuses out of the sediment where it is oxidized in the overlying oxic water column (as quantified by $F_{\text{red}}$).

$F_{\text{red}}$ was calculated based on the sediment pore-water concentrations of reduced substances and converted to oxygen equivalents according to Eq. 2. $F_{\text{red}}$ varied from only $0.018 \text{ g DO m}^{-2} \text{d}^{-1}$ to $0.086 \text{ g DO m}^{-2} \text{d}^{-1}$ with a mean of $0.049 \pm 0.024 \text{ g DO m}^{-2} \text{d}^{-1}$ among the different measurement sites (Fig. 8; Table 2), which is only 3.6% of the total hypolimnetic oxygen depletion in Lake Geneva and is
extremely small compared to medium-sized lakes with comparable eutrophication histories (Matzinger et al. 2010; Steinsberger et al. 2017). $F_{\text{red}}$ estimated in cores from the deepest lake zones of ~ 300 m, varied between ~ 0.06 $g_{\text{DO}}$ m$^{-2}$ d$^{-1}$ and 0.09 $g_{\text{DO}}$ m$^{-2}$ d$^{-1}$. Cores taken at sites between 40 m and 200 m depth showed generally lower $F_{\text{red}}$ fluxes (0.02–0.05 $g_{\text{DO}}$ m$^{-2}$ d$^{-1}$). An exception is core LG7, taken at depth 200 m (0.07 $g_{\text{DO}}$ m$^{-2}$ d$^{-1}$), where $F_{\text{red}}$ is comparatively high, but still very low in absolute terms. This particular site, however, may be influenced by the close proximity of the City of Evian or the inflow of the river Dranse which could lead to locally increased organic matter concentrations due to wastewater inputs. Unfortunately, no microprofile measurements were performed to characterize sediment-water oxygen dynamics at this location as the steep bathymetry is not well suited for microprofiler stability. At sites where parallel microprofile and sediment core measurements were obtained (LG1, LG2, and LG3; Table 2), $F_{\text{red}}$ was consistently less than 10% of SOU. The most dominant reduced substances were CH$_4$ and NH$_4^+$ (Table 2), which is consistent with measurements in other Swiss lakes (Matzinger et al. 2010; Steinsberger et al. 2017).

**BBL turbulence**

BBL turbulence depends on the wind forcing prior to the measurement period. During microprofiling, the highest BBL turbulence was measured at 133 m; the lowest was at 45 m due to calm conditions shortly before the microprofiler deployment. All measurements were performed under hydrodynamic conditions typical for Lake Geneva, as shown by the comparison with the Delft3D model flows in the BBL (Fig. 5b). While DO flux and BBL thickness varied at each site depending on turbulent conditions, mean values of DO fluxes were considerably higher at the shallowest measurement site which also had the lowest BBL currents. In turn, BBL thickness tended to increase with depth.

**Long-term monitoring data**

For the years 1970–2012, monthly DO concentration data were interpolated spatially and temporally and the areal hypolimnetic mineralization (AHM) rate ($g_{\text{DO}}$ m$^{-2}$ d$^{-1}$; Fig. 2) as well as the TOD rate ($g_{\text{DO}}$ m$^{-3}$ d$^{-1}$) per depth layer were determined based on the linear decrease in DO during summer stagnation (Fig. 3). While TOD is a measure of the DO depletion per volume in every depth layer during summer (Fig. 2), AHM integrates the summer DO depletion (for ~ 180 d) over the entire hypolimnion (from the sediment to a water depth of 15 m; see definition in Fig. 2). Although AHM varied considerably with a mean value of 1.34 ± 0.34 $g_{\text{DO}}$ m$^{-2}$ d$^{-1}$, AHM shows no significant trend during the entire observed period. An analysis of the vertical structure of DO depletion rates shows that TOD decreases with depth down to ~ 150 m. Below 150 m depth, TOD increases again (Fig. 1). At each depth, the fraction of oxygen depletion caused by SOU (Fig. 7) was found to be significantly lower than TOD.

**Discussion**

**SOU and sediment oxic zone depth**

Despite strong variability in the physical boundary conditions, all microprofile measurements showed a consistent decrease of SOU with lake depth (Fig. 7). Measurement dates ranged from March to October, mostly during summer (June–August). Because of temporal variations and differing lake depths, DO concentrations in the BBL varied between 10 mg L$^{-1}$ and 7.8 mg L$^{-1}$ in the measurements above 150 m. In the measurement at 224 m depth, BBL DO concentration was 4.5 mg L$^{-1}$. Since the microprofiler was always deployed well below the thermocline, BBL temperatures varied only by ~ 0.8°C and did not alter organic matter mineralization and oxygen depletion significantly (Gudas et al. 2010). DO fluxes are partially dependent on the difference in DO concentrations between SWI and the BBL above the DBL (Eq. 1). Hence, lower BBL DO concentrations directly imply lower DO fluxes into the sediment. Nevertheless, the variation of measured SOU with lake depth was too high to be a result of this effect only. During measurements, the standard derivation of SOU was typically on the order of ~ 30% and some individual measurements were found to be more than 100% higher than the mean values. Per Eq. 1, these variations can partially be explained by (1) varying DO concentration differences between SWI and bulk BBL (Fig. 6) and/or (2) varying DBL thicknesses due to changes in BBL turbulence (Lorke et al. 2003; Bryant et al. 2010a; Schwefel et al. 2017). Based on microprofile measurements in Lake Alpnach, increased SOU has been shown to directly correlate with increased turbulence and decreased DBL thickness (Bryant et al. 2010a,b). However, the relationship between BBL turbulence and SOU could not explain the systematic decrease of SOU with lake depth in Lake Geneva. The estimated differences in the averaged SOU at 43 m and 224 m depth were ~ 700% while changes in BBL current speed were only ~ 50% (Fig. 5). During measurements at the shallowest site, which had the highest SOU, currents and, in turn, BBL turbulence was even lower than at the deeper sites, all of which had lower SOU rates (Fig. 5b).

DO depletion in the sediment depends on the amount of organic matter and its quality (Steinsberger et al. 2017). While parts of organic material are easily degradable (i.e., labile) and decompose relatively quickly, other parts degrade over longer time scales (Kristensen and Holmer 2001; Arndt et al. 2013; Bouffard and Perga 2016). In the shallower regions of a lake, the labile fraction is decomposed quickly in the water column or precipitates quickly to the sediment. Conversely, the organic matter in the deeper sediments consists mainly of more refractory organic carbon and the mineralization rate decreases with increasing lake depth (Sobek...
et al. 2009; Arndt et al. 2013). Similar observations were made for a marine system in the St. Lawrence Estuary (Alkhatab et al. 2012).

The deepening of the sediment oxic zone with increasing lake depth is fully consistent with this interpretation. While easily mineralized organic matter is more abundant in shallower sediments, the observed decrease in SOU with depth in Lake Geneva is strong evidence that the deeper parts of the lake are covered with more refractory organic matter which leads to lower mineralization activity and subsequently deeper sediment oxic zones. A similar deepening of the oxic zone was noticed in the studies of Müller et al. (2003) of the eutrophic Lake Sempach and by Maerki et al. (2013) reported an average SOU of 0.46 gDO m⁻² d⁻¹, whereas measurements in sediment cores obtained at the center of the lake resulted in a SOU of only ~0.2 gDO m⁻² d⁻¹ (Smith and Matisoff 2008) indicating a considerable decrease in SOU with lake depth.

In addition to implications for F_red and sediment-water oxygen dynamics, the sediment oxic zone depth is of particular interest for fish eggs, which only survive under oxic conditions. Given the typically thin oxic zone in the upper sediments, fish egg survival is not guaranteed even if DO concentrations in the overlying water remain relatively high. Fish eggs sinking even slightly into the sediment can easily experience anoxic conditions which are detrimental to their survival (Ventling-Schwank and Müller 1991). Swiss federal law therefore requires a minimum of 4 mg L⁻¹ DO in natural waters to facilitate natural fish reproduction. In Lake Geneva, however, the Lake Char (Salvelinus umbla) population is still not able to naturally reproduce and is artificially stocked, despite decades of reoligotrophication during which hypoxia occurred only rarely (Caudron et al. 2014). The thin sediment oxic zone observed during the current study, which was found to be particularly prevalent in the shallower hypolimnetic sediments, could explain this phenomenon (Ventling-Schwank and Müller 1991).

In Lake Geneva, the majority of soluble, reduced substances are mostly oxidized within the thin oxic zone which prevents these species from diffusing into the water column. Thus, three sediment cores (S1, S2, S3) were taken in close proximity to the microprobe measurement sites (between 100 m and 400 m distance) to allow for comparison of SOU and F_red. At 40 m depth, F_red was less than 5% of SOU (Fig. 8). This fraction increases with increasing lake depth, as SOU decreases while F_red remained relatively constant, but was still below 10% at 133 m. Thus, reduced substances might play a more important role in the deep hypolimnion. At 224 m, SOU was only 0.14 gDO m⁻² d⁻¹. However, SOU varied considerably between profiles at this site and the bulk BBL DO concentration was with 4.5 mg L⁻¹ lower than average, both of which could have led to an underestimation of SOU. The averaged F_red of the eight cores (Table 2) was 0.049 gDO m⁻² d⁻¹ or only 3.6% of the total AHM of 1.34 gDO m⁻² d⁻¹.

Ultimately, reduced substances were found to play only a minor role for the overall hypolimnetic oxygen depletion, expressed as AHM, in Lake Geneva despite high primary production. Müller et al. (2012a) reported relative contributions of up to 80% of AHM in very shallow small- or medium-sized lakes. In Steinsberger et al. (2017), a direct relationship is documented between the relative importance of reduced substances, mean hypolimnion depth, and the organic carbon content of the sediment. Deeper hypolimnias of lakes correspond to larger DO reservoirs and smaller sediment surface to water volume ratios (z). The mean hypolimnion depth of Lake Geneva is much higher than those of the lakes discussed in Müller et al. (2012a) and hypoxic conditions are found only occasionally in the deepest layers; hence, the contribution of F_red to AHM is low. The increased sediment oxic zone depth in the deeper hypolimnion of Lake Geneva confirms that organic matter mineralization is largely completed in the upper oxic zone of the lake and oxic respiration is the dominant pathway of organic matter mineralization. Consequently, F_red values are quite low.

The results of this deep lake study are in perfect agreement with observations from shallow lakes (Müller et al. 2012a) and explain how the very high AHM rate in Lake Geneva is a consequence of the large deep hypolimnetic reservoir of DO which facilitates persistently high DO concentrations that drive the mineralization of settled organic matter. As deep lakes are less likely to become anoxic above the sediment, they favor the more efficient process of oxic respiration for organic matter mineralization in the sediment.

Water column mineralization

Oxygen depletion was also quantified by the monthly DO profiles measured by CIPEL. The mean AHM calculated below a depth of 15 m (corresponding to a hypolimnion area of 534 km²; i.e., 92% of the surface area) between 1970 and 2012 was 1.34 ± 0.34 gDO m⁻² d⁻¹. With a depletion period during summer of ~180 d, this corresponds to an annual loss of 1.29 × 10⁸ t of DO per summer stratification period. Assuming a Redfield ratio of 138 moles of O₂ per 106 mineralized moles of carbon C (Redfield 1958), this relates to a mineralization rate of 370 mg C m⁻² d⁻¹. Graham et al. (2016) reported an annual gross sedimentation flux of organic matter between 285 mg C m⁻² d⁻¹ and 595 mg C m⁻² d⁻¹ for the years 2009 and 2010, which is in excellent agreement with our estimates based on the DO consumption.

While the volume-averaged AHM is in good agreement with the expectations based on the organic matter flux, it gives no information about the vertical distribution of DO depletion. TOD during summer decreased rapidly in the first 100 m, remained constant over ~150 m, and then increased again toward the maximum depth of the lake (Fig. 1). According to Livingstone and Imboden (1996), TOD can be
separated into contributions proportional to (1) the water volume and (2) the sediment surface. While the areal contribution depends on lake geometry, specifically on the sediment surface to water volume ratio (Livingstone and Imboden 1996), the volumetric contribution is independent from the morphology. The strong increase in z with depth explains the increase in TOD below 250 m. The model of Livingstone and Imboden (1996), however, assumes the volumetric and areal fraction of the TOD to be constant over the hypolimnion and, therefore, cannot explain the observed decrease in the upper layers. The trend of SOU decreasing with lake depth, as shown in Fig. 7, partially explains the decreasing TOD. However, even if the oxygen depletion caused by the sediment (zSOU) is subtracted from TOD, the remaining fraction, which can now be interpreted as water column mineralization (WCM), still shows a decreasing trend in the upper hypolimnion. As the quality of organic matter does not significantly differ between the water column and the upper sediment, it is reasonable to assume that the decrease in WCM with depth is a result of the same process causing decreasing SOU with depth, with organic matter mineralization largely occurring in the more oxic zones via oxic respiration.

Conceptual model of oxygen depletion

The observed variations in SOU and TOD with depth were used to build the foundation of a conceptual model of basin-scale DO depletion based on the deductive model of Livingstone and Imboden (1996). TOD can be separated in a water-volume-proportional term, WCM (g m\(^{-3}\) d\(^{-1}\)), and a sediment area-proportional term which consists of z, SOU, and F\(_{\text{red}}\) (g\(_{\text{DO}}\) m\(^{-2}\) d\(^{-1}\)):

\[
\text{TOD} = \text{WCM} + z(\text{SOU} + F_{\text{Red}}) \tag{3a}
\]

where z [m\(^{-1}\)] again is the sediment surface to water volume ratio (Fig. 2). In the following analysis, F\(_{\text{Red}}\) will be considered as a part of the SOU since the net effect of SOU and F\(_{\text{Red}}\) on TOD is system-analytically the same and F\(_{\text{Red}}\) was found to be negligible for most parts of Lake Geneva. Equation 3a then simplifies to

\[
\text{TOD} = \text{WCM} + z \text{SOU} \tag{3b}
\]

Assuming that the DO depletion rate is a sum of (1) an exponentially decreasing contribution caused by easily degradable organic material and (2) a constant refractory contribution of slowly degrading organic matter, SOU, and WCM can be described as:

\[
\text{SOU} = S_1 + S_2 e^{-(zS_0/Z)} \tag{4}
\]

\[
\text{WCM} = W_1 + W_2 e^{-(zW_0/Z)} \tag{5}
\]

where S\(_1\) and W\(_1\) are the contribution of the slowly degrading part of the organic matter and S\(_2\) and W\(_2\) describe the contribution of the fast degrading fraction at the lower end of the productive zone (depth z\(_0\), here set to 30 m) which decrease exponentially with depth z. The degradation depth scale z\(_s\) indicates where the contribution of the fast degrading part is reduced to 1/e of the value at z\(_0\). The degradation depth scale and is assumed to be the same for WCM and SOU.

The parameters S\(_1\), S\(_2\) and z\(_s\) were based on observational results from microprofile measurements. The best fit was found for S\(_1\) = 0.18 ± 0.06 g m\(^{-2}\) d\(^{-1}\); S\(_2\) = 1.45 ± 0.18 g m\(^{-2}\) d\(^{-1}\), and z\(_s\) = 30 ± 8 m. Estimated values of zSOU were subtracted from the mean TOD values determined by the long-term data and the result was considered to be the fraction of TOD caused by WCM. Equation 5 was then fitted to the residual, leading to values of W\(_1\) = 0.0055 ± 0.0004 g m\(^{-3}\) d\(^{-1}\); W\(_2\) = 0.0058 ± 0.0012 g m\(^{-3}\) d\(^{-1}\). In Fig. 9, the model result is shown in comparison to the values of the monitoring data.

Unlike Livingstone and Imboden (1996), this model takes the vertical variability in TOD into account. This allows for a more accurate description of the vertical structure of oxygen depletion. Rhodes et al. (2017) showed that the Livingstone and Imboden (1996) model describes the oxygen depletion in Lake Constance below 150 m well. However, our results show that the highest variability in oxygen depletion can be observed above ~ 150 m. Müller et al. (2012a) describe the AHM based on a box model which takes F\(_{\text{Red}}\) and the SOU in the upper layer of the sediment into account. Their model neglects spatial variabilities and does not consider the mineralization within the water column explicitly. Hence, it
reproduces the AHM of shallow eutrophic lakes very well, but fails to describe the processes in deep lakes. A comparison of the findings of Müller et al. (2012a,b) with the results of this study shows an important difference between shallow and deep lakes: while the oxygen depletion in shallow lakes is mainly controlled by the sediments with high values of $F_{\text{red}}$, TOD of large lakes is dominated by oxic respiration. This implies that the mineralization of organic matter is mostly completed in the upper part of the sediments in deep lakes and changes in productivity might affect oxygen depletion more rapidly. Additional studies focused on the relative importance of SOU for the TOD in different lakes based on Eqs. 1–5 and the influence of changes in primary production might further enhance understanding of the development of oxygen depletion rates.

As the model relies on several fit parameters ($S_1$, $S_2$, $W_1$, $W_2$, and $z_a$), the exact quantification of an individual parameter is complex given the relatively large variations in the data. The parameters $S_1$, $S_2$, and $z_a$ were determined experimentally; thus, the estimation of these three parameters is based on relatively few measurements from the northern part of the deep basin Lake Geneva and assumes horizontal homogeneity. In particular, the high error in the parameters $S_1$ and $z_a$ may be attributed to the large uncertainty of SOU in the deeper layers of the lake, where only one measurement was performed. Additional sources of uncertainty in these parameters include possible localized influences that were outside the scope of this study, such vertical variability resulting from Rhône River influent or nutrient loading due to wastewater effluent.

Regardless of intrinsic uncertainties in the fit parameters, the model is able to sufficiently reproduce TOD and allows for the separation of the volumetric and areal contributions. The model deviates in regions with large gradients of $z_a$ namely near 50 m and at maximum depth. At these depths, the increased SOU is smoothed by the effect of turbulent vertical diffusion. In theory, Eqs. 1–5 are only valid for non-diffusive water columns. However, this is not always strictly fulfilled, especially in deep layers where turbulent diffusion and DO gradients are comparatively large.

Between 100 m and 250 m depth, the proposed model approximates TOD with a root mean square error of only 7%. Larger values are observed above and below these depths. Below 250 m, the error is mainly due to the neglect of vertical diffusion. Diffusion leads to a net flux of DO from regions with low depletion rates to regions with higher depletion rates. Since the highest depletion rates are expected in the deepest layers with enhanced values for $z_a$, neglecting diffusion leads to an underestimate of SOU in these regions (as shown in Fig. 9). Similarly, the local bathymetry with an increase in $z$ at 50 m leads to an overestimation in the model above ~100 m. Increased mixing due to gyre formation and basin-scale internal waves may also explain parts of these deviations since the resulting DO transport to the deeper layers is not considered.

According to this simplified model, $68\% \pm 6\%$ of TOD takes place in the interior of the water column while $32\% \pm 6\%$ occurs at the sediment surface. These results compare well with values found in the literature. Bouffard et al. (2013) found SOU to be responsible for $51\% - 53\%$ of the TOD for Lake Erie (maximum depth: 64 m). Beutel et al. (2007) reported $40\% - 50\%$ for the San Vicente Reservoir in California (maximum depth: 58 m). In general, the importance of SOU decreases with lake depth since smaller values of $z_a$ favor mineralization in the water column. Consequently, the AHM in more shallow lakes can be explained solely by SOU as exemplified by Müller et al. (2012a).

**Conclusions**

The different contributors to the TOD in the hypolimnion of Lake Geneva were measured and conceptually explained. For the first time, SOU was measured autonomously across the SWI via microprofiling at up to 224 m depth in lakes. In parallel, sediment cores were retrieved to measure the flux of reduced substances from deeper sediment layers into the overlying water column. The findings were complemented by more than 40 yr of historical oxygen monitoring data. We conclude as follows:

- The SOU rate decreased from $1.08 \, {g_{\text{DO}} \over \text{m}^2 \cdot \text{d}}$ at 43 m to only $0.14 \, {g_{\text{DO}} \over \text{m}^2 \cdot \text{d}}$ at 224 m depth. As SOU decreased, the thickness of the sediment oxic zone increased. This observation can be attributed to decreasing availability of labile organic carbon. Besides the implications for the oxygen budget of the entire lake, the varying SOU and the corresponding sediment oxic zone are highly relevant for the lake ecosystem as fish egg survival depends on oxic conditions in the upper sediment. Within the sediment, the flux of reduced substances ($F_{\text{red}}$) was extremely low at all eight sampling locations. This confirms that oxic respiration of organic carbon, as opposed to oxidation of reduced substances originating from anoxic mineralization, is the dominant pathway of DO consumption in deep lakes.

- The analysis of long-term monitoring data suggests that WCM shows a similar decrease with depth. A basin-scale depletion model was developed to explain the variation of the TOD with depth based on parametrizations for SOU, $F_{\text{red}}$, WCM, and lake bathymetry.

- Based on the developed basin-scale depletion model, the relative importance of SOU and WCM can be estimated. Compared to shallow lakes, the relative importance of WCM ($\sim 70\%$ of TOD) is high. This agrees with the expectation that SOU becomes less important with increasing lake depth and, in turn, a decreasing ratio of sediment surface to water volume.
• The exponential decrease of oxygen depletion in the upper hypolimnion of Lake Geneva provides evidence of the critical role of the quality of organic matter in oxygen depletion processes and carbon cycling. Detailed analyses of the fraction of labile and refractory organic matter at different lake depths could further improve understanding of oxygen depletion in deep lakes.

References


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Conflict of Interest
None declared.