Revisiting the applicability and constraints of molybdenum- and uranium-based paleo redox proxies: comparing two contrasting sill fjords

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Abstract. Sedimentary molybdenum (Mo) and uranium (U) enrichments are often used as redox proxies to reconstruct bottom water redox changes. However, these redox proxies may not be equally reliable across a range of coastal settings due to varying depositional environments. Fjords vary greatly in their depositional conditions, due to their unique bathymetry and hydrography, and are highly vulnerable to anthropogenic and climatic pressures. Currently, it is unknown to what extent Mo and U sequestration is affected by variable depositional conditions in fjords. Here, we use pore water and sequential extraction data to investigate Mo and U enrichment pathways in sediments of two sill fjords on the Swedish west coast with contrasting depositional environments and bottom water redox conditions. Our data suggest that sedimentary authigenic Mo and U pools differ between the two fjords. At the (ir)regularly dysoxic (oxygen = 0.2–2 mL L\(^{-1}\)) Gullmar Fjord, authigenic Mo largely binds to manganese (Mn) oxides and to a lesser extent to iron (Fe) oxides; Mo sulfides do not play a major role due to low sulfate reduction rates, which limits the rate of Mo burial. Authigenic U largely resides in carbonates. At the (ir)regularly euxinic (oxygen = 0 mL L\(^{-1}\); total hydrogen sulfide $\geq$ 0 mL L\(^{-1}\)) Koljö Fjord, authigenic Mo is significantly higher due to binding with more refractory organic matter complexes and Mo-Fe-sulfide phases. Uranium is moderately enriched and largely bound to organic matter.

We found no direct evidence for temporal changes in bottom water redox conditions reflected in Mo and U enrichments at either Gullmar Fjord or Koljö Fjord. While sulfidic bottom waters favor Mo sequestration at Koljö Fjord, enrichment maxima reflect a combination of depositional conditions rather than short-term low-oxygen events. Our data demonstrate that secondary pre- and post-depositional factors control Mo and U sequestration in fjords to such an extent that bottom water redox conditions are either not being systematically recorded or overprinted. This explains the large variability in trace metal enrichments observed in fjords and has implications for applying Mo and U as proxies for environmental redox reconstructions in such systems.

1 Introduction

Sedimentary molybdenum (Mo) and uranium (U) enrichments are frequently used as (paleo) redox proxies to reconstruct changes in bottom water oxygen (O\(_2\)) due to their redox-sensitive geochemical behavior (Algeo and Lyons, 2006; Jokinen et al., 2020b; Bennett and Canfield, 2020). However, the reliability of these redox proxies may be biased by inadequate understanding of Mo and U enrichment pathways and secondary depositional environmental factors (Bennett and Canfield, 2020; Jokinen et al., 2020b; Paul et
al., 2023). Besides bottom water redox conditions, secondary factors such as “the basin reservoir effect” and equilibrium with FeMoS$\sum$ (Algeo and Lyons, 2006; Helz, 2021), particulate iron (Fe) and manganese (Mn) (oxy)(hydr)oxide “shuttling” (Fe and Mn oxide shuttling hereafter; Crusius et al., 1996; Algeo and Tribovilllard, 2009), reoxygennation events (Zheng et al., 2002a, b; Morford et al., 2009), the depth and intensity of the sulfate–methane transition zone (SMTZ) in the sediment (Jokinen et al., 2020b), sedimentation rate (Algeo and Maynard, 2004; Liu and Algeo, 2020), and local detrital background (Van der Weijden, 2002; Brumsack, 2006) may considerably control authigenic Mo and U sequestration in modern coastal sediments (Jokinen et al., 2020b; Bennett and Canfield, 2020; Paul et al., 2023).

Bottom water deoxygenation is expanding in coastal areas globally due to rising anthropogenic and climatic pressures (Breitburg et al., 2018; Conley et al., 2011; Meier et al., 2022). The severity of deoxygenation varies in response to individual properties of coastal depositional environments, such as water mass restriction, temperature- and salinity-induced density gradients, productivity, and sedimentation rate. Standard thresholds to designate the degree of deoxygenation are dysoxic (O$_2$ $<$ 2 mL L$^{-1}$), suboxic (O$_2$ $<$ 0 mL L$^{-1}$), and euxinic (O$_2$ = 0 mL L$^{-1}$; total hydrogen sulfide, $\sum$H$_2$S $>$ 0 mL L$^{-1}$) after Algeo and Li (2020) and references therein.

Fjords are particularly sensitive to anthropogenically and climatically induced environmental changes, due to their unique morphological, hydrological, and sedimentological characteristics (Howe et al., 2010; Bianchi et al., 2020 and references therein). Fjords are formed by glacial erosion from Late Cenozoic ice sheets in mid-to-high latitudes. Typically, fjords are long, narrow, deep, and steep-sided U-shaped estuaries that often have one or more sills (Pickard and Stanton, 1980; Syvitski and Shaw, 1995). Such sill fjords often experience a strongly limited water mass exchange between the deep basin(s) and the coastal ocean, resulting in episodic to permanent bottom water deoxygenation. Deoxygenation in fjords is further aggravated by anthropogenic climate change (e.g., ocean warming) and high riverine and coastal runoff of nutrients and organic matter (OM), which all lead to strong vertical water mass stratification, increasing eutrophication and primary productivity, and a higher O$_2$ demand than O$_2$ supply to the bottom water upon aerobic degradation of OM (Aksnes et al., 2019; Boone et al., 2018; Darelus, 2020). Yet, high sediment accumulation and organic carbon (C$_{org}$) burial rates (e.g., Bianchi et al., 2020; Smith et al., 2015) make fjords effective spatial and high-temporal-resolution sedimentary archives of past environmental changes (e.g., Nordberg et al., 2001; Harland et al., 2004; Howe et al., 2010; Asteman et al., 2018). Thereby, fjords are ideal to test and apply trace metal proxies for investigating deoxygenation (Russell and Morford, 2001; Goldberg et al., 2012; Brinkmann et al., 2023b).

Recent research suggests that Mo and U enrichment factors (EFs) in fjord sediments show a large range (Fig. 1c and d) and less accurately record bottom water redox changes compared to less dynamic restricted basin sediments, for which Mo and U EFs have mostly been applied (Paul et al., 2023). That study showed that such limitations can be partially explained by particulate Fe and Mn oxide shuttling and pore water chemistry. However, neither the mechanisms nor extent to which Fe and Mn oxide shuttling and pore water chemistry control Mo and U sequestration in fjord sediments is fully understood. This has implications for interpreting, for example, trace metal redox proxy data derived from fjords compared to other non-fjord depositional environments.

Here, we investigate Mo and U sequestration pathways in two sill fjords with contrasting bottom water redox conditions and depositional environments using pore water and sequential extraction data of Mo, U, Mn, Fe, calcium (Ca), aluminum (Al), and sulfide, complemented with historical data of O$_2$ conditions and their relationship to climatic indices. We assess the applicability and constraints of Mo- and U-based redox proxies for reconstructing deoxygenation in fjord-type systems. Using these data, we aim to explain the observed wide ranges in Mo and U enrichments in fjord settings (Paul et al., 2023). Our study demonstrates that improving the understanding of sedimentary Mo and U redox dynamics in different coastal settings is essential for a more reliable application of Mo- and U-based redox proxies for environmental reconstructions in fjord-type systems.

2 Materials and methods

2.1 Study area and bathymetrical characteristics

Gullmar Fjord and Koljö Fjord are two adjacent sill fjords on the Swedish west coast (Fig. 1a). Gullmar Fjord (Swedish: Gullmarsfjorden) is 29 km long and 1–3 km wide and has a maximum depth of 120 m (Alsbäck Deep; Lindahl and Hernroth, 1988). Koljö Fjord belongs to an open-ended fjord system surrounding the Orust and Tjörn islands (Björk et al., 2000; McQuoid and Nordberg, 2003; Fig. 1a). Compared to Gullmar Fjord, Koljö Fjord is shallower (maximum depth of 56 m) and more restricted: it has three shallow sills to the adjacent Havsten Fjord (S1) at 12 m water depth, Skagerrak (S2) at 8 m water depth (Nordberg et al., 2001), and Gullmar Fjord (S3) at < 5 m depth (Filipsson and Nordberg, 2004a).

2.2 Oxygenation history and deoxygenation drivers

Over the past century, deoxygenation has been frequently recorded in both fjords. However, the severity and duration of deoxygenation differs between the two fjords, as evident from trace metal proxy data (Paul et al., 2023). These show lower sedimentary Mo and U enrichments in Gullmar Fjord (Fig. 1c and d, light green colored violins) – indicating less reducing conditions – compared to Koljö Fjord with higher
sedimentary Mo and U enrichments (Fig. 1c and d, pink colored violins) – indicating more reducing conditions. Several factors controlling oxygenation in both fjords have been proposed, including limited water mass exchange related to bathymetrical characteristics (e.g., presence of sills and narrow and deep basins), human activities (e.g., eutrophication), and natural variability (e.g., atmospheric drivers). These will be discussed in the following sections.

2.2.1 Hydrographic characteristics controlling water mass renewal and deoxygenation

At Gullmar Fjord, fresh water enters from the Örekilsälven river, and marine water enters from the Kattegat and Skagerrak to which Gullmar Fjord opens across a sill at 42 m water depth (Harland et al., 2006, Fig. 1a). The mixed inflow of fresh and marine water results in a strong and persistent thermohaline stratification. Brackish surface water (24–27) and almost fully marine salinities in the deep water (34–35; Nordberg et al., 2001; Arneborg, 2004) are separated by a variable pycnocline between 15–20 m below sea surface (Svensson, 1984). Deep waters are renewed and reoxygenated annually (usually in late Northern Hemisphere winter or spring; Nordberg et al., 2000). Strength and duration of bottom water renewal and reoxygenation at Gullmar Fjord are driven by the variability in the predominant wind direction and forcing, partially controlled by the North Atlantic Oscillation (NAO: normalized pressure differences between the Azores High and the Icelandic Low; Hurrell, 1995; Chen and Hellstrom, 1999; Björk and Nordberg, 2003). When prevailing northeasterly-to-easterly winds are dominant over Scandinavia, the fjord is more oxygenated due to upwelling of highly saline Skagerrak deep waters along the Swedish coast, which ventilate the fjord (Harland et al., 2006). Conversely, when westerly winds are dominant, the fjord is less oxygenated due to a downwelling regime along the coastline. Annual bottom water exchange at Gullmar Fjord via the 42 m deep sill between Lysekil and Fiskebäckskil (Fig. 1a) leads to strong seasonal variability in bottom water O2. Despite seasonally low O2 levels and episodic dyoxia, ΣH2S has never been detected in the fjord during the past 70 years (Fig. 2, upper O2 panel; Filipsson and Nordberg, 2004b).

The hydrography of Koljö Fjord (and the other fjords in the Orust and Tjörn island system, Fig. 1a) is dominated by brackish Kattegat–Skagerrak surface water originating from the Baltic Sea, while freshwater input is of minor importance as no major river discharges into the fjord (Björk et al., 2000; Filipsson et al., 2005). Mixing of the brackish surface waters (15–27) and more saline deep water (27–30), as well as deep water renewals, are less frequent and effective at Koljö Fjord compared to Gullmar Fjord. A strong pycnocline (between 15–25 m water depth) and shallow sill depths prevent direct inflow of deep waters from Skagerrak (Gustafsson and Nordberg, 1999; Filipsson and Nordberg, 2004a). Instead, deep waters enter via the adjacent Havsten Fjord, across a deeper sill between Havsten Fjord and Skagerrak at 20 m depth, before they reach Koljö Fjord (Harland et al., 2004). Deep waters at Koljö Fjord are renewed but not necessarily reoxygenated with a variable frequency (from annual to several years; Gustafsson and Nordberg, 1999). This has led to intermittently euxinic water masses between 15–20 m, typically during fall and winter, at least since the 1960s (Fig. 2, lower O2/ΣH2S panel) – potentially already longer, although no monitoring data are available before this date (Rosenberg, 1990). The strength and occurrence of low O2 conditions appear to be inverted between both fjords, despite their close proximity and connection to each other (sill S3, Fig. 1a). Shallow sill depths and dense saline deep waters prevent inflow of saline and O2-rich deep water into Koljö Fjord when environmental conditions favor reoxygenation at Gullmar Fjord (cold winters and warm summers). Only when bottom water salinity has decreased to a certain level (e.g., <28.5), below which thermohaline stratification is weakened, is ventilation re-enabled (typically for mild, humid winters with limited ice cover and cold summers; Björk and Nordberg, 2003; Nordberg et al., 2001). At both fjords, natural processes, such as deep water exchange related to the fjord’s bathymetry and hydrography, and weather variability (wind strength and direction) play a large role in controlling environmental changes and deoxygenation (Nordberg et al., 2001; Filipsson and Nordberg, 2004b).

2.2.2 Sampling and sample processing

Sediment cores were collected aboard R/V Skagerrak during September 2018. Prior to core recovery, dissolved O2, temperature, and salinity profiles of the water column were recorded by CTD (conductivity–temperature–depth; Brinkmann et al., 2022). From each site, GF-117 (115 m water depth) and KF-43 (41.5 m water depth) (see Fig. 1), one set of duplicate cores were taken using a GEMAX™ twin-barrel short gravity corer (modified Gemini corer, 9 cm internal diameter, core length 20–60 cm, from Oy Kart Ab, Finland). The duplicate cores were sampled for bottom water and pore water, as well as solid-phase geochemical analyses. Two series of bottom water and pore water were collected using Rhizon™ samplers (pore size 0.12–0.18 μm) at 2 cm vertical resolution. The first series was collected for elemental analyses and the second series for pore water ΣH2S analysis. The samples were collected into 10 mL polyethylene syringes through predrilled holes (diameter 4 mm, e.g., Jokinen et al., 2020a) immediately after core retrieval. The syringes for the ΣH2S analysis were prefilled with 1 mL of 10% zinc acetate solution to precipitate the ΣH2S as zinc sulfide (ZnS; Jilbert et al., 2018). Samples for elemental analyses were transferred into 15 mL polypropylene centrifuge tubes, acidified with 1 M HNO3, and stored in the dark at 4 °C until further analysis.
Cores for solid-phase analyses were sliced at 0.4–2.0 cm intervals on deck immediately after core retrieval. Each sediment slice was transferred to a plastic bag, which was submerged into water to remove the remaining air, sealed, and then transferred into gastight glass jars. To prevent oxidation artifacts (Kraal et al., 2009), the jars were flushed with nitrogen (N₂) and stored in a dark environment at −20 °C until subsampling for the sequential extraction procedure. Subsampling of wet sediment samples was conducted under strict oxygen-free conditions inside a N₂-flushed glove bag.

Each subsample was refrozen for 24 h at −20 °C and subsequently freeze-dried under vacuum for 48 h prior to sequential extraction. Freeze-dried samples (instead of wet samples) were chosen for simultaneous and accurate determination of water and salt contents, porosity, and trace metal speciation on the same sample.

For estimating the water [g] and salt contents [g], as well as porosity [cm³ cm⁻³], each sample was pulverized and homogenized, and samples were weighed in between each step using the bottom water salinity and the assumed solid-phase
2.3 Pore water analyses

2.3.1 Major and trace elemental concentrations

The acidified bottom water and pore water samples from both fjords were analyzed for Mo and U by inductively coupled plasma mass spectrometry (ICP-MS, Thermo Scientific XSeries 2, Department of Earth Sciences, Utrecht University) and for Al, Mn, Fe, and S by inductively coupled plasma optical emission spectrometry (ICP-OES, Thermo Scientific iCAP 6000, Faculty of Forestry and Agriculture, University of Helsinki). Dissolved Fe and Mn are considered to be present as Fe$^{2+}$ and Mn$^{2+}$, although some Mn$^{3+}$ (Madison et al., 2013) or colloidal and nanoparticulate Fe and Mn might also be present (Boyd and Ellwood, 2010; Raiswell and Canfield, 2012). Due to acidification of the pore water samples causing the release of $\Sigma$H$_2$S, dissolved S is assumed to be present primarily as sulfate (SO$_4^{2-}$; Jilbert and Slomp, 2013). Pore water $\Sigma$H$_2$S contents were analyzed spectrophotometrically (670 nm). This method is based on the dissolution of the ZnS precipitate and subsequent quantitative complexation of S as methylene blue (Jilbert et al., 2018). Measurements were calibrated with a series of standard solutions of hydrated sodium thiosulfate (Na$_2$S$_2$O$_3$ × 5H$_2$O). Subsequently, the stock solution of Na$_2$S$_2$O$_3$ × 5H$_2$O was back titrated to determine the exact concentration of S in the solution (Burton et al., 2008).

2.3.2 Diffusive flux calculations

The diffusive fluxes of Mo and U ($F_{\text{Diff}}$) were determined using Fick’s first law of diffusion (Eq. 1, Boudreau, 1997):

$$ F_{\text{Diff}} = -\varphi(0) D_S \frac{\partial c}{\partial x} \tag{1} $$

where $\varphi(0)$ is the porosity of the surface sediment, $D_S$ is the molecular diffusion coefficient near the sediment water interface (SWI), $\partial c/\partial x$ denotes the concentration gradient between the bottom water and the uppermost pore water sample. $D_S$ was determined from the seawater diffusion coefficient $D_{SW}$ (Eq. 2). Values for $D_{SW}$ for Mo and U were obtained from Li and Gregory (1974), following Morford et al. (2009), who assumed the diffusion coefficient of U to approximate that of Mo rather than the value for the UO$_2^{2+}$ complex. Based on the Stokes–Einstein relationship, $D_{SW}$ was corrected for ambient temperature, salinity, and pressure using an extended version of the `diffcoeff` function (Sulugambari et al., 2017) in the R package `marelac` (v. 2.1.10) (Soetaert et al., 2010). Pore water salinity and temperature were assumed to equal the deepest bottom water value determined by CTD. Subsequently, $D_S$ was corrected for tortuosity (Eq. 2; Boudreau, 1997).

$$ D_S = \frac{D_{SW}}{1 - \ln(\varphi^2)} \tag{2} $$

2.4 Solid-phase analyses

2.4.1 Sequential trace metal extraction

Aliquots of $\sim$100 mg of freeze-dried sediment were used for solid-phase fractionation using a combination of different extraction methods (Table 1), based closely on Jokinen et al. (2020a). We acknowledge the long-standing debate about the validity of using freeze-dried vs. wet sediments for sequential extraction of trace metals (e.g., Kersten and Förster, 1986; Hjorth, 2004). However, Jokinen et al. (2020a, b) observed no evidence for remobilization of highly redox-sensitive elements such as arsenic (As), as discussed by Huang et al. (2015), and concluded that their Mo and U data were reliable. By following the same sample handling measures as Jokinen et al. (2020a, b), we consider the potential for significant introduction of artifacts due to freeze-drying to be low. The Al, Ca, Fe, Mn, Mo, S, and U contents were fractionated in the following pools: F1 – weakly
sorbed metal (Me) species; F2 – carbonates, acid volatile sulfur (AVS), Mn(II) phosphates, and labile Me-OM complexes; F3 – Fe (oxy)(hydr)oxides, Mn (oxy)(hydr)oxides, and labile Me-OM complexes; F4 – refractory Me-OM complexes; F5 – pyrite; and F6 – silicates.

All solutions and reagents used in the sequential extraction were ultrapure for trace metal analysis and prepared with Milli-Q water to avoid contamination. Since the mineral phases extracted in F1–F3 are very redox-sensitive, solvents used in these steps were purged with N$_2$ for 30 min prior to extraction (the acetic acid and sodium citrate solution for F3 was only purged prior to adding sodium dithionate; otherwise, purging would lead to loss of volatile S compounds, which are required to reductively dissolve metal oxides), and solvents were added to the samples under constant N$_2$ gas flow (F1–F3).

The Al, Ca, Fe, Mn, Mo, S, and U contents in the first five fractions were determined using ICP-MS (Agilent 7800 ICP-MS) at Hellabs (University of Helsinki). Blank corrections were applied to all extracts to correct for background contamination (see Table S1 for details on the blank corrections). Additionally, Ca and S contents in F1 were determined in F2, since data from almost all samples fell below the detection limit as determined from a 10 mg of the residual samples was microwave-digested at 200 °C for 20 min (Rock method) on a CEM Mars 6 at Hellabs (University of Helsinki) using 5 mL 65 %–70 % HNO$_3$, 3 mL 34 %–37 % HCl$_3$, and 3 mL 48 % HF. After cooling at room temperature, the vessels were opened, and 30 mL of 4 % boric acid was added to each vessel for HF neutralization. Afterwards, the vessels were closed tightly, placed into the microwave, and neutralized at 170 °C for 15 min (boric acid HF neutralization method). The sample digests were then analyzed for Al, Ca, Fe, Mn, Mo, S, and U concentrations by ICP-MS (triple quadrupole QQQ ICP-MS) at Hellabs (University of Helsinki). Total contents of Al, Ca, Fe, Mn, Mo, S, and U were determined by the sum of all six fractions assuming that 100 % is extracted.

2.4.2 Carbon and nitrogen contents

Aliquots of ~0.25 g of freeze-dried sediment were decalcified using two wash steps of 1 M HCl as described in Van Santvoort et al. (2002). After drying and re-powdering, the decalcified samples were analyzed for organic carbon ($C_{\text{org}}$) and total nitrogen contents on a LECO 2000 CNS analyzer (Ecosystems and Environment Research Programme, Helsinki University). The results were normalized against the international analytical standard sulfamethazine. The certified value for sulfamethazine is 51.8 wt % for C and 20.1 wt % for N. The obtained mean values for the analyses were 51.5 wt % and 20.3 wt % with a standard deviation of 0.4 wt % and 0.2 wt %, respectively. Average analytical uncertainty (relative standard deviation, RSD %) based on sediment sample duplicates ($n = 3$) was < 4 wt % for both C and N. For determination of the $C_{\text{org}}$ content in each sample, measured C and N contents were corrected for weight loss upon decalcification and salt content.

### 2.4.3 Organic matter source determination

Fjord systems receive OM loading from terrestrial organic carbon ($OC_{\text{terr}}$) sources (e.g., plant material, soil, and weathering of bedrock) and marine biogenic organic carbon ($OC_{\text{phyt}}$) sources (phytoplankton production, either autochthonous or allochthonous) (Smith et al., 2015; Prebble et al., 2018). A widely used tool to quantify the sources of OM in the aquatic environment is the C/N (or N/C) ratio (e.g., Thornton and Mcmanus, 1994; Wehrmann et al., 2014; Faust and Knies, 2019). The contribution of $OC_{\text{phyt}}$ (Eq. 3) and $OC_{\text{terr}}$ (Eq. 4) to the total OM loading in estuary-type depositional environments can be approximately quantified using simple two-endmember mixing models, based on the molar N/C ratios of bulk OM (Goñi et al., 2003). Here, we use endmember values of ($N/C_{\text{terr}} = 0.04$ (terrestrial-C$_3$-plant-derived) and ($N/C_{\text{phyt}} = 0.13$ (riverine–estuarine phytoplankton), as per Jilbert et al. (2018).

\[
\%OC_{\text{phyt}} = \frac{(N/C_{\text{sample}} - N/C_{\text{terr}})}{(N/C_{\text{phyt}} - N/C_{\text{terr}})} \times 100, \quad (3)
\]

\[
\%OC_{\text{terr}} = 100 - \%OC_{\text{phyt}}. \quad (4)
\]

We acknowledge that these endmember values approximate a large potential range of N/C values for both phytoplankton-derived and terrestrial OM. In this study, we report the results of the calculation including ranges as given for the maximum and minimum endmember combinations shown in the fields of Goñi et al. (2003) (i.e., ($N/C_{\text{terr}} = 0.02–0.05$ and ($N/C_{\text{phyt}} = 0.13–0.17$).

#### 2.4.4 Calculation of authigenic Mo and U accumulation rates

To determine the authigenic Mo and U accumulation rates ($TM_{\text{MAR}}$; Eq. 7), first the mass accumulation rate (MAR; Eq. 5) and then the authigenic trace metal concentrations of Mo and U were calculated, here expressed as the excess trace metal concentration ($TM_{XS}$; Eq. 6).

\[
\text{MAR (g cm}^{-2}\text{ yr}^{-1}) = \text{SAR \times \rho \times (1 - \varphi),} \quad (5)
\]

where SAR is the mean sediment accumulation rate (cm yr$^{-1}$), $\rho$ is the dry bulk density (2.65 g cm$^{-3}$) of sediments, and $\varphi$ is the mean sediment porosity (cm$^3$ cm$^{-3}$) at
Table 1. Sequential extraction procedure for trace metals. Adapted from Jokinen et al. (2020a); an additional nitric acid step was added, designated as F5, in order to extract pyrite as described in Claff et al. (2010).

<table>
<thead>
<tr>
<th>Code</th>
<th>Fraction</th>
<th>Solvent</th>
<th>Time</th>
<th>Targeted minerals phase</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>Exchangeable</td>
<td>MgCl₂ (1 M), pH8(^a)</td>
<td>0.5 h</td>
<td>Weakly sorbed Me species</td>
<td>Tessier et al. (1979)</td>
</tr>
<tr>
<td>F2</td>
<td>Acid-soluble</td>
<td>Sodium acetate (1 M), pH 4.5</td>
<td>6 h</td>
<td>Carbonates, Iron monosulfide (FeS), Mn(II) phosphates(^b), Labile Me-OM complexes</td>
<td>Tessier et al. (1979); Cornwell and Morse (1987); Lenstra et al. (2021a); Jilbert et al. (2018)</td>
</tr>
<tr>
<td>F3</td>
<td>Reducible</td>
<td>Sodium dithionite (5 %), Acetic acid (0.35 M), Sodium citrate (0.2 M), pH 4.8</td>
<td>4 h</td>
<td>Labile and crystalline Fe (oxo)hydr/oixides (i.e., ferrihydrite and lepidocrocite)</td>
<td>Poulton and Canfield (2005); Hermans et al. (2019b); Lalonde et al. (2012)</td>
</tr>
<tr>
<td>F4</td>
<td>Organic</td>
<td>Ashing at 550 °C HCl (1 mM), pH 0</td>
<td>2 h</td>
<td>Refractory Me-OM complexes</td>
<td>Ruttenberg (1992)</td>
</tr>
<tr>
<td>F5</td>
<td>Strong-acid-soluble</td>
<td>HNO₃ (65 %–70 %) Milli-Q wash step(^c)</td>
<td>2 h/0.5 h</td>
<td>Pyrite (FeS₂)</td>
<td>Claff et al. (2010)</td>
</tr>
<tr>
<td>F6</td>
<td>Residual</td>
<td>HNO₃ (65 %–70 %) + HCl (34 %–37 %) HF (48 %), Boric acid (4 %)</td>
<td>20 min until 24 h/20 min/RT/15 min</td>
<td>Silicates</td>
<td>This study</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Sum of all phases</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) pH adjusted using Mg(OH)₂. \(^b\) Assumed based on the extraction protocol from Lenstra et al. (2021a), who used ascorbic acid to extract Mn(II) phosphates. \(^c\) An additional wash step was introduced here to remove residual concentrated nitric acid from the sample tubes. \(^d\) RT = room temperature.

each site.

\[ \text{TM}_{XS} = \text{TM}_{sample} - (\text{TM/Al}_{\text{standard}}) \times \text{Al}_{\text{sample}}, \quad (6) \]

where \( \text{TM}_{sample} \) is the trace metal concentration in the sediment sample, \( \text{Al}_{\text{sample}} \) is the Al concentration in the sediment sample, and \( \text{TM/Al}_{\text{standard}} \) is the ratio between the trace metal and Al in a standard, typically upper continental crust (UCC) values (Rudnick and Gao, 2014). Finally, Mo and U accumulation rates (\( \text{TM}_{\text{MAR}}, \mu \text{mol TM}_{\text{auth}} \text{m}^{-2} \text{yr}^{-1} \)) were estimated as

\[ \text{TM}_{\text{MAR}} = \text{MAR} \times \text{TM}_{XS} \times 10000, \quad (7) \]

where MAR is the mass accumulation rate (Eq. 5), \( \text{TM}_{XS} \) (\( \mu \text{mol g}^{-1} \)) is the authigenic trace metal enrichment in each core in (Eq. 6), and 10,000 is the conversion factor from \( \text{cm}^{-2} \) to \( \text{m}^{-2} \). A small set of Mo\(_{XS} \) values (5 out of 30) in the middle section of the GF-117 core (10.625–23.75 cm) were negative, which were omitted from the MAR estimation.

3 Age vs. depth model

Age models for sediments at both coring locations are available from previous sampling campaigns conducted between 1996 and 2001, using \(^{210}\text{Pb}\) dating and applying the constant rate of supply (CRS) model (Nordberg et al., 2000, 2001; Asteman et al., 2018), as well as biostratigraphy (Filipsson and Nordberg, 2004b). According to these studies, sedimentation rates for Gullmar Fjord are more difficult to estimate than for Koljö Fjord due to possible bioturbation artifacts. At Gullmar Fjord average sedimentation rates have been estimated to fluctuate between \(~0.70\) and \(~0.90\) cm yr\(^{-1}\). To account for compaction and bioturbation, previous studies have assumed \(~0.90\) cm yr\(^{-1}\) for the upper 15 cm and \(~0.70\) cm yr\(^{-1}\) for the remainder of the sediment core at Gullmar Fjord (Filipsson and Nordberg, 2004b; Asteman et al., 2018). At Koljö Fjord, a previous study assumed a sedimentation rate of \(~0.40\) cm yr\(^{-1}\) for the upper 25 cm and \(~0.24\) cm yr\(^{-1}\) for the remainder of the sediment core (Nordberg et al., 2001). Sediment \( \text{C}_{\text{org}} \) profiles at our sampling location in Koljö Fjord are comparable between sampling campaigns over recent decades and show distinct fluctuations around a value of \(~6\) wt \%, related to salinity variations (> or < 28–29, respectively) and the presence of laminae or lack thereof (Nordberg et al., 2001). Filipsson and Nordberg (2004a) used these relationships to establish an age model for sediment profiles from different Koljö Fjord sampling sites.
For developing our own age models, we assume similar sedimentation rates at each site to those reported previously, as a starting point. In the case of Koljö Fjord, we further refine the sedimentation-rate-based age model using the Corg-based age model of the sediment core collected in 1998, K6A (same location as KF-43). Tuning of Corg content versus sediment depth was performed in the time series tuning and analysis program QAnalyseries 1.5.1 Win (Kotov and Paelike, 2018) – a development of AnalySeries 1.1.0 (Paillard et al., 1996). For dated Corg sediment profiles of K6A and KF-43, we refer to the Supplement (Fig. S2a). Since no ages were available from the K6A sediment core for the top 0.5–8.5 cm (8.5 cm ≈ 1998 in KF-43), the assumed average sedimentation rate of ~0.40 cm yr⁻¹ for the upper 25 cm was used construct the age model for this interval. According to our sedimentation-rate-based age model for Gullmar Fjord, the sediment core from site GF-117 covers approximately the last 80 years (0–59 cm, Fig. S2b). Using the combined sedimentation-rate- and Corg-based age model for Koljö Fjord sediment core KF-43, we estimate an approximate time coverage of the last 160 years (0–49 cm, Fig. S2b).

4 Results

4.1 Pore water geochemistry

At the (ir)regularly dysoxic Gullmar Fjord, SO₄²⁻ remains relatively constant throughout the sediment core (~22–24 mM) (close to the modern global seawater value of ~28 mM), while ΣH₂S remains below the detection limit (Fig. 3a). Dissolved Mn rapidly increases below the SWI up to 288 µM at 8.5 cm depth, followed by a gradual decrease down to ~110 µM at depth and simultaneous moderate release of dissolved Fe²⁺+, reaching a maximum of ~26 µM at 16.5 cm depth. Dissolved Mo (Mo₅₉) primarily follows Mn²⁺ with depth, and once Fe²⁺ release commences, Mo follows Fe²⁺+ with depth. Local maximum Mo concentrations (230 and ~240 nM) are found at 8.5 and 14.5 cm, respectively – coinciding with Mn²⁺ and Fe²⁺+ peaks. At depth, Mo gradually decreases, except for a third smaller peak, which is also visible in Fe²⁺+ at 34.5 cm depth. This specific pattern is also visible in the dissolved U (U₅₉) profile; the remaining profile, however, shows a somewhat diverging trend relative to Mo. Besides an initial decrease below the SWI, the U maxima in the upper sediment column show a slight offset to Mo maxima at 6.5 cm (~19 nM) and 12.5 cm (~14.5 nM), respectively. Furthermore, while Mo decreases in the deeper sediment, U increases below its minimum at 20.5 cm (~5 nM) and continues to increase at depth, reaching two local maxima at 26.5 cm (~19 nM) and 36.5 cm (~23 nM), respectively. Overall, U concentration is 10-fold lower than that of Mo.

At the (ir)regularly euxinic Koljö Fjord, SO₄²⁻ gradually decreases with depth from ~20 mM down to ~13 mM, while ΣH₂S rapidly increases below the SWI to ~370 µM at 11 cm depth and remains high between ~200–400 µM throughout the sediment core (Fig. 3b). The linear gradient in the pore water shows an ΣH₂S efflux into the water column (582 µmol m⁻² d⁻¹) and an influx of SO₄²⁻ into the sediment (~18 mmol m⁻² d⁻¹). The strongly reducing character of the pore water is further illustrated by a sharp drop in Fe²⁺+ at the SWI (~2–4 µM), below which Fe²⁺+ remains extremely low with depth. Moreover, Mn²⁺ is 10-fold lower compared to Gullmar Fjord; highest concentrations are found between 2.5 and 15 cm (~12–30 µM) and below 30 cm (~14–30 µM), respectively. Molybdenum and U profiles resemble each other more closely compared to Gullmar Fjord, showing at least three distinct coinciding peaks: below the SWI, at 24 cm, and at 28 cm depth.

4.2 Solid-phase geochemistry

Solid-phase geochemistry strongly differs between the two fjords (Fig. 4). While Corg in Koljö Fjord is 2 times greater than in Gullmar Fjord, both fjords have similar molar N/C ratios of 0.08–0.11 (or as molar C/N ratio: 8.7–13.3) throughout the sediment cores. These ranges plot between phytoplankton and terrestrial derived OM (Borodvskiy, 1965; Meyers, 1994; Gohi et al., 2003; Lamb et al., 2006), as expected from fjord settings. Based on the two endmember mixing model, in both fjords the contributions of OCterr : OCphyt to the total Corg loading are ~40% : 60% with an absolute range of ±7%–24% around these values, depending on the endmembers chosen (Sect. 2.4.3). We also note the possibility of diagenetic alteration of sediment C/N ratios influencing the estimates (e.g., Van Mooy et al., 2002), although this effect likely falls within the error ranges of the endmembers.

4.2.1 Gullmar Fjord

Manganese

Total Mn contents correspond to those from previous studies on Gullmar Fjord sediments (Engström et al., 2005; Goldberg et al., 2012). Highest Mn enrichments are found in the surface sediments (14 to 399 µmol g⁻¹ ≡ 0.1 wt%–2.2 wt%), primarily in F3 as Mn (oxy)hydr)oxides (Mn oxides hereafter) and secondarily in F2 as Mn carbonates (Fig. 4a). Below 10 cm depth, Mn is almost exclusively associated with F2. Notably, there are two distinct Mn peaks at ~20 and ~40 cm depth. Fractions F1 (weakly sorbed metal species), F4 (OM complexes), F5 (pyrite bound), and F6 (residual phase, i.e., silicates) do not play a major role as host phases for Mn in this system.

Iron

While total Fe concentrations are 2 times greater than Mn (731 to 897 µmol g⁻¹ ≡ 4.1 wt%–5.0 wt%, Fig. 4a), signif-
Figure 3. (a) Gullmar Fjord and (b) Koljö Fjord downcore pore water profiles of major and trace pore water constituents: sulfate (SO$_4^{2-}$), ΣH$_2$S, Fe, Mn, Mo, and U. Estimated benthic fluxes (in nmol m$^{-2}$ d$^{-1}$) for Mo and U are provided. Positive benthic fluxes (+, upward facing arrow) refer to benthic release, and negative fluxes (−, downward facing arrow) refer to sedimentary uptake. The white filled symbols indicate the samples used for the benthic flux estimation. Bottom water values not following a linear gradient with surface sediment values were omitted from the calculation (i.e., Mo at both fjords and U at Gullmar Fjord).

Significant trends with depth in sediment are less pronounced in Fe compared to Mn. The shape of the total Fe profile is mostly impacted by F2 – Fe carbonates (i.e., siderite and ankerite) – with the co-occurrence of two peaks at approximately the same depths as the Mn peaks. Fraction 3 – Fe (oxy)(hydr)oxides (Fe oxides hereafter) – shows a modest downward decreasing trend after an ~7 cm thick subsurface peak. Other fractions are either missing (F1) or show negligible variation (F4–F6).

Molybdenum

Sedimentary Mo is strongly coupled to Mn cycling (Mn oxides) at the surface sediment in weakly sorbed metal species (F1 and F3, Fig. 4a). Except for the massive surface Mo enrichment, Mo remains below 10 nmol g$^{-1}$ on average. Remarkably, the gradual increase of F1 and F3 below 24 cm depth cannot be linked to changes in Mn content. Overall, Mo and Mn appear to be decoupled at depth, because the F2 peaks in Mn do not match the Mo profile. Fraction F2 is entirely absent, in accordance with a previous study (Jokinen et al., 2020b). We note that this result is not a consequence of analytical challenges. Although the sodium acetate matrix has a comparatively high detection limit for Mo (see Sect. 2.5.1), the value of this detection limit is equivalent to approximately 1 nmol of Mo per gram of sediment, a negligible value in comparison to the other fractions; hence, the absence of Mo in F2 is considered genuine.

Uranium

Decoupling is also apparent for Mo and U to the extent that U is sequestered in completely different phases than Mo. Uranium is largely associated with the residual (silicate) fraction F6 (~53%), followed by similar proportions of carbonates (F2) and refractory OM complexes (F4, Fig. 4a). Both F4 and F6 do not show a significant trend with depth, while F2 increases with depth (particularly below the Mn F2 peak at 20 cm depth), accompanied by F1 and F3. These trends are similar to those visible in the Mo solid-phase data.

4.2.2 Koljö Fjord

Manganese

The solid-phase Mn content at Koljö Fjord is as much as 20 times lower compared to Gullmar Fjord (9 to 25 µmol g$^{-1}$ ≡ 0.05 wt%–0.14 wt %, Fig. 4b). Moreover, the proportions between the six fractions are different: Mn is primarily enriched in F6, closely followed by F4, and total proportions of F1 and F2 make up only one-third of those of F4 and F6. All four fractions share distinct enrichment peaks, albeit none of these peaks are present in all fractions simultaneously. Strikingly, only in F4 and F6 do peaks occur
below 30 cm, whereas the other fractions show similar concentrations. All Mn peaks are also present in the solid-phase Fe extraction data at the same depth intervals (F2–F6), albeit less pronounced except for the two subsurface maxima. Manganese in F3 shows the same two peaks visible for Mn in F2, F4, and weakly in F6. However, given that F3 has the lowest total concentration of all fractions (< 1 nmol g \(^{-1}\)), it probably only plays a subordinate role for authigenic sequestration of Mn at Koljö Fjord. The same applies to F1, which is not preserved under sulfidic conditions.

Iron

Total Fe contents are slightly more elevated compared to Gullmar Fjord (607–1311 µmol g \(^{-1}\) ≡ 3.4 wt %–7.3 wt %; Fig. 4b). By contrast to Mn, Fe in F4 and F6 constitute ~60 % of the total Fe host phases. While F4 does not show any clear trend throughout the sediment core, the size of F6 shows a clear separation between strong enrichments at the top and at the bottom of the core, separated by a less enriched central part of the sediment core (~ half of the top and bottom enrichment). The third largest Fe pool consists of Fe carbonates (i.e., siderite or ankerite) or iron monosulfide (FeS) extracted in F2; however, below 30 cm, this pool disappears.

Molybdenum

Molybdenum strongly follows Mn and Fe, showing enrichment maxima at similar sediment depths, albeit Mo peaks are more pronounced (particularly between 10 and 20 cm, and at 45 cm depth, respectively, Fig. 4b) and range between ~200 and 750 nmol g \(^{-1}\). Molybdenum peaks are dominated by F1 followed by equal proportions of F3 and F4 – both likely sulfurized OM. Despite high pore water ΣH\(_2\)S, the pyrite pool F5 is unexpectedly low and of similar size as the residual fraction F6. Fraction 2 is also absent at Koljö Fjord (see Sect. 4.2, Gullmar Fjord).

Uranium

Uranium partially covaries with Mo with respect to the occurrence of peaks (Fig. 4b). However, below 25 cm U gradually increases with depth with two peaks at 33 cm and at the core bottom (~49 cm), whereas Mo drops below 30 cm and only sharply increase again from 40 to 45 cm. Besides these differences, U host phases are contrasting those of Mo and differ from Gullmar Fjord – except for F1 and F5 being the smallest fractions. Approximately 40 % of the U content is associated with F2; the remaining 60 % is divided among the others: F6, similar portions in F3 and F4, and F1 and F5 make up only 2 % and 1 %, respectively, of the total extractable U.

5 Discussion

5.1 Iron and manganese cycling and sequestration mechanisms

Sedimentary Mn and Fe geochemical cycling, contents, and speciation differ between Koljö Fjord and Gullmar Fjord (Figs. 3 and 4). These differences are likely caused by distinct depositional environmental processes within the fjords. Such processes include riverine input and salinity-driven flocculation of Mn and Fe oxides during estuarine mixing processes at the freshwater–marine interface (Sholkovitz, 1978; Brinkmann et al., 2023b), shelf-to-basin shuttling (e.g., Lenz et al., 2015a; Lenstra et al., 2020, 2021b), gravitational focusing of suspended material (sedimentary Mn oxide enrichments are usually highest in the deepest part of a basin; Hermans et al., 2019b), subsequent refluxing (reductive dissolution–oxidative precipitation) of Mn and Fe oxides between the oxic/dysoxic redox interface (within the water column or the sediment; e.g., Adelson et al., 2001; Sulu-Gambari et al., 2017), and enhanced sequestration of Fe compared to Mn under suboxic and particularly sulfidic conditions (Hermans et al., 2019b). Occurrence, strength, and interaction between such processes are sensitive to temporal variability in seasonal changes in water mass stratification, lateral transport of sediments, and redox conditions (Lenz et al., 2015a; Sulu-Gambari et al., 2017; Scholz et al., 2019).

5.1.1 Manganese

At Gullmar Fjord, sedimentary Mn mostly consists of nonsilicate Mn species – Mn oxides (F3) and Mn carbonates (F2) (Goldberg et al., 2012). Enrichment peaks of Mn are found in the surface sediments, present as Mn oxides (Fig. 4a). Oxic water conditions, which prevailed prior to our sampling campaign (Fig. 2), likely stimulated precipitation of Mn oxides from dissolved Mn\(^{2+}\) upon contact with O\(_2\), and their subsequent shuttling to the seabed (Dellwig et al., 2018; Lenstra et al., 2021b). Upon reductive dissolution of Mn oxides, Mn\(^{2+}\) is released to the pore water (Fig. 3a) and subsequently reprecipitates as Mn carbonates upon contact with bicarbonates (Calvert and Pedersen, 1996). Besides a strong subsurface maximum in Mn\(^{2+}\) at Gullmar Fjord (Fig. 3a), dissolution of Mn oxides continues (albeit slower) deeper in the sediment within the zone of Fe(III) reduction (Fig. 3a, Goldberg et al., 2012), as evident by elevated Mn\(^{2+}\) (~100 µM) at depth and the presence of two comparatively small Mn oxide peaks at ~20 and ~40 cm depth, respectively. At the same depth intervals, our data also show two Mn peaks in F2, which likely consist of Mn carbonates (Fig. S4) – although some Mn(II) phosphates might also be present (Hermans et al., 2019b, 2021). Distinct layers of authigenic Mn carbonates are a common observation in dysoxic–suboxic sediments beneath oxic bottom waters in many coastal marine environments (e.g., Huckriede and Meischner, 1996; Lenz et
Figure 4. (a) Gullmar Fjord and (b) Koljö Fjord downcore solid-phase distribution of $C_{\text{org}}$ (divided into riverine–estuarine phytoplankton-derived (blue) and terrestrial-plant-derived (yellow)) and elemental Ca, Mn, Fe, Mo, and U contents (divided into six different fractions (F1–F6) based on the sequential extraction scheme) (Table 1). In each $C_{\text{org}}$ plot, the light-shaded area (light yellow, light blue) denotes the absolute range in $OC_{\text{phy}}$ and $OC_{\text{terr}}$ fractions based on the maximum ($N_{\text{C}}^{\text{terr}}$ and $N_{\text{C}}^{\text{phyt}}$) endmembers (dashed line) and minimum ($N_{\text{C}}^{\text{terr}}$ and $N_{\text{C}}^{\text{phyt}}$) endmembers (dash-dot line), as reported in Goñi et al. (2003). The solid line denotes the $OC_{\text{terr}}$: $OC_{\text{phyt}}$ contributions calculated when using the ($N_{\text{C}}^{\text{terr}}$) = 0.04 and ($N_{\text{C}}^{\text{phyt}}$) = 0.13 endmembers, as per Jilbert et al. (2018). A summary of bottom water $O_2$ history at both fjords is provided on the right. The bottom water redox conditions were derived from the fjord’s monitoring data (Fig. 2; SMHI, 2022) and correlated to selected depth intervals in the sediment cores using the estimated age–depth models (Sect. 3, Fig. S2a, b). The abbreviations used to describe the average bottom water (BW) redox conditions are defined as follows (in order of appearance from top to bottom): dyx = dysoxic, ox = oxic, epi. = episodical, seas = seasonal, re-oxy = reoxygenation, eux = euxinic, and sub = suboxic.

al., 2015b; Lenstra et al., 2020). Formation of Mn carbonate enriched layers at Gullmar Fjord (and similar depositional environments) may either represent relic shifts in the redox boundary within the sediment (Goldberg et al., 2012) as described in Burdige (1993) or are a result of changes in Mn input related to varying intensity in Mn oxide shuffling – where stronger Mn oxide shuffling promotes Mn carbonate accumulation (e.g., Lenz et al., 2015a; Lenstra et al., 2021b).

At Koljö Fjord these processes also occur but are less efficient due to high sulfate reduction rates, inducing $\Sigma$H$_2$S release to the pore water (Fig. 3b), which in turn alters the vertical zonation of electron acceptors used for OM degradation (Burdige, 1993). Under sulfidic pore water conditions, Mn oxide formation is restricted to sufficiently oxygenated zones in the water column (Brewer and Spencer, 1971), which at Koljö Fjord occur above the pycnocline ($\sim$15–25 m water depth; SMHI, 2022). When these Mn oxides sink through the suboxic water column, they begin to reductively dissolve before reaching the sediment (Burdige, 1993; Scholz et al., 2017). Only a small fraction of Mn oxides may survive the dissolution process in the water column, which is the case here, as inferred by the lower Mn oxide peak close to the sediment surface ($\sim$1.5 cm). Subsequently, these Mn oxides are available for rapid conversion into Mn carbonates in the sediment (Lenz et al., 2015a; Lenstra et al., 2021a). Coinciding Ca and Mn peaks in F2 throughout the sediment core suggest that past fluctuation in the redox conditions have allowed for conversion of Mn oxides to Mn carbonates. However, the more reducing water column and pore water conditions prevent a long-term build-up of these two Mn host phases (Fig. 4b). This explains why the majority of sedimentary Mn resides in the silicate fraction (F6) and refractory OM complexes (F4) and the 20 times lower total Mn contents at Koljö Fjord compared to Gullmar Fjord, where more oxygenated conditions and deeper water depth promote Mn oxide formation and gravitational focusing on the surface sediments.

5.1.2 Iron

Compared to Mn, total sedimentary Fe contents at Gullmar Fjord show hardly any trend with depth (Fig. 4a), analogous to estimates by Goldberg et al. (2012). The largest Fe frac-
tion, F6, probably represents Fe bound to illite, which is the most common clay mineral in this area (Hassellöv et al., 2001), and the smallest Fe fraction, F5, is likely a residual from a previous phase rather than actual pyrite, as $\Sigma H_2S$ was below detection limit. Fractions F2 and F3 show the strongest variability among all fractions. As no separate extraction of labile Fe oxides was performed, F3 likely consists of a mixture of both labile (i.e., ferrihydrite and lepidocrocite) and crystalline (i.e., goethite and hematite) Fe oxides (Table 1). Based on experiments performed by Poulton and Canfield (2005), we cannot rule out that a minor fraction of labile Fe oxides ($\sim$ 1%–2%) may have already been extracted in the previous sodium acetate step (F2). With regards to F3, we observe a surface enrichment in the upper 10 cm, which is likely due to the formation of labile Fe oxides. This assumption is based on the Fe and Mn pore water profiles, showing that the F3 peak coincides with maximum dissolution of Mn oxides (release of $Mn^{2+}$), which itself catalyzes labile Fe oxide formation by oxidation of $Fe^{2+}$ (Figs. 3a and 4a; Wang and Van Cappellen, 1996). With onset of reductive dissolution of Fe oxides below $\sim$ 10 cm depth, these labile Fe oxides are being readily dissolved. Underpinned by the results by Goldberg et al. (2012), we therefore infer that below 10 cm F3 mostly contains crystalline and refractory Fe oxides and to a lesser extent labile Fe oxides. Two distinct Fe peaks are present in F2 that overlap with the Mn peaks at $\sim$ 20 and $\sim$ 40 cm depth. Given the apparent low sulfate reduction rates and release of dissolved $\Sigma H_2S$ required for iron monosulfide (FeS) formation (Canfield et al., 1993; Aller, 1994), we do not expect any FeS present in F2. Instead, Fe peaks likely consist of Fe carbonates, which typically coprecipitate with rhodochrosite and calcite (e.g., Wittkop et al., 2020). Correspondingly to the Mn carbonate peaks, those Fe carbonate peaks likely represent past maxima in Fe oxide accumulation in response to enhanced Mn and Fe oxide refluxing (e.g., Lohan and Bruil, 2008; Lenstra et al., 2021b).

Total Fe contents at Koljö Fjord are comparable to Gullmar Fjord (Figs. 1c, d and 4), which agrees with their geochemical redox behavior under euxinic and dysoxic conditions, respectively (Bennett and Canfield, 2020; Paul et al., 2023). Moreover, molar Mo / U bottom water and surface sediment ratios are elevated at both fjords (Fig. S4) with respect to the molar Mo / U ratio in average seawater ($\sim$ 7.5, Algeo and Tribovillard, 2009), indicating a greater mobility and subsequent sequestration of Mo compared to U (Scholz et al., 2013). Indeed, our sequential extraction data reveal greater authigenic Mo sequestration at both fjords relative to U (Figs. 1c, d and 4).

5.2 Molybdenum and uranium in Gullmar Fjord

Molybdenum

At Gullmar Fjord, the molar Mo / U ratio in the pore water is $\geq$ 7.5 and is gradually rising until 20 cm sediment depth (Fig. S4), which is consistent with the extraction data showing a dominance of Mo and Mn co-enrichments (F3) over U in the upper 10 cm. This distinct Mo-Mn covariation is likely caused by Fe and Mn oxide shuttling from the water column to the sediment surface, favored by ambient water column redox conditions. Under such conditions, Mo (as molybdate, $MoO_4^{2-}$) has a strong affinity to Mn and Fe oxides. By attaching to these oxides, Mo is removed from the water column and shuttled to the sediment surface (Berrang and Grill, 1974; Scholz et al., 2013; Dellwig et al., 2021). As $O_2$ is consumed within the upper 2–5 mm of the sediment (Brinkmann et al., 2023b) – which is typical for non-bioturbated or mildly bioturbated coastal marine sediments underlying a relatively oxygenated water column (Ghid et al., 2003, Slomp et al., 2013; Hermans et al., 2019a) – first Mn and then Fe oxides are being reductively dissolved, by which Mo is subsequently released to the pore water (Fig. 3a, e.g., Sulu-Gambari et al., 2017). A comparison between Mo accumulation rates ($Mo_{MAR}$) of the upper 10 cm and Mo diffusive...
fluxes (\(\text{Mo}_{\text{benthic flux}}\)) reveals that most of the released Mo will likely be buried in the sediment. In this low-sulfide system and thus in the absence of sulfide-mediated pathways for authigenic Mo sequestration, however, a fraction of \(\text{Mo}_{\text{diss}}\) might diffuse upwards and escape burial, indicated by a positive \(\text{Mo}_{\text{benthic flux}}\) (Table 2). This is a common mechanism observed in coastal depositional environments subject to periodic or seasonal reoxygenation events, such as the major Baltic inflows (MBIs, e.g., Scholz et al., 2013; Dellwig et al., 2021) or in response to the collapse of seasonal water mass stratification (Sulu-Gambari et al., 2017).

With the dissolution of Mn and Fe oxides and subsequent conversion of such into Fe and Mn carbonates (F2), solid-phase Mo concentrations drop to background levels (below 10 cm, Fig. 4a). This observation highlights the importance of Mn oxides as carrier phases for Mo in surface sediments underlying seasonally dysoxic bottom water (e.g., Sulu-Gambari et al., 2017) but also that more reducing conditions are required to permanently sequester Mo in the sediment (e.g., Tribovillard et al., 2006; Jokinen et al., 2020b). Below 20 cm, however, solid-phase Mo concentrations gradually increase again with depth, while pore water Mo concentrations show a decreasing trend. The solid-phase increase in Mo is mostly manifested in F1 (almost 20 times higher than the background). A similar observation was made by Jokinen et al. (2020b) in sulfidic sediments in the Finnish archipelago, underlying anoxic water column. The authors inferred that the occurrence of Mo in F1 deeper in the sediment represents a precursor phase of Mo sequestration following the iron sulfide pathway (Vorlicek et al., 2018; Helz and Vorlicek, 2019). In contrast to Jokinen et al. (2020b), the pore waters at Gullmar Fjord are depleted in \(\Sigma\text{H}_2\text{S}\) (at least down to 40 cm depth), due to removal of any produced \(\Sigma\text{H}_2\text{S}\) by reactive Fe oxides in the sediment (Goldberg et al., 2012). In turn, thiolation of molybdate is strongly limited (Helz et al., 1996; Erickson and Helz, 2000). As molybdate is the more likely pore water species of Mo throughout the core (Goldberg et al., 2012), we suspect that most Mo is attached to Fe oxides or weakly sorbed to other mineral phases.

Uranium

Uranium (as uranyl complex with mostly carbonate or phosphate; Langmuir, 1978) may be absorbed to Fe oxides and, to a lesser extent, Mn oxides in surface sediments (McKee et al., 1987; Morford et al., 2007; Brennecka et al., 2011; Singh et al., 2012; Dang et al., 2016). However, shuttling-induced removal and sediment surface focusing, as evident for Mo, is believed to be a less important control for U sequestration in marine sediments (Dellwig et al., 2021). Instead, diffusion across the SWI is regarded as the primary transport pathway of U to the sediment (Barnes and Cochran, 1991; Klinkhammer and Palmer, 1991; Algeo and Maynard, 2004; McManus et al., 2005). Our \(U_{\text{MAR}}\) and benthic flux estimates suggest that, while diffusion across the SWI appears to drive U sequestration in the surface sediments (negative flux, Table 2), the presence of an additional particulate flux is possible. This is inferred from MARs that are 5 times greater than the diffusive flux (Table 2). One possibility is the deposition of U as particulate non-lithogenic U (PNU) associated with OM (Hirose and Sugimura, 1991; Zheng et al., 2002b). However, preservation of such PNU is unlikely at Gullmar Fjord due to surface water \(O_2\) concentrations \(>200\mu\text{M}\). Alternatively, the additional particulate U flux originates from U adsorption to Fe (and Mn) oxides close to the SWI and subsequent shutting of U to the sediment surface.

According to thermodynamics, reducing conditions are required for permanent sequestration of U as particle reactive U(IV) uraninite (Veeh, 1967; Bonatti et al., 1971; Anderson et al., 1989; Klinkhammer and Palmer, 1991). Besides abiotic U(IV) reduction, U may also be reduced by biotic reactions in the presence of dissimilatory metal-reducing bacteria and/or sulfate-reducing bacteria, leading to the formation of both crystalline uraninite (e.g., Bargar et al., 2008; Sharp et al., 2009; Lee et al., 2010) and monomeric (non-crystalline) non-uraninite (e.g., Lovley et al., 1991, 1993; Fredrickson et al., 2000; Bhattacharyya et al., 2017).

In contrast to Mo, pore water U does not increase with the onset of Mn oxide dissolution but rather shows a decreasing trend (Fig. 3a). Coinciding with maximum Fe oxide dissolution (\(\text{Fe}^{2+}\) release), this decreasing trend is amplified and continues until \(\sim20\) cm, at which most \(\text{Fe}^{2+}\) has been consumed. The coinciding drawdown of pore water U with increasing \(\text{Fe}^{2+}\) is a typical observation in both experimental and field data of marine sediments (Cochran et al., 1986; Klinkhammer and Palmer, 1991; Zheng et al., 2002a) and has been attributed to a combination of Fe and U reduction commencing at similar redox potentials and efficient removal of particle reactive U(IV) from pore water (Cochran et al., 1986; McKee et al., 1987; Zheng et al., 2002a). According to U reduction kinetics, however, \(\text{Fe}^{2+}\) is unlikely to act as a direct abiotic U(IV) reductant. More likely are crystalline Fe oxides, which have shown to reduce U(VI) to U(IV) at near-neutral pH (Ginder-Vogel and Fendorf, 2008). At Gullmar Fjord, crystalline Fe oxides (F3) are observed from \(\sim10\) cm and onwards. They remain relatively constant throughout the sediment core, by which they could be involved in U(VI) reduction and U(IV) precipitation.

Our geochemical data reveal that U mostly resides in clay minerals (F6) or associated with refractory OM complexes (F4). However, U in F4 and F6 strongly covaries with Al in F4 and F6 (Fig. S6) and remains relatively constant throughout the sediment core. When U(VI) builds complexes with OM, it becomes unresponsive to microbially mediated U(VI) reduction (despite favorable reduction conditions), which decreases the pool of reactive and mobile U species (Ortiz-Bernad et al., 2004; Fuller et al., 2020). Therefore, we infer that U in F4 (OM-bound) and F6 (clay-bound) is of detrital origin and largely unreactive rather than representing reactive authigenic U phases.
Instead, possible host phases of authigenic U are expected in F2, e.g., associated with Mn carbonates (F2), which particularly below 24 cm strongly coaly with each other. Presumably, U extracted in F2 represents monomeric non-uraninite (Fu et al., 2018; Jokinen et al., 2020b). Water-column-monitoring phosphate (PO$_4^{3-}$) data indicate benthic release of PO$_4^{3-}$ during the sampling month (5 µmol L$^{-1}$ at 110 m; SMHI, 2022), suggesting high pore water PO$_4^{3-}$ concentrations. Numerous studies have observed that PO$_4^{3-}$ inhibits the formation of crystalline uraninite (Bernier-Latmani et al., 2010; Boyanov et al., 2011; Alessi et al., 2014; Morin et al., 2016), supporting the presence of non-uraninite in F2 (Jokinen et al., 2020b). Remarkably, F2 and F3 follow a very similar downcore enrichment pattern (Fig. S7). In contrast to U in F2, U in F3 likely comprises crystalline uraninite, as inferred by laboratory experiments (Fu et al., 2018, uraninite extracted with hydroxylamine). To explain this covariation and downward increase, it could be argued that U in F2 and F3 represents a continuum of U host phases extractable in both F2 and F3. Monomeric non-uraninite and crystalline uraninite are known to have a high affinity to form complexes with OM, by which OM could be a possible candidate as a common U host phase (Alessi et al., 2014; Bone et al., 2017). However, neither F4 nor C$_{org}$ shows a meaningful correlation with F2 or F3. Thus, association of U in F2 (non-uraninite) and F3 (crystalline uraninite) with OM is unlikely at this site (Figs. 4a, S7).

5.2.2 Molybdenum and uranium in Koljö Fjord

Molybdenum

In contrast to Gullmar Fjord, solid-phase Mo/U ratios at Koljö Fjord are elevated throughout the sediment core relative to average seawater, suggesting enhanced authigenic Mo, which is in line with elevated Mo-EFs above the crustal background and compared to U-EFs (Fig. 1c). Bottom water redox conditions are subject to inter–annual fluctuations, resulting in a range from oxic to sulfidic conditions (Fig. 1b). Such fluctuations favor formation of Fe and Mn oxides, which are the most efficient scavenging and transport carriers of Mo in waters with variable oxic–suboxic boundaries (Scholz et al., 2013; Bertine and Turekian, 1973; Algeo and Lyons, 2006; Wagner et al., 2017). The six-fold higher Mo accumulation rates (Mo$_{MAR}$) of the upper 10 cm compared to the Mo benthic influx support this particulate shuttling transport mechanisms for Mo at Koljö Fjord. In contrast to Gullmar Fjord, the sulfidic bottom and pore water permit efficient Mo burial and limit rerelease to the bottom water (Sulu-Gambari et al., 2017; Lenstra et al., 2019).

In the sediment, the close relationship between Mo burial and Mn and Fe oxide refluxing is demonstrated by the corresponding fluctuations in Mo with Fe, particularly Mn enrichments (Fig. 4b). However, in sulfidic pore water, Mn oxides are negligible as host phases due to reductive dissolution of such (Fig. 3b). Instead, our data suggest that Mo weakly bound to S phases (F1, i.e., thiomolybdate intermediates) may serve as transitional host phases after release of Mo from Mn oxides. The presence of these particle reactive thiomolybdate intermediates agrees with pore water $\Sigma$H$_2$S concentrations > 11 µM and required subsequent replacement of O atoms with S atoms during subsequent thiolation (MoO$_4^{2-}$ to MoS$_4^{2-}$; Helz et al., 1996; Erickson and Helz, 2000), and it explains the gradual decrease in pore water Mo / U molar ratios.

Likely candidates for more permanent host phases of Mo are F3 and F4. These comprise crystalline Fe oxide (F3), which in contrast to labile Mn oxides can survive sulfidic conditions (Hermans et al., 2021), as well as the labile and more refractory OM pool (F3 and F4). Both fractions are of similar size and generally follow the same pattern as F1, suggesting that Mo initially stored in labile thiomolybdate intermediates on mineral surfaces (e.g., Helz et al., 1996; Vorlikee et al., 2018) may also become more permanently incorporated into Fe oxides and OM (e.g., Chappaz et al., 2014; Dahl et al., 2017). Organic-bound host phases for Mo are a common finding in different marine sediments underlying suboxic–euxinic water columns (e.g., Huerta-Diaz and Morse, 1992; Algeo and Lyons, 2006; Scholz et al., 2013). Other permanent Mo host phases frequently discussed are pyrite (e.g., Huerta-Diaz and Morse, 1992; Sundby et al., 2004; Chappaz et al., 2014) and Fe-Mo-S colloids, such as FeMoS$_4$ (Vorlikee et al., 2018; Helz and Vorlikee, 2019; Helz, 2021). Both mineral phases are expected to be extracted in F5 (Table 1). While adsorption of Mo onto pyrite surfaces has been considered a possible Mo sequestration pathway (Huerta-Diaz and Morse, 1992), in practice correlations between Mo and Fe or total S contents are often very weak (Lyons et al., 2003; Algeo and Maynard, 2004; Scholz et al., 2013). Consistent with these studies, at Koljö Fjord, the correlation between F5 Mo and Fe : S or S is not significant ($R^2 = 0.02$ or $R^2 = 0.03$, respectively). Thus, we infer that pyrite is probably negligible as a Mo host phase. Although Fe-Mo-S colloids are more likely to host Mo in F5, given the small fraction size relative to the sum of all fractions (median ~ 8 wt %), the total contribution of Mo sequestered as Fe-Mo-S colloids is of minor importance.

Uranium

In contrast to Mo, benthic U fluxes are very similar between Gullmar Fjord and Koljö Fjord, illustrating that diffusion – as the key sequestration pathway of U – commences in a similar manner under either oxic–dysoxic or suboxic–sulfidic bottom waters. Similarly, estimated U$_{MAR}$ of the upper 10 cm are only slightly elevated above those at Gullmar Fjord but also about 5 times greater than the U diffusive flux (Table 2). This implies that U water column dynamics (particular transport to the sediment surface) behave relatively similarly between the two fjords. Despite a strong pycnocline and
Consequently, the results suggest that besides sulfidic crystalline uraninite U(IV), and oxidized U(VI) complexes, that at Koljö Fjord OM-bound U likely consists of a mixture of reactive (prone to remobilization) monomeric non-uraninite U(IV), unreactive (less prone to remobilization), crystalline uraninite U(IV), and oxidized U(VI) complexes.

As Koljö Fjord’s pore water becomes sulfidic just below the SWI, U\textsubscript{diss} is removed from the pore water within the upper 5 cm (Fig. 3b). Within the same depth interval, a small fraction of U is (loosely) sorbed U(VI) to mineral phases (F1, Fig. 4b). However, with onset of reducing conditions, F1 rapidly decreases and is replaced by more refractory U(IV) mineral phases extracted in F2–F4. The majority of authigenic U resides in F2, followed by F3 and F4 with similar fraction sizes (Fig. 4b). Remarkably, all three fractions follow a similar downward trend (Fig. S7). They also show a strong covariation with C\textsubscript{org}, particularly with terrestrial C\textsubscript{org} (Figs. 4b, S7). As refractory metal-OM complexes are expected to be extracted in F2–F4 (Table 1), these findings imply that, unlike at Gullmar Fjord, the majority of authigenic U at Koljö Fjord is bound to labile and refractory OM complexes or C\textsubscript{org}-coated minerals (Bone et al., 2017). Although some crystalline Fe oxides are present at Koljö Fjord (Sect. 5.1.2) to serve as potential reduction agents of U(VI), the strong inverse correlation between pore water $\sum$H\textsubscript{2}S and U\textsubscript{diss} – a phenomenon observed in the Black Sea water column (Rolison et al., 2017) – suggests that U(VI) reduction is dominantly enzymatically mediated by sulfate-reducing bacteria (Lovelvy et al., 1993; Fletcher et al., 2010).

Sulfate reduction also has implications for the type of U(IV) mineral phases precipitating from solution, since SO\textsubscript{4}\textsuperscript{2−} favors non-uraninite formation (Fletcher et al., 2010; Fuller et al., 2020). In other words, even under sulfidic conditions, U(VI) associated with OM may only be partially reduced to crystalline uraninite U(IV); therefore, the latter may not be the dominant U phase despite favorable reduction conditions (Cumberland et al., 2018; Fuller et al., 2020). This infers that at Koljö Fjord OM-bound U likely consists of a mixture of reactive (prone to remobilization) monomeric non-uraninite U(IV), unreactive (less prone to remobilization), crystalline uraninite U(IV), and oxidized U(VI) complexes (Sharp et al., 2011; Alessi et al., 2012; Jokinen et al., 2020b). Consequently, the results suggest that besides sulfidic conditions the presence of OM may increase the preservation potential of U in fjord sediments.

Despite the reduction potential of reduced S species (i.e., FeS or pyrite; Bargar et al., 2013; Cumberland et al., 2021) on U(VI), U does not form direct bonds with such species (Choppin and Jensen, 2006; Bone et al., 2017). As supported by the apparent decoupling between U in F5 (pyrite extractable) and Fe : S, Mo, Fe, and S in F5, such decoupling suggests that U neither forms Fe-S precipitates nor absorbs directly onto other metal-Fe-S phases, such as pyrite or Mo-Fe-S colloids. In contrast to the decoupling of Mo and U in F5 (and all other fractions), both trace metals show a very similar enrichment pattern in F1 (Fig. S8). As the Mo-U covariation pattern is absent at Gullmar Fjord and thiomolybdate intermediates are the most logical host phase for Mo in F1 at Koljö Fjord, this could suggest that U is sorbed to those thiomolybdates. However, the relatively consistent $\sim 2$ cm offset – particularly below 20 cm – between Mo and U in F1 makes the presence of a common host phase very unlikely (Fig. S8). Thus, the covariation pattern must have a different source. Bone et al. (2017) explained U-S correlations by associations between thiol-S species with OM to which U is absorbed. This indirect bonding of U to S species via organic coatings could indeed explain why U in F5 is decoupled from (Mo)-(Fe)-S species but present in F1.

All OM complexes are expected to be extracted in F4; thus, no OM associations are expected in F5. In accordance with Gullmar Fjord, we observe that F6 in U covaries with F6 in Al, suggesting that U is of detrital origin. Uranium(VI) may be transported to the basin sorbed to clay minerals, such as illite, which is the dominant clay mineral (grain size fraction $< 2 \mu$m) in Gullmar Fjord and Skagerrak (Hassel- löv et al., 2001). Sorption of U(VI) to clay minerals has been demonstrated in laboratory experiments (Bachmaf and Merkel, 2011; Mei et al., 2022). Strikingly, U residing in F6 is believed to represent the most refractory U host phase – crystalline uraninite (Jokinen et al., 2020b; F5 in their protocol). Our data do not support this hypothesis. This may be partially explained by the additional extraction step we conducted to separate the pyrite fraction (F5) and silicate fraction (F6).

### Table 2. Estimated authigenic Mo and U mass accumulation rates (TM\textsubscript{MAR}) of the upper 10 cm (whole core in brackets) and benthic fluxes (TM\textsubscript{benthic flux}) at Gullmar Fjord (GF-117) and Koljö Fjord (KF-43). Positive benthic fluxes (+) refer to benthic uptake, and negative fluxes (−) refer to benthic release. TM\textsubscript{MAR} values were estimated using total solid-phase Al, Mo, and U concentrations derived from the sum of F1–F6 in the sequential extraction data. All values are given in µmol m\textsuperscript{−2} yr\textsuperscript{−1}. Note that Mo\textsubscript{benthic flux} and U\textsubscript{benthic flux} provided here are the same as those provided in Fig. 2 given in nmol m\textsuperscript{2} d\textsuperscript{−1}.

<table>
<thead>
<tr>
<th>Site</th>
<th>Mo\textsubscript{MAR} (UCC)</th>
<th>Mo\textsubscript{benthic flux}</th>
<th>U\textsubscript{MAR} (UCC)</th>
<th>U\textsubscript{benthic flux}</th>
</tr>
</thead>
<tbody>
<tr>
<td>GF-117</td>
<td>87.15 (69.49)</td>
<td>+25.13</td>
<td>9.52 (20.56)</td>
<td>−2.04</td>
</tr>
<tr>
<td>KF-43</td>
<td>238.91 (332.33)</td>
<td>−39.21</td>
<td>11.39 (17.98)</td>
<td>−2.12</td>
</tr>
</tbody>
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https://doi.org/10.5194/bg-20-5003-2023
5.3 Key authigenic Fe, Mn, Mo, and U sequestration mechanisms at Gullmar and Koljö fjords

In summary, our geochemical data suggest that sedimentary authigenic Mo and U sequestration in Gullmar Fjord and Koljö Fjord is strongly controlled by differences in composition and availability of pore water and solid-phase species, particularly $\Sigma$H$_2$S and OM, as well as Fe and Mn oxide shuttling (Fig. 5). At Gullmar Fjord, sediment geochemistry is governed by particulate shuttling of Fe and particularly of Mn. Here, Fe and Mn oxides represent the key carrier and sedimentary host phases for authigenic Mo. First, Mo is sequestered by Mn oxide in the surface sediments; with progressing diageneses, the Mo host phase switches to poorly crystalline oxides and crystalline Fe oxides deeper in the sediment. While crystalline Fe oxides are involved in U(VI) reduction to U(IV), ultimately U builds its own authigenic mineral phases either associated with PO$_4^{3-}$ or carbonate. A considerable fraction of U appears to be of detrital origin, which is largely unreactive and immobile.

At Koljö Fjord, sedimentary elemental dynamics are controlled by sulfate reduction and subsequent release of $\Sigma$H$_2$S to the pore water. Given the sulfidic conditions, authigenic sequestration of Mo in Mn oxides is strongly limited; instead, Mo is buried in sulfide phases, such as thiomolybdates and Mo-Fe-S colloids. Authigenic U sequestration at Koljö Fjord is governed by a combination of pore water $\Sigma$H$_2$S and high OM contents. Here, U largely resides with OM, either as an OM complex or adsorbed to organic coatings of other mineral phases, including sulfides.

5.4 Applicability and constraints of Mo and U as paleo-environmental proxies in fjord environments

Fjords are dynamic depositional systems characterized by high sedimentation rates, $C_{\text{org}}$ loading, and sensitivity to weather and climatic changes (e.g., Howe et al., 2010; Bianchi et al., 2020), which theoretically makes them desirable for paleo-environmental reconstructions. However, Mo and U redox proxy signals may be partially or entirely overwritten by secondary depositional environmental factors (Algeo and Lyons, 2006; Scholz et al., 2018; Jokinen et al., 2020b; Paul et al., 2023).

To verify whether authigenic Mo and U enrichments in fjord sediments can be reliably used as archives for changes in bottom water $O_2$, we assess to what extent distinct features in the trace metal speciation from Gullmar Fjord and Koljö Fjord can be explained by hydrographic variability (e.g., occurrence of inflow events or seasonal water mass exchange) or are a result of secondary controls as pre-depositional (e.g., Fe and Mn oxide shuttling and water mass restriction) or post-depositional (e.g., oxidative remobilization and pore water geochemistry) factors.

5.4.1 Molybdenum

Sedimentary Mo enrichment patterns are frequently used to reconstruct temporal environmental changes in coastal marine settings related to deoxygenation or reoxygenation events, or (semi)regular variability in water column stratification and redox conditions, ranging from seasonal (e.g., Egger et al., 2016; Sulu-Gambari et al., 2017; Dellwig et al., 2021) up to centennial timescales (e.g., van Helmond et al., 2018; Scholz et al., 2018). In non-euxinic settings, the efficiency of Mo$_{\text{diss}}$ removal and subsequent Mo sequestration into surface sediments is strongly coupled to ambient Mn redox dynamics, coupled to the redox variability in the water column and bottom water. In principle, short-term redox fluctuations in the bottom water promote Mn oxide refluxing, increasing the Mo flux to the sediment and subsequent surface sediment enrichment (Lenz et al., 2015b; Sulu-Gambari et al., 2017; Scholz et al., 2017; Dellwig et al., 2018).

Between 2010–2018, Gullmar Fjord’s bottom water was largely dysoxic and only periodically interrupted by short-term oxygenation events (Figs. 2 and 4a). Likely combined with higher Mn oxide input into the fjord, these redox fluctuations promoted both Mn oxide (F3) and authigenic Mo accumulation (F3) in the subsurface (∼ upper 10 cm) sediments (Fig. 4a). As Mn oxides diagenetically convert into Mn carbonates over time (Burdige, 1993; Huckriede and Meischner, 1996), the presence of two additional Mn carbonate peaks (F2) at ∼1990–2000 (∼17–24 cm) and ∼1960–1970 (∼35–40 cm) suggest that there has been more than one period of enhanced Mn oxide refluxing in Gullmar Fjord, likely associated with oxygenated periods during gradually basin-wide deoxygenation after the 1960s (Fig. 2). One would expect that this trend towards more dysoxic conditions and enhanced Mn oxide shuttling to have resulted in enhanced authigenic Mo enrichments in the surface sediment at that time. However, no record of such a trend is visible throughout the sediment core, possibly due to poor Mo preservation potentials under the ambient low $\Sigma$H$_2$S pore water concentrations at Gullmar Fjord.

To permanently preserve fluctuations in Mo sequestration, associated with, for example, seasonal changes in water mass stratification, it requires both high Mo supply by Mn oxide shuttling and pore water $\Sigma$H$_2$S concentrations > 11 µM (Egger et al., 2016; Sulu-Gambari et al., 2017) to lock Mo in sediment, e.g., by the formation of stable Mo sulfides (Helz et al., 1996). Manganese oxide supply into Gullmar Fjord is naturally exceptionally high compared to other coastal marine systems worldwide (Burdige, 1993; Aller, 1994; Jokinen et al., 2020b; Lenstra et al., 2021b), and due to gravitational focusing of these Mn oxides at the deeper fjord (our study site; Brinkmann et al., 2023b), Mo supply is expected to have been sufficiently high throughout the past 80 years. Instead, upon reductive dissolution of Mn oxides under suboxic conditions in the surface sediments, insufficiently high pore water $\Sigma$H$_2$S fuels benthic release of Mo$_{\text{diss}}$, thereby reducing
Figure 5. Summary of key authigenic Mo (left columns) and U (right columns) sequestration mechanisms at Gullmar Fjord ((ir)regularly dysoxic) and Koljö Fjord ((ir)regularly euxinic). Mo(IV)$_\text{diss}$ denotes the dissolved phase molybdate and U(VI)$_\text{diss}$ denotes dissolved uranyl complexes with, for example, carbonate or phosphate. All sedimentary Mo and U phases are present as solids. Within each panel, left to right indicates increasing content of solid phases shown with symbols. For detailed description of sequestration mechanisms, see Sect. 5.2.

sedimentary Mo sequestration (Figs. 3a and 4a, Goldberg et al., 2012). Therefore, neither the initial signs of deoxygenation nor the temporal variability in shuttling is recorded by Mo, which impedes the applicability of Mo as a redox proxy to reconstruct environmental changes at Gullmar Fjord.

In contrast to Gullmar Fjord, pore waters at Koljö Fjord are sufficiently sulfidic to permit permanent sequestration of Mo (Fig. 4b). Five distinct enrichment peaks in Mo were found at Koljö Fjord, all of which cover a period of at least 10 years: ∼1860–1870, ∼1923–1940, ∼1944–1962, ∼1972–1994, and ∼2005–2013 (Fig. 4b). This eliminates the possibility that those Mo enrichment peaks are coupled to seasonal or episodical short-term redox fluctuations, either linked to rapid reoxygenation events as described for the Baltic Sea (Scholz et al., 2018; Dellwig et al., 2018) or seasonal variability in water column stratification and redox conditions, as described for a former estuary with comparable seasonal redox dynamics as at Koljö Fjord (Egger et al., 2016).

Instead, it is more likely that Mo enrichment peaks represent longer-term environmental changes. Although water column data show a trend towards more reducing bottom water conditions over the last century, the three Mo peaks for which monitoring data are available (∼1944–1962, ∼1972–1994, and ∼2005–2013) seem to have been deposited under different environmental conditions. Between 1944–1962, the Mo peak was formed during non-euxinic conditions, while between 1972–1994 and 2005–2013 bottom waters were largely suboxic–euxinic, interrupted by oxygenation events of variable intensity and duration (Fig. 2). Therefore, Mo sequestration has in the last century migrated from the pore water – which must have been sulfidic shortly below the sediment–water interface to prevent benthic escape of Mo$_\text{diss}$ (Erickson and Helz, 2000) – to the bottom water (and water column). Seemingly, both mechanisms result in similarly strong Mo sequestration and preservation at Koljö Fjord, which is in line with high Mo contents in coastal sediments with a shallow SMTZ, underlying an oxygenated water column (Jokinen et al., 2020b). Such substantial effects of pore water chemistry on Mo sequestration complicate the application of Mo as a single proxy to reconstruct distinct environmental changes in this (and similar) fjord system(s).

5.4.2 Uranium

Uranium is considered a more sensitive recorder of mild deoxygenation compared to Mo due to a higher reduction–oxidation potential of U relative to Mo (Lovley et al., 1991; Zheng et al., 2002a; van Helmond et al., 2018). According to this, the U record at Gullmar Fjord shows more variability than Mo, specifically in the reactive U(IV) monomeric non-uraninite pool associated with (Mn) carbonates (F2). Given the strong dependency of Mn oxide and carbonate accumulation/preservation on redox changes, U in F2 may provide some information about environmental changes. In fact, U in F2 shows more variability than Mn carbonates, implying that U might have recorded more redox changes. However, individual U peaks cover at least decade-long timescales; as the redox changes occurred on shorter (seasonal) timescales, U peaks cannot be linked to individual/seasonal oxygenation events (Figs. 2 and 4a). Either U did not record these short-term redox changes or the original redox signal has been overprinted by post-depositional redox-induced remobilization (e.g., Wang et al., 2013).

At Koljö Fjord, the timing of U minima and maxima is comparable to that of Mo, although the onset of enrichment
peaks seems to occur slightly earlier in U (Fig. S8). This suggests that U might have recorded the timing of redox variability more reliably than Mo and which – in addition to Gullmar Fjord – would support the idea of U being a more sensitive redox recorder (e.g., van Helmond et al., 2018). Overall, the three broader U peaks occurring between ~1920 and ~1980 (Fig. 4b) coincide with longer periods of oxic–dysoxic intervals intercepted by somewhat periodic suboxic–euxinic intervals (Figs. 2 and 4b). During these intervals, U in F1 shows a strong covariation with Fe in F2 (Fig. S9a) and Fe and Mn in F3 (Fig. S9b), suggesting a stronger particulate flux of U by Fe (and Mn) oxides. In contrast to Mo, the (for Koljö Fjord) unusual long oxic period between 1993 and 1997 is detectable by a clear U minimum (F1) and coinciding small Mn and Fe oxide (F3) enrichment peaks (Fig. S9b), highlighting the better preservation conditions for labile Mn and Fe oxides. Sequestration of loosely sorbed (labile) U was probably limited by higher bottom water O2, which reduced the reduction potential into more refractory phases. The most probably limited by higher bottom water O2.

5.5 Implications for using Mo- and U-based environmental (redox) proxies in fjord sediments

Our findings suggest that using Mo and U enrichments in fjord sediments as archives of environmental changes hosts certain pitfalls. While environmental changes over decadal timescales may be identifiable, higher frequency events (seasonal or episodical) are either not reliably recorded by Mo and U, due to non-steady-state depositional conditions, or overprinted by post-depositional processes. Thus, neither at Gullmar Fjord nor at Koljö Fjord is it possible to accurately reconstruct environmental (redox) changes on these timescales using Mo and U enrichments. This is in line with observations made at the Canadian fjord Saanich Inlet (Algeo and Lyons, 2006), where the occurrence of high-frequency events partially restricts the applicability of the Mo /TOC ratio (where TOC represents total organic carbon) – a widely applied measure for the degree of basin restriction (Algeo and Lyons, 2006). Unsurprisingly, in our sedimentary records from Gullmar Fjord and Koljö Fjord, significant correlations between TOC and Mo are lacking as well (Fig. S10). Therefore, when describing the degree of water mass restriction in fjord settings, it is not advisable to solely consider the Mo /TOC ratio, as this approach may result in false interpretations.

Despite these limitations, sedimentary Mo and U enrichments may still be applicable as environmental (redox) proxies when considering total Mo and U (enrichment factors), expressed as ranges over the whole sediment core (Fig. 1c and d) or as Mo-EF and U-EF cross-plots (Fig. 6, Algeo and Tribovillard, 2009). In fact, sedimentary Mo-EF and U-EF covariation patterns at Gullmar Fjord and Koljö Fjord (Fig. 6) clearly support the key sequestration mechanisms inferred by our sequential extraction data (Fig. 4). Both sites
follow the Fe and Mn oxide shuttling signature, although it is more distinct at Gullmar Fjord, given the more intense Fe and Mn oxide cycling. This is reflected by a clear division into higher Mo–EF/U–EF ratios, associated with the shuttling-induced surface maxima of Mo (0–7 cm, bold reversed triangles, Fig. 6) and lower Mo–EF/U–EF ratios in the remaining core (> 7 cm).

The central role of $\sum H_2S$ in controlling Mo sequestration and burial efficiency is demonstrated by the clear separation of Gullmar Fjord and Koljö Fjord by the $1 \times$ molar Mo : U seawater line, dividing non-sulfidic (bottom water) sites (Koverhar and Gullmar Fjord) from (ir)regularly sulfidic sites (Saanich Inlet, Cariaco Basin, Skurusundet, Black Sea, and Koljö Fjord). This distinct Mo–EF and U–EF pattern corresponds well to our geochemical data revealing that despite strong Mo shuttling by Fe/Mn oxides (i.e., at Gullmar Fjord), ultimately inadequate bottom water and pore water chemistry (here: low $\Sigma H_2S$) is the key factor explaining lower Mo–EFs compared to other (ir)regularly dysoxic sites (e.g., Lilla Värtan, Stockholm Archipelago), by limiting permanent Mo sequestration. Correspondingly, total U enrichments are highly sensitive to the presence of dissolved and solid-phase OM, carbonate, and phosphate phases. All these phases can result in precipitation of refractory and inert U mineral phases, which can help to explain both lower U–EFs under more reducing conditions and higher U–EFs under less reducing conditions. Evidently, pore water geochemistry (i.e., composition and concentration of dissolved phases) dictates permanent Mo and U sequestration in our fjord sediments, regardless of the bottom water redox condition and the impact of secondary pre-depositional factors (i.e., Fe and Mn oxide shuttling).

6 Conclusions and outlook

Our trace metal sequestration extraction-based case study highlights how environmental and post-depositional factors may obscure direct information about past redox conditions stored in sedimentary Mo and U enrichments in fjord settings. We further complement the current understanding of Mo and U sequestration in fjord sediments, albeit with considerable limitations regarding U speciation. This demonstrates the urge for future studies, ideally combining both (1) a modified sequential extraction scheme with targeted U extraction steps to differentiate between crystalline uraninite and monomeric non-uraninite species and (2) microanalytical techniques, such as synchrotron-based X-ray absorption spectroscopy (e.g., XANES, EXAFS) or nano secondary ionizing mass spectrometry (nanoSIMS), providing more information on metal speciation and host phases (e.g., oxidation states, coordination number, and isotopic composition). Despite these limitations, the key findings from this study are the following:

- Mo enrichment factors (EFs) are applicable as a comparative redox proxy for whole sediment core data in fjord settings, when considering a certain degree of diagenetic overprinting of initial redox signals.
- U enrichment factors (EFs) are applicable with caution as a comparative redox proxy for whole sediment core data in fjord settings, due to considerable diagenetic overprinting of initial redox signals.
- Permanent U sequestration is more complex than Mo; even in less reducing sediments, more refractory mineral phases may be formed and vice versa strongly reducing sediment may not guarantee stronger U enrichments.
- Temporal variability in Mo and U enrichments can be used to detect environmental changes over decadal timescales, whereas higher-frequency events (seasonal or episodical) are likely not being recorded or are overprinted by post-depositional processes.
- In dynamic fjord-type settings, it is advised to restrain from reconstructing environmental changes based on individual Mo and U enrichment profiles; however, existing environmental (redox) proxies, such as Mo–EF and U–EF covariation patterns, may still retain reliable informative value.

Data availability. Research data are available from Zenodo (https://doi.org/10.5281/zenodo.8399270, Paul, 2023) or upon request to the corresponding author.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/bg-20-5003-2023-supplement.

Author contributions. KMP, MH, and TJ designed the study. SAJ, IB, and HLF organized the fieldwork and carried out the sampling. KMP, SAJ, and TJ conducted the pore water analyses. KMP and MH executed the sequential extractions. KMP, MH, and TJ interpreted the data and wrote the paper, with comments provided by SAJ, IB, and HLF.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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