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Key Points:

- A substantial fraction of sediment organic carbon (~30%) was bound to iron in two freshwater reservoirs
- Short-term periods (2–3 weeks) of hypoxia decreased iron-bound organic carbon and total organic carbon in reservoir sediments
- Multiannual periods of hypoxia increased total organic carbon in sediment, likely through decreased rates of respiration

Supporting Information:

Supporting Information may be found in the online version of this article.

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Effects of Hypoxia on Coupled Carbon and Iron Cycling Differ Between Weekly and Multiannual Timescales in Two Freshwater Reservoirs

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Abstract Freshwater lakes and reservoirs play a disproportionate role in the global organic carbon (OC) budget, as active sites for carbon processing and burial. Associations between OC and iron (Fe) are hypothesized to contribute substantially to the stabilization of OC in sediment, but the magnitude of freshwater Fe-OC complexation remains unresolved. Moreover, global declines in bottom-water oxygen concentrations have the potential to alter OC and Fe cycles in multiple ways, and the net effects of low-oxygen (hypoxic) conditions on OC and Fe are poorly characterized. Here, we measured the pool of Fe-bound OC (Fe-OC) in surficial sediments from two eutrophic reservoirs, and we paired whole-ecosystem experiments with sediment incubations to determine the effects of hypoxia on OC and Fe cycling over multiple timescales. Our experiments demonstrated that short periods (2-4 weeks) of hypoxia can increase aqueous Fe and OC concentrations while decreasing OC and Fe-OC in surficial sediment by 30%. However, exposure to seasonal hypoxia over multiple years was associated with a 57% increase in sediment OC and no change in sediment Fe-OC. These results suggest that the large sediment Fe-OC pool (~30% of sediment OC in both reservoirs) contains both oxygen-sensitive and oxygen-insensitive fractions, and over multiannual timescales OC respiration rates may play a more important role in determining the effect of hypoxia on sediment OC than Fe-OC dissociation. Consequently, we anticipate that global declines in oxygen concentrations will alter OC and Fe cycling, with the direction and magnitude of effects dependent upon the duration of hypoxia.

Plain Language Summary Freshwater lakes and reservoirs (hereafter: lakes) play a remarkably important role in the global carbon cycle, as important sites for both carbon sequestration and greenhouse gas emissions. The extent to which lakes sequester carbon versus release greenhouse gases depends upon many factors, including associations between carbon and iron that can help to preserve carbon in sediment. However, global declines in freshwater oxygen concentrations have the potential to affect these chemical complexes. Here, we added oxygen to the bottom waters of a lake to test how changes in oxygen concentration affect carbon and iron cycling. We found that over short timescales (weeks), low oxygen conditions decreased the amount of carbon in sediment by breaking apart associations between iron and carbon that help retain carbon in sediment. However, over long timescales (years), low oxygen conditions appeared to *increase* carbon burial by decreasing the rate at which carbon inputs were decomposed. These results suggest that declining oxygen concentrations in lakes around the world may have important effects on global carbon cycling, with the direction and magnitude of the impact depending on the duration of low oxygen conditions.

1. Introduction

Freshwater lakes and reservoirs are increasingly recognized as hotspots in the global carbon cycle (Bastviken et al., 2011; Battin et al., 2009; Carey, Hanson, et al., 2022; Raymond et al., 2013; Tranvik et al., 2018). Due to high organic carbon (OC) loading from the surrounding watershed, lakes and reservoirs likely bury more OC than coastal sediments each year (Dean & Gorham, 1998; Knoll et al., 2013; Mendonça et al., 2017; Pacheco et al., 2014; USGCRP, 2018). Much of this OC remains sequestered in the sediments, especially in reservoirs, which may bury organic carbon at over six times higher rates than natural lakes (per unit area; Mendonça et al., 2017). However, OC inputs can also be respired to carbon dioxide and methane, making lakes and reservoirs a source of greenhouse gas emissions equivalent to 20% of the global emissions from fossil fuels (Deemer et al., 2016; DelSontro et al., 2018). The balance between carbon burial and emission in freshwater systems is



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Writing – review & editing: Madeline E. Schreiber, B. R. Niederlehner, Arpita Das, Nicholas W. Hammond, Mary E. Lofton, Heather L. Wander, Cayelan C. Carey controlled by numerous factors, notably including associations between OC and iron (Fe). To refine global carbon budgets and manage water resources in a changing world, it is critically important to quantify the role of Fe in OC cycling in lakes and reservoirs against the backdrop of rapidly-changing environmental conditions.

Recent research suggests that associations between OC and iron (Fe) may play a critical role in OC sequestration, though the importance of Fe in freshwater OC dynamics is currently unresolved. Fe can promote OC stability through multiple mechanisms, including occlusion of OC in aggregates, which can result in physical inaccessibility to microbial degradation and subsequent burial of OC in deeper soil or sediment horizons (Kleber et al., 2015 and references therein). Consequently, protection of OC through complexation with Fe may facilitate OC sequestration over decades to millennia (Kleber et al., 2015; Lalonde et al., 2012 and references therein). To date, most research on coupled OC and Fe cycling has focused on terrestrial and marine environments, where it has been shown that Fe-OC complexation can serve as an important mechanism for stabilizing OC, the so-called "rusty carbon sink" (e.g., Barber et al., 2017; Hemingway et al., 2019; Kramer & Chadwick, 2018; Lalonde et al., 2012). In contrast to terrestrial and marine ecosystems, only a few studies have explicitly examined Fe-OC in freshwater lakes and reservoirs. Peter & Sobek, 2018 analyzed Fe-OC in surficial sediment from five boreal lakes that spanned a gradient of oxygen conditions and found that less than 11% of sediment OC was bound to Fe, in comparison with $\sim 20\%$ across a range of primarily marine sediments (Lalonde et al., 2012). Furthermore, Peter and Sobek (2018) found no association between Fe-OC content in sediment and oxygen in overlying water when comparing across lakes. However, it should be noted that the lakes in that study were particularly high in dissolved OC (DOC) concentrations (9-42 mg/L DOC), and may not be representative of all freshwater ecosystems. Bai et al. (2021) studied Fe-OC along a salinity gradient in a subtropical tidal wetland and similarly found that freshwater areas had lower levels of sediment Fe-OC (18% of sediment OC in freshwater and 29% in saltwater), but these results were attributed primarily to wetland plant characteristics, which may not be relevant in the bottom waters of lakes and reservoirs. Differences in sediment Fe-OC between freshwater and saltwater environments may be expected based on water column characteristics, as increasing ionic strength can increase aggregation and flocculation of Fe, with differential effects depending on the quantity and quality of organic matter (Beauvois et al., 2021; Heerah & Reader, 2022; Herzog et al., 2020).

Despite limited research on Fe-OC in freshwater sediments, there are multiple reasons to expect that Fe may play an important role in OC sequestration in some freshwater ecosystems. Concentrations of Fe and DOC are strongly correlated in many freshwaters (Björnerås et al., 2017; von Wachenfeldt et al., 2008; Weyhenmeyer et al., 2014), and aqueous Fe concentrations are correlated with sediment OC accumulation in boreal lakes (Einola et al., 2011). Moreover, it is well-documented that DOC can be released from lake sediments under low-oxygen conditions, and this sediment flux is often attributed to reductive dissolution of Fe (Brothers et al., 2014; Kim & Kim, 2020; Lau & del Giorgio, 2020; Peter et al., 2017; Yang et al., 2014). Still, few studies have examined whether reactions involving Fe-OC complexes are the driving force for observed correlations between dissolved Fe and OC (but see Peter & Sobek, 2018). Furthermore, it remains unknown how the Fe-OC cycling occurring on sub-annual time scales may affect OC sequestration on the multi-annual timescales relevant for global carbon budgets.

Over short timescales (days to weeks), Fe-bound OC (Fe-OC) complexes are sensitive to the redox conditions of the surrounding environment (Figure 1). Fe-OC complexes form under oxic conditions (Riedel et al., 2013), as Fe(III) is more effective at complexing with organic matter than Fe(II) (Nierop et al., 2002). Under hypoxic conditions (low oxygen, defined here as <2 mg/L following, e.g., Yang et al., 2014), OC can be released from Fe-OC complexes through Fe(III) reduction and dissolution (e.g., Pan et al., 2016; Patzner et al., 2020; Skoog & Arias-Esquivel, 2009), which can either result directly from hypoxia or through resultant increases in pH that promote OC release (Kirk, 2004; Thompson et al., 2006). Given these conflicting patterns—i.e., that Fe-OC complexes can be preserved over decades to millennia and yet may be unstable under the reducing conditions which commonly occur on day to month timescales in aquatic sediments—it remains unclear how changing oxygen dynamics will affect coupled OC and Fe cycling in freshwater ecosystems.

Currently, the duration of bottom-water hypoxia is increasing in many lakes and reservoirs around the world (Bartosiewicz et al., 2019; Jane et al., 2021; Jenny et al., 2016; Williamson et al., 2015), which could have varying consequences for OC sequestration (Figure 1). In many dimictic lakes and reservoirs, bottom-water hypoxia is interrupted by oxic conditions during spring mixing and fall turnover, resulting in dynamic oxygen conditions on the week to month scale. Combined, these short-term patterns sum to determine the net role of lakes and reservoirs in the global carbon cycle over multiannual timescales. Periods of hypoxia have the potential to decrease OC





Figure 1. Conceptual diagram describing the hypothesized effects of changing oxygen conditions on coupled iron (Fe) and organic carbon (OC) interactions in a lake or reservoir. Under oxic conditions (top), complexation of Fe and OC (both through coprecipitation and adsorption) leads to increased concentrations of Fe-OC in sediments (increased Fe-OC protection), though oxic conditions may also lead to increased OC respiration rates. Under hypoxic conditions, reductive dissolution of Fe(III) in Fe-OC complexes increases dissolved concentrations of Fe(II) and OC in the water column while decreasing the amount of Fe-OC in sediment (decreasing Fe-OC protection), though hypoxia may also decrease OC respiration rates. The net effect of these processes on OC sequestration remains unknown, motivating this study. This figure is a simplification of complex interactions happening on a whole-ecosystem scale, and focuses on hypothesized dominant processes operating on the timescale of days to years.

sequestration through reductive dissolution of Fe(III) in Fe-OC complexes (Chen et al., 2020; Huang et al., 2021; Patzner et al., 2020). However, hypoxia also has the potential to increase OC sequestration by decreasing the rate of OC respiration (Carey, Hanson, et al., 2022; Carey et al., 2018; Hargrave, 1969; Peter et al., 2017; Sobek et al., 2009; Walker & Snodgrass, 1986), particularly if Fe-OC complexes are resistant to, or protected from, changes in oxygen concentrations in overlying water. Decreased OC respiration rates under hypoxic conditions is thought to occur primarily because respiration is less thermodynamically favorable in the absence of oxygen (e.g., Arndt et al., 2013; LaRowe & Van Cappellen, 2011). Because reductive dissolution of Fe(III) in Fe-OC complexes and decreased OC respiration under hypoxic conditions would have divergent effects on total OC sequestration, understanding the relative importance of these two processes across multiple timescales is critical for predicting the effect of hypoxia on OC sequestration in the bottom waters of lakes and reservoirs (Figure 1).

Analyzing the complex effects of oxygen on coupled OC and Fe cycling requires multiple experimental approaches. Field surveys have been effective at identifying correlations between OC and Fe (Björnerås et al., 2017; von Wachenfeldt et al., 2008; Weyhenmeyer et al., 2014). However, these observational approaches have limited capacity for identifying causal relationships. Whole-ecosystem experiments may be highly effective at identifying



real-world impacts of freshwater oxygen on Fe and OC dynamics, while allowing for important ecosystem-scale processes such as turbulence and external loading (Carpenter, 1996; Dzialowski et al., 2014; Schindler, 1998). However, high levels of variability on a whole-ecosystem scale may limit the detection of subtle changes in OC and Fe processing. Small-scale incubations may be particularly useful for identifying changes that result from hypoxia (i.e., increased DOC and Fe release from sediment, decreased levels of Fe-OC, changes in sediment OC). However, small-scale incubations are limited by fouling and changes in microbial communities, among other microcosm effects, and do not reflect the full suite of processes that interact to control OC and Fe cycling in lakes and reservoirs. Consequently, integrating multiple approaches can provide complementary information on Fe-OC dynamics across spatial and temporal scales and overcome the limitations of single-approach studies.

To analyze how hypoxia impacts OC and Fe cycling over multiple scales, this study paired whole-ecosystem oxygen manipulations with laboratory incubations. We had two objectives: (a) characterize Fe-OC (operationally defined as dithionite-extractable OC) levels in sediment of two iron-rich reservoirs, and (b) analyze how hypoxia affects coupled OC and Fe cycling over both short-term (2–4 weeks) and multiannual timescales. Through this work, we aimed to provide insight on how increasing prevalence and duration of hypoxia in lakes and reservoirs may affect the critical role of these ecosystems in the global carbon cycle.

2. Methods

2.1. Study Sites: Falling Creek and Beaverdam Reservoirs

Falling Creek Reservoir (FCR; 37.30°N, 79.84°W) and Beaverdam Reservoir (BVR; 37.31°N, 79.81°W) are small (FCR: 0.12 km², 9.3 m deep; BVR: 0.39 km², 11 m deep), eutrophic drinking water reservoirs located in southwestern Virginia, USA (Hounshell et al., 2021; Figure 2). Both reservoirs are located in deciduous forested catchments and both are dimictic, with summer stratified periods that typically last from May to October. BVR is located 3 km upstream of FCR and serves as the primary inflow source for FCR. Fe levels are high in surface water and groundwater from this region as a result of weathering and erosion of Fe-rich metamorphic rocks (53,466 mg/kg dry weight in sediment; Chapman et al., 2013; Krueger et al., 2020; Woodward, 1932). The bedrock underlying both reservoirs is layered pyroxene granulite (Virginia Division of Mineral Resources, 2003), and the primary soil series in the reservoir catchments is Edneytown Loam (Soil Survey Staff, 2022). The Köppen-Geiger climate designation of this region is Cfa: humid subtropical climate (Kottek et al., 2006). Both reservoirs have been owned and operated for drinking water provision by the Western Virginia Water Authority (WVWA) since their construction (FCR: 1898, BVR: 1872; Gerling et al., 2016; Hamre et al., 2018).

A suite of variables are routinely sampled in FCR and BVR as part of a long-term monitoring program; all data analyzed in this manuscript are available in the Environmental Data Initiative (EDI) repository with detailed metadata (Carey, Lewis, McClure, et al., 2022; Carey, Wander, Howard, et al., 2022; Carey, Wander, McClure, et al., 2022; Lewis, Niederlehner, et al., 2022; Lewis, Schreiber, et al., 2022; Schreiber et al., 2022).

2.2. Whole-Ecosystem Oxygenation Experiments

In 2012, FCR was equipped with a side-stream supersaturation hypolimnetic oxygenation (HOx) system to improve water quality in the reservoir (Gerling et al., 2014). This type of HOx system functions by withdrawing water from the bottom of the reservoir, adding concentrated, pressurized oxygen gas (95% purity) to supersaturate the water with dissolved oxygen (DO), and then returning the oxygenated water at the same depth and temperature. Previous work in FCR has shown that the HOx system effectively increases DO concentrations throughout the hypolimnion without altering temperature or decreasing thermal stability (see Gerling et al., 2014). From 2013 to 2019, the HOx system in FCR was operated at variable rates, maintaining an oxygenated hypolimnion for at least part of the summer (Carey, Thomas, & Hanson, 2022). Conversely, oxygenation was reduced in 2020 and 2021, resulting in median hypolimnetic DO concentrations <1 mg/L throughout the summer stratified period. To assess the effects of multiannual changes in oxygen availability on OC and Fe-OC in sediment, we compared sediment core and sedimentation trap data from summer 2019 (which had a history of high-oxygen conditions during the preceding 6 years) to summer 2021 (which followed a summer of hypoxic conditions in 2020; Figure S1 in Supporting Information S1). Sediment data were not collected in 2020 due to the COVID-19 pandemic.

To assess how short-term changes in hypolimnetic DO concentrations impact Fe-OC on a whole-ecosystem scale, we operated the HOx in FCR on a variable schedule throughout the summer of 2019 (Carey, Thomas, &





Source: Esri, Maxar, Earthstar Geographics, and the GIS User Community

Figure 2. Falling Creek Reservoir (FCR; 37.30°N, 79.84°W) and Beaverdam Reservoir (BVR; 37.31°N, 79.81°W) are eutrophic drinking water reservoirs located in southwest Virginia, USA (data from Carey, Lewis, Howard et al. (2022), following Woelmer et al. (2022)).

Hanson, 2022). Oxygen was added in approximately 2-week intervals at a rate of 25 kg O_2 day⁻¹ to the whole hypolimnion. Between oxygenation periods, we allowed the hypolimnion to become hypoxic over periods of at least 2 weeks without oxygenation. Because hypolimnetic volume varied throughout the summer (generally decreasing throughout the summer as the thermocline deepened), the mean concentration of oxygen added to the whole hypolimnion throughout an oxygenation period in 2019 ranged from 0.80 to 0.90 mg L⁻¹ day⁻¹.

BVR does not have a HOx system and experiences seasonal hypoxia from May through November (Hounshell et al., 2021). Consequently, BVR serves as a reference ecosystem to analyze the effects of oxygenation in FCR.

2.2.1. Oxygen

We monitored DO concentrations throughout the full water column approximately two times per week in FCR and one time per week in BVR (Carey, Lewis, McClure, et al., 2022). High-resolution (~1 cm) depth profiles were taken using a conductivity, temperature, and depth profiler (CTD; Sea-Bird, Bellevue, Washington, USA) equipped with a DO sensor (SBE 43; Carey, Lewis, McClure, et al., 2022) from the reservoir's surface to the sediments. We also measured dissolved oxygen using a YSI ProODO DO probe when the CTD was not available due to maintenance (YSI Inc. Yellow Springs, Ohio, USA; Carey, Wander, McClure, et al., 2022). YSI measurements were taken at discrete 1 m depth intervals. For a comparison of YSI and CTD measurements, see Carey, Hanson, Thomas, et al. (2022).

2.2.2. Hypolimnetic Fe and DOC

We collected water samples for DOC and Fe analysis at the deepest site in each reservoir with a 4-L Van Dorn sampler (Wildlife Supply Company, Yulee, FL, USA). Samples were collected once per week at seven depths in FCR (0.1, 1.6, 3.8, 5.0, 6.2, 8.0, and 9.0 m), which corresponded to the reservoir's extraction depths, and five

depths in BVR (0.1, 3.0, 6.0, 9.0, and 11.0 m). In 2019, we conducted a limited amount of additional sampling in FCR on a second day each week, and these measurements included DOC from 0.1, 1.6, 5.0, and 9.0 m depths.

We analyzed DOC by filtering water samples through a 0.7-µm glass fiber filter into an acid-washed bottle, which was rinsed with the filtered water three times before sample collection. The filtered samples were frozen for less than 6 months before analysis on an OC analyzer (Elementar Vario TOC cube, following APHA standard method 5310B; American Public Health Association, 2018b).

We collected both total and dissolved (filtered through 0.45-µm filters) samples for Fe. Samples were preserved in the field using trace metal grade nitric acid and analyzed using ICP-MS (Thermo Electron X-Series, Waltham, MA, USA) following APHA Standard Method 3125-B (American Public Health Association, 2018a; Krueger et al., 2020; Munger et al., 2019; Schreiber et al., 2022).

2.2.3. Fe-OC in Sediments

We analyzed the concentration of Fe-OC in surficial sediments from both FCR and BVR on multiple dates throughout the summer stratified periods of 2019 and 2021. In 2019, sediment cores in FCR were collected immediately before the HOx system was turned on or off, resulting in the most oxic or hypoxic conditions during that SSS activation or deactivation interval, respectively. Sediment cores at BVR were taken once in the middle of summer and once approximately two weeks before fall turnover in 2019. In 2021, sediment core samples were taken from both reservoirs on the same dates, approximately once per month. Additional sediment core samples were collected in March 2021, when both reservoirs were unstratified and had oxic hypolimnia.

On each sampling date, we collected four replicate hypolimnetic sediment cores using a K-B gravity sediment corer (Wildlife Supply Company, Yulee, FL, USA). Cores were collected in the deepest part of each reservoir, approximately 20 m from where water samples were taken. In 2019, each core was capped and kept on ice while transported back to the lab, where the top 1 cm of sediment from each core was immediately extruded, collected, and frozen in scintillation vials for future analysis. In 2021, cores were extruded in the field, and the samples were kept on ice while being transported back to the lab.

2.2.4. Sediment Traps

To determine the amount of Fe-OC and total OC in samples of material settling from the water column (i.e., not estimate deposition rates), we deployed 19-L buckets approximately 1 m above the sediments at the deepest point of each reservoir (8 m at FCR and 10 m at BVR). These sediment traps were deployed from June to December 2021 and sampled every two weeks by slowly bringing the bucket to the surface, decanting and discarding water from the bucket, collecting up to 5 L of the remaining water and particulate matter, and transporting this material back to the lab on ice. Upon arriving at the lab, we allowed the particulates to settle for approximately 5 min before decanting and discarding as much water as possible and filling four 50-mL centrifuge tubes with the remaining material. The samples were centrifuged for 10 min at 3,100 rpm, then combined into one vial and frozen for later analysis. No sediment traps were deployed for Fe-OC analysis in 2019.

2.3. Microcosm Incubations

To isolate the effects of oxygen from other interacting factors that affect Fe and OC on a whole-ecosystem scale, we conducted 6-week microcosm incubations using hypolimnetic sediment and water from FCR. Incubations were conducted in 177-mL glass jars with two-piece gasket-sealed lids (Verones brand; Figure S2 in Supporting Information S1), after extensive pilot testing revealed that these jars were highly effective at maintaining hypoxic conditions when sealed (DO concentrations <0.5 mg/L in this experiment) and oxic conditions when uncapped. We started the experiment with 102 microcosms split evenly into oxic (uncapped) and hypoxic (capped) treatments. After two weeks (similar to the 2019 whole-ecosystem HOx manipulation), we switched the treatment of approximately half of the remaining microcosms, generating two additional oxygen regimes: hypoxic-to-oxic and oxic-to-hypoxic. Starting on week two, there were consequently a total of four oxygen regimes: hypoxic, oxic, hypoxic-to-oxic, and oxic-to-hypoxic.

To set up the experiment, we collected sediment and water from the deepest site in FCR on 30 June 2021, when the hypolimnetic DO concentrations were <0.5 mg/L. Water was collected from 9 m depth using a Van Dorn sampler, and sediment was collected from the same location using an Ekman sampler. Samples were transported



on ice back to the lab, then homogenized by stirring and shaking. We used a syringe to add the sediment slurry (20 mL) to each jar, then slowly added 150 mL of hypolimnetic water, making an effort to minimize sediment disturbance. We stored the microcosms in an unlit incubation chamber at 15°C for the duration of the experiment, which corresponded to warm, end-of-summer conditions in the hypolimnion of FCR (Carey, Lewis, McClure, et al., 2022).

2.3.1. Microcosm Sampling

Microcosms were sampled destructively for DO, total and dissolved Fe, total and dissolved OC, pH, sediment OC, and sediment Fe-OC. For the continuous oxic and hypoxic treatments, we sampled 3–6 replicates two times per week for 4 weeks (6 replicates: days 2, 6, 9, 13; 3 replicates: days 16, 20, 23). We added additional sampling for the hypoxic-to-oxic and oxic-to-hypoxic treatments: these treatments were sampled for the first three days after switching the oxygen regime (days 14, 15, 16), twice the following week (days 20, 23), and one more time a total of 4 weeks from when treatments were switched (day 34), with three replicates analyzed per sampling event. All microcosms under a hypoxic treatment were sampled in an anaerobic chamber which maintained mean ambient oxygen conditions <200 ppm (Coy Laboratory, Grass Lake, MI, USA) to reduce oxygen exposure during sampling.

To begin sampling a microcosm, DO was measured using a YSI DO probe. While measuring DO, we used the probe to gently swirl the water in the microcosm, homogenizing the water sample while minimizing sediment disturbance. Next, we used an acid-washed syringe to withdraw 30 mL of water for total OC (TOC), 13 mL for total Fe, 30 mL of water for DOC, and 13 mL for dissolved Fe analyses. DOC samples were filtered through a 0.7-µm glass fiber filter, and dissolved Fe samples were filtered through 0.45-µm filters. After taking samples for Fe and DOC, we withdrew as much water as possible without disturbing the sediment and measured pH from this sample in a separate container using an Ohaus Starter 300 pH probe (Parsippany, NJ, USA). Finally, we swirled the sediment with remaining water (approximately 1–5 mL) and poured this mixture into a 20 mL glass EPA vial, which we then froze for Fe-OC analysis. Hypoxic microcosms were stored in the anaerobic chamber for approximately 2 hours before analysis to ensure oxygen concentrations in the chamber were sufficiently low before opening the jars. Oxic microcosms were sampled immediately after removal from the incubator.

All microcosm samples were analyzed following standard methods. We stored TOC and DOC samples in bottles that had been acid-washed and rinsed three times with the sample water. All DOC and TOC samples were frozen for <6 months prior to analysis on an OC analyzer (Elementar Vario TOC cube, following Standard Method 5310B; American Public Health Association, 2018b) Fe samples were preserved using trace metal grade nitric acid and analyzed using the ferrozine method (Gibbs, 1979). We also analyzed Fe samples from days 16 and 23 using inductively coupled plasma mass spectrometry (ICP-MS). All microcosm data are published with complete metadata in the Environmental Data Initiative repository (Lewis, Niederlehner, et al., 2022).

2.4. Fe-OC Analysis

We analyzed the amount of Fe-OC in both the whole-ecosystem and microcosm sediment samples using the citrate bicarbonate dithionite (CBD) method (Figure S3 in Supporting Information S1). This method was first described for marine systems by Lalonde et al. (2012) and has since been adapted for freshwater lakes by Peter & Sobek, 2018. It is important to note that our measurement of Fe-OC as the percentage of OC that is extractable using the CBD method is an operational definition (Fisher et al., 2021). CBD extractions have documented inefficiencies when extracting crystalline hematite (Adhikari & Yang, 2015; Thompson et al., 2019) and carboxyl-rich compounds (Fisher et al., 2020). While Fe is the primary reducible metal that associates with OC, other metals, including aluminum (A1) and calcium (Ca), may also release OC during CBD extractions. However, previous work in soils found that CBD-extracted aluminum was approximately an order of magnitude lower than CBD-extracted Fe, and therefore quantitatively much less important (Sondheim & Standish, 1983). Moreover, we found that Fe was present in much (\geq 5 times) higher quantities than A1 and Ca in water samples across all of our sediment incubation treatments (Lewis, Niederlehner, et al., 2022), further justifying our use of the operational term Fe-OC. We used the CBD method to enable comparisons both between oxygen treatments and with other published work that used the same general approach (e.g., Lalonde et al., 2012; Peter & Sobek, 2018).

Following the CBD method, each sediment sample was freeze-dried and divided into three treatments: initial, reduction, and control (Figure S3 in Supporting Information S1). "Initial" samples received no treatment and



were used to measure the OC content of the sediment. "Reduction" samples were treated with a metal-complexing agent (trisodium citrate) and reducing agent (sodium dithionite) in a buffered solution (sodium bicarbonate) to measure how much Fe and OC were released as a result of Fe reduction. Control samples were used to account for the release of OC in the reduction treatment that resulted from processes other than Fe reduction. They were treated with the same buffer (sodium bicarbonate) and sodium chloride in the same ionic strength as the trisodium citrate and sodium dithionite of the reduction treatment.

For both the control and reduction treatments, we measured 100 mg of homogenized, freeze-dried sediment into 15-mL polypropylene centrifuge tubes (Falcon Blue, Corning Inc., Corning, NY, USA). We then added 6 mL of either control or reduction buffer solution (0.11 M sodium bicarbonate) to each tube. The reduction buffer contained 0.27 M trisodium citrate, while the control buffer contained 1.6 M sodium chloride. After heating samples to 80°C in an oven, 0.1 g sodium dithionite was added to the reduction samples and 0.088 g sodium chloride was added to control samples. Samples were kept at 80°C for an additional 15 min, then centrifuged for 10 min at 3100 RPM. The supernatant was discarded. This extraction process was repeated two more times for both treatments, resuspending the sediment pellet each time by vortexing with buffer solution (Peter & Sobek, 2018).

Following the extraction step, samples were rinsed three times using OC- and Fe-free artificial lake water. Artificial lake water was prepared by diluting Artificial Hard Water from Marking and Dawson (1973) to 12.5% with Type I reagent grade water. We added 3 mL of artificial lake water to each tube and resuspended the sediment pellet using a vortex. Samples were then centrifuged for 10 min at 3100 RPM, and the supernatant was discarded.

After extraction and rinsing, all sediment samples (including those in the initial treatment) were dried and acid-fumigated for 48 hr to remove remaining citrate and bicarbonate (Harris et al., 2001). Samples were then run on a CN analyzer (Elementar VarioMax, Ronkonkoma, NY, USA) to determine the amount of OC per unit mass of sediment. In these calculations, we adjusted sediment mass to account for Fe loss during control and reduction treatments (Lewis, Niederlehner, et al., 2022; Lewis, Schreiber, et al., 2022; Peter & Sobek, 2018; Text S1 in Supporting Information S1). The amount of OC removed with Fe reduction (CBD-extractable OC) was calculated as the difference between the OC content of the control and reduction samples and expressed as a percentage of the initial OC content of the sediment.

2.5. Data Analysis

All analyses were performed in R (version 4.0.3; R core team 2020) using packages tidyverse (Wickham et al., 2019), lubridate (Grolemund & Wickham, 2011), ggpubr (Kassambara, 2020), egg (Auguie, 2019), rstatix (Kassambara, 2021), akima (Akima et al., 2022), colorRamps (Keitt, 2022), rLakeAnalyzer (Winslow et al., 2019), and tseries (Trapletti et al., 2022). All novel analysis code is archived as a Zenodo repository (Lewis, 2022).

2.5.1. Sediment Fe-OC Characterization

We calculated summary statistics to describe iron-bound organic carbon and total organic carbon in surficial sediment (2019 and 2021) and settling particulate material (2021 only) across both reservoirs. We then pooled data from both reservoirs to analyze the difference between settling material and surficial sediments using Welch's *t*-tests. Because data were unavailable for settling material in 2019, the comparison of settling material to surficial sediment was limited to 2021 data only.

2.5.2. Whole-Ecosystem Experiments: Short-Term Responses

We used Welch's *t*-tests to assess whether sediment properties differed between the 2-week periods of HOx activation compared to HOx deactivation during summer 2019 in FCR. Sediment time series did not exhibit significant temporal autocorrelation, justifying this approach (Lewis, Schreiber, et al., 2022).

To qualitatively assess whether oxygenation experiments led to differences in water column chemistry, we overlayed plots of DOC and Fe from the deepest sampling depth in each reservoir with dissolved oxygen at the same depths throughout the summer stratified period of 2019.

2.5.3. Whole-Ecosystem Experiments: Interannual Differences

We assessed whether there were significant differences in sediment properties among the four reservoir-years— BVR in 2019 (hypoxic), BVR in 2021 (hypoxic), FCR in 2019 (oxic) and FCR in 2021 (hypoxic). First, we used



Levene tests to assess homogeneity of variance among reservoir-years (Table S1 in Supporting Information S1). While Fe-OC (both per unit sediment and as a percentage of sediment OC) met the ANOVA assumption of homogeneous variance, total sediment OC did not. Consequently, we used one-way ANOVAs and Tukey post hoc tests for Fe-OC metrics, but used Welch one-way ANOVAs and Games-Howell post-hoc tests, both of which account for unequal variances, for sediment OC (Tables S2 and S3 in Supporting Information S1).

2.5.4. Microcosm Incubations

We used one-way ANOVAs and Tukey post-hoc tests to assess whether sediment properties differed between microcosm treatments, after testing for homogeneity of variance using Levene tests (Table S4 in Supporting Information S1). For this analysis, we used data from days 20 and 23 (pooled together because replicates were sampled destructively), as these were the final days when data were available for all treatments.

Equilibrium speciation-solubility calculations were conducted for day 23 of the microcosm experiments using the Spece8 module of Geochemists' Workbench (GWB; Aquatic Solutions LLC, Champaign, IL, USA) and the wateq4f thermodynamic database (Ball & Nordstrom, 1991). The goal of the calculations was to assess the predicted speciation of Fe in the presence of OC under the environmental conditions of each microcosm treatment (following Oyewumi & Schreiber, 2017). Environmental conditions considered in this analysis included pH, DO, temperature, DOC, major cations (Ca, Na, K), Fe, and major anions (Cl, SO₄; bicarbonate was not measured so we calculated bicarbonate concentrations via charge balance). Data for major cations and anions were drawn from ICP-MS analyses (described in 2.3.1 Microcosm sampling). We assumed that DOC consisted primarily of humic acids (operationally defined within the wateq4f database) for the calculations.

3. Results

3.1. Fe-OC Comprised a Substantial Portion of Sediment OC and a Smaller Proportion of OC in Settling Particulate Matter

A substantial proportion of sediment OC was associated with Fe in both FCR and BVR. In FCR (averaged across 2019 and 2021), one g of surficial sediment contained a mean of 481 µmol Fe-OC (\pm 138, 1 SD), 31 \pm 8% of the total sediment OC pool (n = 30). BVR had slightly lower Fe-OC than FCR on average, and one g of surficial sediment contained a mean of 418 \pm 121 µmol Fe-OC, 24 \pm 7% of the total sediment OC pool (n = 20). Total OC comprised 9 \pm 3% of sediment mass in FCR and 10 \pm 1% of sediment mass in BVR.

Levels of Fe-OC, both as a fraction of sediment mass and as a fraction of total sediment OC, were significantly higher in sediment core samples than in settling material collected in hypolimnetic traps (Figure 3). In 2021, averaged across both reservoirs, one g of the hypolimnetic surficial sediments contained a mean of 443 \pm 133 µmol of Fe-OC (n = 28), 69% higher than settling material collected in the traps, which contained a mean of 262 \pm 143 µmol Fe-OC (n = 17; $t_{32} = -4.24$, p < 0.001; Figure 3a). A mean of 24 \pm 6% of the total sediment OC pool was bound to Fe in sediments (n = 28), while only 9 \pm 4% of sediment OC was bound to Fe in settling material (n = 17; $t_{43} = -10.44$, p < 0.001; Figure 3c). Total OC was 60% higher in settling material ($\mu = 16.5 \pm 3.3$) than in surficial sediments ($\mu = 10.3 \pm 1.6$; $t_{20} = 7.33$, p < 0.001; Figure 3b).

3.2. Short-Term Hypoxia Decreased Sediment OC and Fe-OC on a Whole-Ecosystem Scale

Intermittent activation of the HOx in FCR in 2019 was associated with substantial changes in sediment characteristics. The amount of Fe-OC per g sediment was 30% lower during hypoxic ($\mu = 394 \pm 173$, n = 9) compared to oxic ($\mu = 560 \pm 70$, n = 7) periods in 2019 ($t_{11} = -2.6$, p = 0.02; Figure 4b). Likewise, the total amount of OC (as a percentage of sediment mass) decreased by 30% during hypoxic ($\mu = 6.0 \pm 1.55$, n = 11) compared to oxic periods ($\mu = 8.5 \pm 0.8$, n = 7; $t_{15} = -4.6$, p < 0.001; Figure 4b). Fe-OC as a percentage of total sediment OC did not significantly change with variation in oxygen during these experiments (oxic: $\mu = 36.7 \pm 3.8$, n = 7; hypoxic: $\mu = 35.7 \pm 7.9$, n = 9; $t_{12} = -0.3$, p = 0.747; Figure 4c).

In the hypolimnion of FCR, total Fe concentrations tended to increase as oxygen decreased and decrease as oxygen increased (Figure 5b). Consequently, Fe concentrations were generally lower in FCR compared to the unoxygenated reference reservoir (BVR) in 2019. Trends in DOC were more variable, though DOC concentrations were typically highest when oxygen concentrations were low in FCR (Figure 5).





Figure 3. Iron-bound organic carbon (Fe-OC; a), total sediment organic carbon (b), and Fe-OC as a percentage of sediment OC (c) all differed significantly between surficial sediment and settling particulate matter. Asterisks indicate statistical significance of the difference between surficial sediment and sediment traps: *** indicates p < 0.001 (Welch's ANOVA). Note that only 2021 data are presented because settling particulate material was not collected in 2019.

3.3. Multiannual Hypoxia Was Associated With Increased Sediment OC

Activation of the oxygenation system increased summer hypolimnetic DO concentrations in FCR from 2014 through 2019, and lower oxygen addition rates allowed for primarily hypoxic conditions in 2020 and 2021 (Figure S1 in Supporting Information S1). BVR exhibited summer hypolimnetic hypoxia throughout the duration of the study (Figure S1 in Supporting Information S1).

In FCR, the amount of OC in surficial sediment increased by 57% as DO concentrations decreased from 2019 to 2021 (Figure 6b; Tables S1 and S2 in Supporting Information S1). Consequently, total OC was lower in FCR than



Figure 4. Iron-bound organic carbon (Fe-OC; a) and total organic carbon (b) in sediment were significantly higher under oxic compared to hypoxic conditions in Falling Creek Reservoir (FCR) during the summer stratified period of 2019. Fe-OC as a percentage of sediment OC did not differ significantly between hypoxic and oxic conditions (c). Here, oxic and hypoxic conditions are classified based upon mean oxygen levels during the 2 weeks preceding sampling (Figure 5, Figure S4 in Supporting Information S1). Statistical significance of differences between oxic and hypoxic periods is indicated using asterisks: * indicates p < 0.05, *** indicates p < 0.001.







Figure 5. Increased dissolved oxygen concentrations (black lines) were associated with decreased dissolved organic carbon (brown lines; a) and total iron (Fe; slate blue lines; b) in the hypolimnia of Falling Creek Reservoir (FCR; left) and Beaverdam Reservoir (BVR; right) during the summer stratified period of 2019. All data presented are from the deepest sampling depth in each reservoir (9 m in FCR and 11 m in BVR).

BVR in 2019, but not in 2021 (Figure 6b; Tables S1 and S2 in Supporting Information S1). However, the amount of Fe-OC per gram of sediment did not change (Figure 6a; Table S1). As a result, the percentage of sediment OC that was bound to Fe decreased from 2019 to 2021 in FCR (Figure 6c; Tables S1 and S3 in Supporting Information S1). None of these three sediment characteristics differed between 2019 and 2021 in BVR (Figures 6d–6f; Tables S1–S3 in Supporting Information S1).

3.4. Experimental Microcosm Incubations Revealed Rapid Effects of Hypoxia on Fe and OC

Experimental microcosm incubations successfully established four distinct oxygen regimes. DO concentrations increased rapidly when hypoxic microcosms were unsealed and decreased rapidly when microcosms were sealed (Figure 7). At the transition from hypoxic-to-oxic conditions, DO concentrations increased to approximately the same level as the continuous oxygen treatment (\sim 7 mg/L) within 1 day. At the transition from oxic-to-hypoxic conditions, DO concentrations decreased below 1 mg/L within 1 day and declined to 0 mg/L by the end of the experiment.

Changes in oxygen conditions were associated with clear but asynchronous changes in aqueous OC and Fe. As microcosms switched from hypoxic-to-oxic conditions, TOC, DOC and total Fe decreased near synchronously, while dissolved Fe decreased below detection within 1 day of oxygen exposure. At the transition from oxic-to-hypoxic conditions, DOC and TOC rapidly increased to the same level as microcosms that had experienced continuous hypoxia (~10 mg/L; Figure 7). However, concentrations of both dissolved and total Fe only began to increase after 3 weeks of hypoxia (Figure 7). Measured DOC and TOC were strongly and linearly





Figure 6. Sediment organic carbon metrics differed significantly in association with oxygen on a multiannual scale. Metrics assessed include iron-bound organic carbon (Fe-OC) (a), total sediment organic carbon (OC; b), and Fe-OC as a percentage of OC (c) in Falling Creek Reservoir (FCR) and Beaverdam Reservoir (BVR). Blue color indicates the reservoir-year with the highest mean oxygen (2019 in FCR) and letters delineate groups that are significantly different (p < 0.05; Tables S1–S3 in Supporting Information S1).

correlated, with DOC representing a mean of $96 \pm 14\%$ of TOC (Figure S5 in Supporting Information S1); thus, we focus our discussion on DOC hereafter, but the same trends apply to TOC.

At the end of the experiment, sediment OC differed significantly among treatments (one-way ANOVA: $F_{3,20} = 9.09$, p < 0.001). Sediment OC was significantly higher in microcosms that started under oxic conditions (oxic: $\mu = 4.6 \pm 0.3$, oxic-to-hypoxic: $\mu = 4.5 \pm 0.3$) than microcosms that started under hypoxic conditions (hypoxic: $\mu = 4.0 \pm 0.0$, hypoxic-to-oxic: $\mu = 4.1 \pm 0.2$; Figure 8). Fe-OC did not differ significantly between treatments as a proportion of sediment mass ($F_{3,20} = 0.51$, p = 0.683) or as a proportion of sediment OC ($F_{3,20} = 2.40$, p = 0.098).

Speciation calculations (Table S5 in Supporting Information S1) based upon observed solution chemistry (Figure S6 in Supporting Information S1) suggest that oxygen conditions had primary control over Fe speciation, with a lesser impact on Fe-OC. The experiments that were maintained under hypoxic conditions had dominant Fe species of Fe(II), FeHCO₃⁺, FeCO₃, and FeSO₄ (all of these species contained Fe in its reduced state, Fe(II)). For all of the microcosms that were exposed to oxygen at any time (hypoxic-to-oxic, oxic-to-hypoxic, oxic), the dominant Fe species were Fe(OH)₃, Fe(OH)₂⁺, Fe(OH)₄⁻, FeOH₂⁺ and FeHumate⁺ (all of these species contained Fe in the oxidized state, Fe(III)). pH remained circumneutral across all treatments (Figure S7 in Supporting Information S1). These results indicate that (a) exposure to oxic conditions at any time in the experiment shifted the dominant oxidation state to Fe(III); (b) under oxic conditions, and to a lesser extent, hypoxic conditions, Fe complexed with DOC.

4. Discussion

Results from our whole-ecosystem manipulations suggest that oxygen affects coupled OC and Fe cycling differently over short-term (weekly) compared to long-term (multiannual) timescales (Figure 9). Short periods of hypoxia decreased total OC and Fe-OC in surficial sediment and increased concentrations of DOC and Fe in overlying water, indicating that a portion of the sediment Fe-OC pool is sensitive to changes in oxygen. However, over longer timescales, low oxygen conditions in FCR from 2019 to 2021 were associated with a 57% increase in sediment OC, indicating that the effects of hypoxia on Fe-OC (i.e., dissociation of Fe-OC complexes) may be outweighed by decreases in respiration rates under hypoxic conditions. Microcosm incubations composed of



Figure 7. Varied oxygen regimes led to differences in water chemistry among experimental microcosm treatments. Metrics assessed include dissolved oxygen (DO), dissolved organic carbon (DOC), total iron (Fe), and dissolved Fe concentrations in microcosms that were sampled destructively over the course of 34 days. Vertical lines indicate when oxygen conditions were switched, creating the oxic-to-hypoxic and hypoxic-to-oxic treatments.

slightly deeper sediment layers showed no significant change in sediment Fe-OC in response to hypoxia, suggesting that buried Fe-OC may be resistant to the effects of hypoxia. Notably, Fe-OC comprised nearly one-third of surficial sediment OC in both FCR and BVR—regardless of oxygen status—which is more than previously reported for freshwater lakes (Peter & Sobek, 2018). Below, we discuss short-term (Section 4.1) and multiannual (Section 4.2) results in the context of previous work, analyze net processing rates across the sediment-water interface (Section 4.3), and discuss why Fe-OC levels may be higher in these reservoirs than other freshwater systems (Section 4.4).

4.1. Short-Term Periods of Hypoxia Lead to Release of Fe-Protected OC and Decrease Total Sediment OC

Both whole-ecosystem and microcosm experiments suggest that short-term periods (i.e., weeks) of hypoxia can dramatically alter OC and Fe cycling. Whole-ecosystem experiments revealed changes in hypolimnetic DOC and Fe, sediment OC, and sediment Fe-OC associated with water-column oxygenation (Figures 4, 5 and 9), while microcosm incubations showed clear differences in aqueous Fe, and DOC and sediment OC among treatments (Figures 7 and 8). The magnitude of these effects was substantial: on a whole-ecosystem scale, 2 weeks of hypoxic conditions decreased both sediment Fe-OC and total sediment OC by a mean of 30%. Declining Fe-OC





Figure 8. Experimental oxygen microcosm treatments altered some, but not all, sediment properties. Metrics assessed include moles of iron-bound organic carbon (Fe-OC) per unit sediment mass (a), total sediment organic carbon (b), and Fe-OC as a percentage of sediment OC (c). Letters delineate treatments that are significantly different (p < 0.05): no treatments were significantly different for Fe-OC metrics (a, c). Days 20 and 23 were chosen for statistical comparisons as the last days in the experiment when data were available from all treatments.

and total OC in sediment, as well as concomitant increases in Fe and OC in overlying water, are consistent with the expectation that hypoxia causes reductive dissolution of Fe(III) in Fe-OC complexes, releasing soluble Fe and DOC on day to week scales (e.g., Carey et al., 2018; Pan et al., 2016; Patzner et al., 2020; Peter et al., 2016; Skoog & Arias-Esquivel, 2009; Figures 1 and 9).

Our results contribute to an accumulating body of evidence that short-term fluctuations in oxygen concentrations have important effects on OC storage in soils and sediment. Previous research has shown that recently-formed



Figure 9. Experimental results suggest that the dominant process through which hypoxia affects sediment organic carbon differs between weekly and multiannual timescales. Left: in microcosm incubations, short-term (weeks) periods of hypoxia led to increased DOC and aqueous Fe, while decreasing sediment OC. On a whole-ecosystem scale, hypolimnetic Fe was closely correlated with oxygen concentrations, and short periods of hypoxia decreased both Fe-OC and OC in sediment. Consequently, Fe-OC protection appears to be a more dominant control on sediment OC sequestration than respiration on short timescales. Right: 2 years of summer hypoxia in FCR led to increased OC in sediment on a whole-ecosystem scale, suggesting that respiration may be a more dominant control on OC sequestration than protection by Fe on this timescale.



Fe-OC associations may be particularly prone to hypoxic release, and reduction of Fe(III) in Fe-OC compounds can increase OC respiration rates during hypoxic conditions (Chen et al., 2020). As a result of these and other processes, periodic fluctuations in oxygen conditions may sustain or stimulate respiration rates relative to both constant oxic and constant hypoxic conditions (Bastviken et al., 2004; Huang et al., 2021). Furthermore, while OC released from Fe is likely to be aromatic and therefore potentially resistant to respiration (e.g., Riedel et al., 2013), this OC is susceptible to photo-oxidation upon release as DOC to the water column. Here, we found substantial decreases in sediment OC and Fe-OC following 2 weeks of hypoxia, with restored OC and Fe-OC after 2 weeks of oxic conditions (Figure 4, Figure S3 in Supporting Information S1), suggesting that at least a fraction of the sediment Fe-OC pool is sensitive to short-term changes in oxygen concentrations in overlying water on a whole-ecosystem scale.

While sediment Fe-OC responded to oxygenation on a whole-ecosystem scale, Fe-OC did not vary significantly among oxygen treatments in experimental incubations, and observed changes in aqueous Fe and DOC in microcosms were asynchronous. These observations suggest that Fe may not be driving the observed oxygen-dependent changes in OC in microcosm sediments. We expect that this difference between microcosm and whole-ecosystem results may derive at least in part from the sediments we analyzed: whole-ecosystem samples were composed of the top 1 cm of sediment from sediment cores, while sediment for the experimental incubations was sampled using an Ekman grab sampler, and therefore included deeper layers of sediment. In soil, deeper horizons are thought to have more stable Fe-OC aggregates (Rumpel & Kögel-Knabner, 2011). Our results suggest that the same pattern may be true in sediments, resulting in burial of stable Fe-OC compounds in deeper sediments over time.

Formation and dissociation of Fe-OC complexes are two of many ways in which Fe and OC are impacted by hypoxia; both Fe and OC can also respond independently to changes in DO concentrations. Fe is oxidized from Fe(II) (soluble) to Fe(III) minerals (insoluble) both biotically and abiotically under oxic conditions (Kappler et al., 2021). Increased microbial biomass may be partially responsible for the increase in sediment OC under oxic conditions in experimental microcosms, as we observed the formation of orange (likely Fe-oxidizing) biofilms on top of the sediment layer in oxic microcosms (Figure S2 in Supporting Information S1). Fe reduction is also often associated with an increase in pH, which may increase the solubility of OC (Tavakkoli et al., 2015). However, pH did not differ consistently between microcosm treatments in this study and remained circumneutral on a whole-ecosystem scale (Figures S7 and S8 in Supporting Information S1). Likewise, other metals (e.g., aluminum, calcium, manganese) may release OC from sediment under hypoxic conditions, though the influence of these alternative metals on OC release is likely less quantitatively important than Fe (e.g., Wang et al., 2021). While these alternative mechanisms likely play a role in Fe and OC release, the decrease in Fe-OC and total OC following inactivation of the oxygenation system in FCR suggests that, at least for some surficial sediments, short periods (~2 weeks) of hypoxia can cause Fe-OC complexes to dissociate and decrease sediment OC burial on a whole-ecosystem scale.

4.2. Over Multiannual Timescales, OC Respiration Rates Play a Greater Role Than Fe-OC in Determining the Net Effect of Hypoxia on Sediment OC Content

Over multiannual timescales (2019–2021), exposure to seasonal hypoxia increased the amount of OC in sediments from FCR by 57% without changing the amount of Fe-OC (Figure 6). This clearly contrasts with shortterm experimental results, which showed decreased OC content following short periods of hypoxia (Section 4.3). While many factors could affect OC and Fe-OC over multiannual timescales, the fact that no comparable effects were seen in the unoxygenated reference reservoir (BVR) suggests that these changes may be attributed to changes in DO concentrations in overlying water. Over 2 years of summer hypoxia, OC levels in sediment from FCR increased to the extent that they were no longer significantly different from the hypoxic reference reservoir (Figure 6), suggesting that legacy effects of oxygen conditions on total sediment OC may diminish after a 2-year interval.

Increases in sediment OC content with increased hypoxic duration are consistent with a reduction in sediment OC respiration rates under hypoxic conditions (Carey et al., 2018; Hargrave, 1969; Walker & Snodgrass, 1986). OC respiration rates decrease under hypoxic conditions because OC must be broken down using alternative electron acceptors, which produce a lower energy yield (Bastviken et al., 2003, 2004). Previous work conducted in BVR found that CO₂ was the dominant terminal electron acceptor in the hypolimnion during hypoxic summer



conditions (producing CH_4), and CO_2 has one of the lowest energy yields of alternate electron acceptors (McClure et al., 2021). As less OC is respired in hypoxic hypolimnetic water and sediments, OC can consequently accumulate more quickly in surficial sediments. Our results suggest that over multiannual timescales, this process (decreased respiration under hypoxic conditions) outweighs the counteracting decrease in Fe protection of OC that we observed during short periods of hypoxia.

Sediment Fe-OC content (per g sediment mass) did not significantly change after 2 years of hypoxic conditions in FCR (Figure 6), indicating that at least a fraction of these compounds are able to withstand fluctuations in environmental redox conditions. Long-term stability of Fe-OC complexes can be promoted by the formation of strong chemical bonds between OC and mineral surfaces, and these bonds continue to form over time through the rearrangement of organic molecules (Kaiser et al., 2007). Likewise, associations between Fe and OC can help to maintain a pool of small (high surface area) Fe particles that are particularly likely to associate with OC, as associations with OC can inhibit the conversion of these particles into larger, more crystalline Fe forms (called Ostwald ripening; e.g., Hiemstra et al., 2019; Zhao et al., 2022). Decreased accessibility to microbial decomposition (e.g., through burial) may further increase the ability of Fe-OC compounds to persist over time (Kaiser & Guggenberger, 2003; Kleber et al., 2015). In FCR, the history of oxic conditions (2013–2019) may have contributed to the formation of particularly stable Fe-OC complexes in sediment, which were then able to withstand two summers of hypoxia.

Importantly, much of the OC that accumulated under hypoxic conditions did not end up being bound to Fe. This result may be due to Fe oxidation state, as sorptive associations between DOC and Fe in sediment are much less likely to form if Fe is in a reduced state (Fe(II); Nierop et al., 2002). Changes in oxygen conditions are also likely to affect the composition of organic matter in sediments and in the water column (e.g., Riedel et al., 2013), potentially altering the capacity to associate with Fe. Because total OC increased following hypoxia and Fe-OC did not change, Fe-OC as a percentage of sediment OC was significantly lower after 2 years of hypoxia than before this hypoxic period. Declines in the percentage of OC that is bound to Fe may have important implications for ecosystem-scale carbon cycling, as OC that is associated with Fe is comparatively more protected from respiration (e.g., Chen et al., 2018, 2020; Hemingway et al., 2019; Kleber et al., 2005). Increased stocks of OC that are not associated with Fe may increase rates of methane production and OC release from the sediment to the water column (e.g., Hounshell et al., 2021), and could increase aerobic respiration rates under subsequent oxic periods (e.g., Chen et al., 2020; Huang et al., 2021).

4.3. Substantial OC and Fe Cycling Occurs at the Sediment-Water Interface

Notably, the OC content of the top 1 cm of sediment was significantly lower than that of settling particulate material in both FCR and BVR, and nearly three times as much of this OC was bound to Fe in sediments compared to settling material (Figure 3). These results imply a substantial level of OC and Fe processing at the sediment-water interface, and they reinforce previous research in suggesting that the sediment-water interface is a hotspot for biogeochemical cycling freshwater lakes and reservoirs (e.g., Dadi et al., 2017; Hanson et al., 2015; Krueger et al., 2020).

From a mass-balance perspective, the difference in Fe-OC between settling material and surficial sediments suggests that Fe-OC complexes are either formed or preferentially preserved (compared to OC that is not associated with Fe) in sediments. Preferential preservation of Fe-OC is well-supported, as complexation with Fe has been shown to decrease OC turnover rates across multiple ecosystems (Eusterhues et al., 2014; Kaiser & Guggenberger, 2003; Kalbitz et al., 2005; Kleber et al., 2005; Lalonde et al., 2012; Mikutta & Kaiser, 2011). However, the difference in Fe-OC between settling material and surficial sediments likely also results in large part from Fe-OC complexes), as Fe(III) levels are much higher in sediments (e.g., Davison et al., 1991) and the composition of OC in sediments may be more preferable for complexation with Fe. While we did not measure OC quality in this study, we anticipate that settling material may have higher autochthonous OC levels and be more rapidly respired, while sediment OC may be enriched in allochthonous aromatic OC, which preferentially associates with Fe (Kramer et al., 2012; Luo et al., 2022; Riedel et al., 2013; Shields et al., 2016). Documenting changes in Fe-OC throughout the process of sediment diagenesis enhances our understanding of OC sequestration, as few if any previous studies have quantified the difference between Fe-OC inputs and stocks in aquatic sediments.



4.4. High Fe-OC Levels Reflect Site-Specific Characteristics

On average, nearly one-third of surficial sediment OC was bound to Fe (dithionite-extractable) across 2 years in FCR and BVR (Figure 6). This percentage is far greater than that documented by Peter and Sobek (2018), where Fe-OC comprised $\leq 11\%$ of total sediment OC across five boreal lakes. Furthermore, the levels of Fe-OC recorded here are higher than the mean of $21.5 \pm 8.6\%$ reported for a broad range of marine sediments (Lalonde et al., 2012). With few other studies analyzing Fe-OC in freshwater lakes and reservoirs to date, our analysis provides new evidence that Fe-OC may play an important role in carbon sequestration in some freshwaters.

Despite having higher Fe-OC levels (as a percentage of total sediment OC) than most aquatic sediments studied to date, other sediment characteristics in FCR and BVR are within the range of those measured in other locations. FCR and BVR have much lower sediment OC content than the boreal lakes analyzed by Peter and Sobek (2018; 14%–38% of sediment mass), but higher sediment OC than the primarily marine sediments analyzed by Lalonde et al. (2012; 0%–7% of sediment mass). Fe concentrations are high in sediment from FCR and BVR, with a mean of 53,466 mg/kg dry weight (Krueger et al., 2020). However, Peter and Sobek (2018) observed low Fe-OC as a percentage of sediment OC ($\mu = 6.7\%$) in one extremely high-Fe lake (Övre Skärsjön; 226,172 mg/kg reducible Fe in sediment). Likewise, pH in FCR and BVR is circumneutral (Figure S8 in Supporting Information S1), well within the range of 5.4–7.6 reported by Peter and Sobek (2018), and both Peter and Sobek (2018) and Lalonde et al. (2012) included a range of oxic and hypoxic sediments in their analyses.

Differences in the percentage of organic matter that is bound to Fe may result from numerous site-specific factors. For example, increasing ratios of Fe:OC and increasing absolute concentrations of Fe and OC can all increase the amount of Fe-OC coprecipitation (Chen et al., 2016; Kleber et al., 2015 and references therein). Likewise, differing Fe forms, Fe-weathering rates, OC quality, and pH may also impact the formation and stability of Fe-OC complexes (Curti et al., 2021; Kaiser & Guggenberger, 2003; Kaiser et al., 2007; Kleber et al., 2015; Luo et al., 2022); these differences may derive from contrasting geology, catchment vegetation, and trophic status, among many other factors. Disentangling the multiple interacting factors that can influence Fe-OC dynamics across sites will require Fe-OC characterization at a greater number and diversity of lakes and reservoirs. Such research will be essential to understanding how freshwater OC sequestration may be affected by global changes in Fe concentrations (Weyhenmeyer et al., 2014), water temperatures (Dokulil et al., 2021; O'Reilly et al., 2015), and pH (Garmo et al., 2014; Gavin et al., 2018; Stoddard et al., 1999), among other factors.

5. Conclusions

Results from this study help reconcile previous Fe-OC research and shed light on how declining oxygen concentrations may impact the role of lakes and reservoirs in the global carbon cycle. Research across terrestrial soils and marine sediments has provided contradictory evidence that Fe-OC complexes are (a) readily dissociated under hypoxic conditions and (b) capable of promoting sediment OC burial into deeper (hypoxic) layers over the course of decades to millennia. Here, we find that the timescale of analysis plays a critical role in understanding the net effect of hypoxia on sediment OC and Fe-OC. Specifically, a portion of the Fe-OC pool in surficial sediment is highly responsive to hypoxia in overlying water on a weekly timescale, resulting in decreased sediment OC. However, over longer timescales, the decrease in OC that results from dissociation of Fe-OC complexes is outweighed by the increase in sediment OC that results from slower respiration rates under hypoxia, and Fe-OC complexes buried in deeper sediment layers may be particularly resistant to hypoxic release. At both timescales, our results reinforce that Fe may serve as one of several important controls over OC cycling and sediment preservation of OC in freshwater ecosystems. As the duration of hypoxia increases in lakes and reservoirs, our results suggest that OC dynamics will respond non-linearly. While short periods of hypoxia may decrease OC burial, increasing prevalence and duration of hypoxia over multiannual timescales has the potential to increase OC burial in freshwater sediment, intensifying the role of freshwaters as sinks in the global carbon cycle.

Data Availability Statement

All data used in this study are available in the Environmental Data Initiative (Carey, Lewis, Howard, et al., 2022; Carey, Lewis, McClure, et al., 2022; Carey, Wander, Howard, et al., 2022; Carey, Wander, McClure, et al., 2022; Lewis, Niederlehner, et al., 2022; Lewis, Schreiber, et al., 2022; Schreiber et al., 2022). Code to reproduce results in this manuscript is available in a Zenodo repository (Lewis, 2022).



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