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 **Investigating the molybdenum and uranium redox proxies in a modern shallow anoxic carbonate rich marine sediment setting of the Malo Jezero (Mljet Lakes, Adriatic Sea)**

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# **Abstract**

 The molybdenum (Mo) and uranium (U) isotope compositions recorded in carbonate rich sediments are emerging as promising paleo-redox proxies. However, the effects of early diagenetic effects within the sediments on these isotope systems are not well constrained. We examined the Mo and U isotopic systematics in anoxic carbonate rich sediments in a semi enclosed karstic marine lake (Malo Jezero) of the Island of Mljet, Adriatic Sea.

 Measurements of water column redox behavior in the lake since the 1950s, have shown a transition from anoxic-sulfidic conditions in the deeper water column to more oxic conditions and anoxia refined to the sediment and pore-waters. A 50 cm long sediment core from the deepest part of the lake, show a transition from moderate to high authigenic Mo and U accumulation with depth, consistent with the changing lake redox environment in the past. In the deep euxinic part of the core, the authigenic Mo and U are isotopically lighter and heavier, respectively, than seawater, following similar systematics as observed in other modern euxinic basins, with high, but non-quantitative, Mo and U uptake into the sediments.

29 Based on Bahamas bank carbonate sediments, it has been suggested that the  $^{238}$ U/ $^{235}$ U ratio is ~+0.25‰ higher compared to seawater from the effects of early carbonate sediment diagenesis and this carbonate vs. seawater off-set is applicable to carbonate rich sediments across the geological past. The shallower part of lake sediment core was deposited under similar redox 33 conditions as the Bahamas sediments, and these sediments show an average  $^{238}$ U/ $^{235}$ U ratio  $+0.31 \pm 0.01\%$  (2SE) higher than seawater. Although the average <sup>238</sup>U/<sup>235</sup>U ratios for these two 35 carbonate rich settings are similar, caution is necessary when inferring seawater  $^{238}$ U/ $^{235}$ U compositions from such sediments, as they contain U from different sources (e.g. diagenetic uptake and carbonate-bound). The Mo isotope compositions within the same Malo Jezero 38 sediments are variable but approaches the seawater composition at low pore-water  $H_2S$ concentrations. This show the potential of using the Mo isotope composition from carbonate rich

- sediments to infer the seawater composition, however, further work is required to establish the
- 41 link between the Mo isotope composition and the chemistry of the pore water environment.
- Keywords: molybdenum, uranium, redox proxies, carbonate sediments, isotopes
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### **1. Introduction**

 Reconstructing the chemistry and redox state of the ocean in the past is critical for understanding the evolution of life (e.g. Chen et al. 2015; Dahl et al. 2010; Lyons et al. 2014). Marine sediments provide geochemical records for such reconstruction. The reliability of identifying good indicators of specific environmental condition (i.e., proxies), and the processes controlling sequestration from an aqueous phase into the sediments, are key points in the "reconstruction chain". The enrichment of several redox-sensitive trace elements, as well as their isotopic composition (e.g. Fe, Mo, Cr, U) in (anoxic) sediments have previously shown to carry important information about the redox-sensitive mechanism of seawater removal and addition to sediments at the time of deposition (e.g. Anbar and Rouxel, 2007; Chen et al. 2015; Dahl et al. 2010; Lyons et al. 2014, Montoya-Pino et al. 2010; Planavsky et al. 2014). Yet, to fully capitalize on the potential of these emerging redox-sensitive proxies for past environments, a thorough understanding of the behavior in modern environments is mandatory.

- 57 The contrasting <sup>238</sup>U/<sup>235</sup>U ratio (reported as  $\delta^{238}$ U the <sup>238</sup>U/<sup>235</sup>U ratio relative to the CRM 145 standard; Andersen et al. 2017) in different marine redox environments showed the promise of 59  $\delta^{238}$ U as a tool to investigate the history of ocean oxygenation at a global scale (Weyer et al. 2008). Both organic-rich sediments and carbonates have been used to reconstruct past variations 61 in oceanic  $\delta^{238}$ U, assuming a quantifiable relationship between the  $\delta^{238}$ U in the geological archive and seawater (e.g. Brennecka et al. 2011; Dahl et al. 2014; Montoya-Pino et al. 2010). Early 63 determination of  $\delta^{238}$ U in modern and fossil carbonates suggested that these materials captured 64  $\delta^{238}$ U identical to seawater (Stirling et al. 2007; Weyer et al. 2008). Further work showed, however, in addition to minor isotope fractionation effects from U incorporation into some carbonates, early 66 diagenetic U addition from anoxic pore-waters may occur and increase the  $\delta^{238}$ U in carbonate rich sediments (e.g. Chen et al. 2016; Chen et al. 2018; Romaniello et al. 2013; Tissot et al. 2018). 68 Consequently, correction factors of  $+0.27 \pm 0.14\%$  and  $+0.24 \pm 0.06\%$ , respectively, were 69 suggested in order to estimate seawater  $\delta^{238}$ U from bulk carbonate sediment  $\delta^{238}$ U compositions (Chen et al. 2018; Tissot et al. 2018).
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72 The Mo isotopic composition (reported as  $\delta^{98}$ Mo, the  $^{98}$ Mo/ $^{95}$ Mo ratio relative to NIST SRM = +0.25‰; Naegler et al. 2014) of organic-rich sediments are used extensively in order to investigate the history of ocean oxygenation (e.g. Arnold et al. 2004; Chen et al. 2015; Dahl, et al. 2010; Dickson et al. 2014; Goldberg et al. 2016; Siebert et al. 2003). Early high-precision Mo isotope work showed large Mo isotope fractionations in marine Fe-Mn oxides, while near- quantitative Mo removal under strongly euxinic (anoxic+sulfidic) conditions in modern Black Sea 78 yielded  $\delta^{98}$ Mo near the seawater composition (+2.3‰) in deposited sediments. This led to the 79 framework that the oceanic  $\delta^{98}$ Mo is largely controlled by the size of the oxidized seafloor and Mn 80 oxide burial, while euxinic sediment archives may record the seawater composition (Barling et al.

81 ; Siebert et al. 2003; Nägler et al. 2011). Additional work showed lower  $\delta^{98}$ Mo than seawater 82 in organic rich sediments at lower  $H_2S$  water column levels, with potential additions of Mo with 83 low  $\delta^{98}$ Mo from particulate water-column shuttle processes (e.g. Arnold et al. 2004; Bura-Nakić et al. 2018; Goldberg et al. 2012; Poulson et al. 2006; Noordmann et al. 2015; Poulson Brucker et 85 al. 2009; Scholz et al. 2017). This add uncertainty when  $\delta^{98}$ Mo in ancient organic-rich sediments 86 is used to reconstruct ocean oxygenation in the past (Arnold et al. 2004; Barling et al. 2001; Chen et al. 2015; Kendall et al. 2017; Siebert et al. 2003). It has also been suggested that non-skeletal 88 carbonates may directly record the  $\delta^{98}$ Mo of the seawater (Vogelin et al. 2009; Vogelin et al. 2010, Czaja et al. 2012). However, the Mo concentration in carbonates are generally low and potentially influenced by early diagenetic conditions within carbonate rich sediments, emphasizing the need 91 to evaluating the processes that effect Mo and its isotopic composition during sedimentation and subsequent burial (Romaniello et al. 2016).

 Coupling the Mo and U isotope redox proxies may provide a better understanding of local vs. 95 global sedimentary  $\delta^{238}$ U and  $\delta^{98}$ Mo signatures (e.g. Andersen et al. 2018; Asael et al. 2013; Bura- Nakić et al. 2018; Kendall et al. 2015; Noordmann et al. 2015). For example, organic rich 97 sediments from modern restricted euxinic basins show systematic  $\delta^{238}$ U vs.  $\delta^{98}$ Mo co-variation, 98 suggested to be dominantly driven by the  $H_2S$  concentration and deep-water overturning rates (Andersen et al. 2018; Bura-Nakić et al. 2018). In this study the Mo and U concentrations and isotope systematics were investigated in modern to near-modern carbonate rich sediments from the marine lake Malo Mljet, Croatia. The deeper part of the seawater-supplied lake has shown variable redox and oxic conditions in the recent past (Lojen et al. 2010; Sondi and Juračić, 2010; Sondi et al. 2016) offering the potential to study U and Mo uptake under changing redox states in a carbonate-dominated sediment matrix. This setting therefor serves as an important analog for examining ancient carbonate sediment archives and the further understanding and utility of the combined Mo and U isotope proxies. 

## **2. Study area**

 The seawater-fed lakes "Malo Jezero" and "Veliko Jezero" ("small" and "big" lake) are located on the western part of Mljet island, in the southern Adriatic Sea (Figure 1). The Mljet island is separated from the mainland by the 8-10 km wide Mljet Canal and Veliko Jezero is connected to 112 the open sea through a shallow channel ('Soline'  $\sim$  2.5 m depth) while another channel ( $\sim$  0.6 m depth) connects Veliko and Malo Jezero (Lojen et al. 2010; Sondi and Juračić, 2010; Sondi et al. 114 2017). Malo and Veliko Jezero are classified as doline lakes, formed as semi-closed depressions in highly permeable limestone karst of Cretaceous age (Schubert, 1909). The karstic depressions, formed during the Mesozoic, have been filled with marine waters since post-glacial sea level rise that intruded seawater through the karst. These submarine karstic connections are found to be active today, detected from sub-bottom echo profiles (Wunsam et al. 1999). Malo Jezero has a 119 surface area of 0.25 km<sup>2</sup> with a maximal depth of approx. 30 m, while Veliko Jezero has a surface 120 area of 1.45 km<sup>2</sup> and a maximal depth of approx. 46 m (Benović et al. 2000).

 Previous studies showed that anhydrous carbonate dominate the mineral composition of the Malo 123 and Veliko Jezero sediments with abundance up to ~70% within Malo Jezero (Lojen et al. 2010;

Sondi et al. 2017; Vuletić, 1953; Figure 2). The ratio between aragonite and calcite calcium

 carbonate polymorphs in the sediment varies, with aragonite the most abundant comprising approximately 60% of total Malo Jezero carbonates, defining the Mljet Lakes system as a site of distinctive authigenic aragonite formation (Sondi and Juračić, 2010; Sondi et al. 2017).

 Hydrographic data for Malo Jezero has been reported since the 1950s (e.g. Vuletić 1953; Buljan and Špan 1976; Benović et al. 2001; Cuculić et al. 2008; Sondi and Jurčić 2010; Vilibić et al. 2010). The pH of the Malo Jezero waters varies between the yearly seasons with lowest pH values during winter (e.g. 8.02 ± 0.01 at the surface in January 2008), and highest pH values during summer (e.g. 8.30 ± 0.01 at the surface in July 2005). In the early 1950s the salinity in Malo Jezero bottom waters was in the range of 36-38 while surface waters were characterized by lower salinity ~29 (Buljan and Špan, 1976). An increase in salinity was noticed in the 1990s with the Malo Jezero bottom waters surpassing salinity of 38.5 (Benović et al. 2000). Spatial and temporal variations of water temperature in the Malo Jezero show typical seasonal variability, reflecting the air temperature changes, with the surface Malo Jezero waters being colder during the winter and warmer during the summer, compared to Adriatic open sea waters. A thermocline appears in early summer, deepening towards early autumn before vanishing during the winter. Records from 1997 to 1999 showed that in the summer months the thermal stratification occurred in the layers between 10-15 m (Malo Jezero) and 15-20 m (Veliko Jezero). During the periods of thermal stratification, the bottom water layers were low in oxygen, with oxygen saturation reaching 144 minimum of 4% (Malo Jezero) and 17% (Veliko Jezero) in October 1997 (Benović et al. 2000).

 Anoxic and euxinic conditions in the water column of Malo and Veliko Jezero has not been observed within the last couple of decades (Jasprica et al. 1995; Benovic et al. 2000). However, vertical sediment profiles of redox sensitive elements within Malo Jezero, indicate the occurrence of euxinic conditions in the bottom water layer of Malo Jezero and a rapid disappearance of such conditions later in the lake history (Sondi et al. 2017). Reported [H2S] (Buljan and Špan, 1976) at 151 25 m water depth for the 1951 to 1961 period were in the range from 6 to 155  $\mu$ mol  $I<sup>-1</sup>$ , showing 152 seasonal variations with the highest [H<sub>2</sub>S] during the winter-autumn period (Supplementary Information Table 1). Also, the lake [H2S] data shows that the extent of anoxia was variable, with the chemocline rising upwards during the winter-autumn period. The variable extent of anoxia is likely linked to the degree of Malo Jezero water mass exchange with Veliko Jezero and the open sea during the year. Sudden environmental change took place in the early 1960s, as a consequence of the enlargement and deepening of the Soline Channel at that time. This resulted in significantly increased water exchange between the open ocean and the lakes with the disappearance of euxinic conditions in Malo Jezero and occurrence of oxic conditions in the bottom waters as determined in July 1961 (Buljan and Špan, 1976). The irregular occurrence of dark layers of different thickness in the laminated sediments deposited before 1960s, indicates that euxinic conditions in Malo Jezero were not permanent, but were interrupted with shorter or longer oxic to suboxic periods (Supplementary Information Table 1 - Buljan and Špan, 1976; Sondi et al. 2017).

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### **3. Sampling**

 Undisturbed sediment cores up to 50 cm long were collected in Malo Jezero using an Uwitec gravity corer in the period 2007-2010 at depths of 30 meter (core C1), 22 meter (core C2) and 13 meter (core C3). In order to extract pore water from sediments, sediment cores were sectioned into 2 cm segments in a glove box under an inert nitrogen atmosphere immediately after sampling. The sampled segments were centrifuged at 4000 x g for 30 minutes and the pore water was extracted using a plastic syringe and filtered through 0.45 µm Minisart cellulose acetate (Sartorius) filters under an inert nitrogen atmosphere. The segments were then freeze-dried and stored until further geochemical analyses were performed. Samples of the surrounding rocks, Cretaceous limestone and Jurassic dolomite, and the two main types of soil samples, *terra rossa* and dark humus rich soil, were also collected. Lake water was collected using a Niskin sampler system at three different depths (0, 12 and 25 m) and stored in acid cleaned plastic bottles.

## **4. Methods**

## 4.1. Sample preparation

 All chemical preparation and sample analyses were performed in the isotope facilities at the Institute of Geochemistry and Petrology, Department of Earth Sciences, ETH Zürich, Switzerland, unless stated otherwise.

 Water sample aliquots (~20 ml) were taken for isotope analyses, aiming for a total of 20–50 ng U and 150–250 ng Mo. Aliqouts were transferred into pre-cleaned Teflon beakers, and spiked with 191 the IRMM–3636  $^{236}$ U– $^{233}$ U double-spike (Richter et al. 2008) aiming for a 1:35 spike to sample 192 ratio, and a Mo– $97$ Mo double-spike (approach adapted from Siebert et al. 2001) aiming for a 193 1:1 spike to sample ratio. These were subsequently dried down (at  $\sim$ 100 °C) and prepared for isotope analyses following methods described in Bura-Nakić et al. (2018). Briefly, following the seawater drying step, a large NaCl precipitate forms. This precipitate is leached using 10 ml 7 N HCl for 24 hours, a treatment which dissolves Mo, U and other metals but minimises dissolution of NaCl. After centrifugation (3500 x g for 10 minutes) the supernatant was removed, dried down and re-dissolved in 5 ml 7N HCl in preparation for column chromatography. The recoveries of 199 both U and Mo in the supernatant were consistently >90% using this procedure (Bura-Nakić et al. 2018).

 Approximately 50–100 mg of the sediment and soil samples were dissolved for analyses. Full dissolution of samples were carried out using standard protocols for silicates, involving mixtures 204 of HF–HNO<sub>3</sub>–HCl and H<sub>2</sub>O<sub>2</sub> in the same manner as described in Andersen et al. (2013). After drying, the samples were dissolved in 10 ml 6 N HCl. One small aliquot was taken to determine elemental concentrations while another aliquot, containing 20–50 ng U and 150–250 ng Mo, was added to pre-cleaned Teflon beakers, spiked with the U and Mo double-spikes, and left to 208 equilibrate on a hotplate before being dried down (at  $\sim$  100 °C). Samples were then re-dissolved in 5 ml 7 N HCl in preparation for column chromatography.

### 4.2. Column chromatography

 A one step purification and U–Mo separation procedure was conducted using RE Resin (Triskem technologies) in custom-made shrink-fit Teflon columns (~0.2 ml resin reservoir) following the procedure of Bura-Nakić et al. (2018). The resin was added to the Teflon columns and pre-216 cleaned using 2 ml 0.1 N HCl–0.3 N HF, rinsed with MQ water, and pre-conditioned with 2 ml 7 217 N HCl. After the samples were loaded in 5 ml 7 N HCl, the matrix was eluted with 10 ml 1 N HCl. 218 The Mo and U fraction were eluted from the resin and collected separately, first with 5 ml 0.2 N HCl, followed by 5 ml 0.1 N HCl–0.3 N HF, respectively. The column chromatography procedure yielded high purity Mo and U fractions with low Mo and U blanks (<13 pg and <22 pg for Mo and 221 U, respectively; Bura-Nakić et al. 2018). Prior to mass spectrometry, the U fractions were dried 222 down and fluxed on a hotplate for 24 hours in a 1 ml 1:1 mixture of concentrated HNO<sub>3</sub> and H<sub>2</sub>O<sub>2</sub>, while the Mo fractions were merely dried down. The purified Mo and U were then re-dissolved in 224 1 ml 0.3 N HNO<sub>3</sub> and 1 ml 0.2 N HCl, respectively, for mass spectrometry. The column separation and purification procedure was tested by processing two open Atlantic Ocean samples, with measured Mo and U isotopic compositions in good agreement with previously reported values. 

4.3. Elemental concentration measurements

 The concentrations of selected elements (see Supplementary Information Tables 2 and 3) were 230 measured in 0.3 N HNO<sub>3</sub> using a Thermo–Finnigan Element XR ICP–MS in both low and medium resolution mode, following measurement protocols in Andersen et al. (2013, 2016). A primary in- house concentration standard was interspersed with three unknowns and a secondary standard (BCR–2) was used to monitor the accuracy and reproducibility of the method. Repeated measurements of BCR–2 gave reproducibilities better than ± 10% (1 S.D.) and mean values within 235 ± 10% of certified concentrations (see Andersen et al. 2016).

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- 4.4. Molybdenum and uranium isotope measurements

 Isotope ratios were measured on a Neptune (Thermo–Finnigan) MC–ICPMS equipped with an AridusII, auto-sampler (CETAC) using a PFA nebulizer and spray chamber (CPI) sample introduction system. Instrumental set-up details are given in Archer and Vance (2008) for Mo isotopes and Andersen et al. (2016) for U isotopes. Analysis of spiked in-house CPI standard (standard/spike ratios in the range of 0.1 to 5) as well as the open-ocean seawater were used to 243 test the reproducibility and accuracy of the Mo isotope method. The long-term average and  $\pm 2$ 244 S.D. reproducibility during the period of this study, gave  $\delta^{98}$ Mo = -0.02± 0.04‰ relative to NIST 245 SRM = +0.25‰ (Bura-Nakić et al. 2018). Analysed seawater samples gave  $\delta^{98}$ Mo of +2.37 ± 246 0.03‰, in agreement with previous data for seawater  $\delta^{98}$ Mo (e.g. Nakagawa et al. 2012). In 247 addition to <sup>238</sup>U/<sup>235</sup>U, <sup>234</sup>U/<sup>238</sup>U ratios were measured and reported as  $(^{234}$ U/<sup>238</sup>U) activity ratios (using half-lifes of Cheng et al. 2013).The verification of the U double spike method was carried out via repeated measurement of the in-house CZ–1 uraninite standard and open-ocean seawater 250 samples. The long-term average and  $\pm 2$  S.D. reproducibility for the CZ-1 standard were -0.04  $\pm$ 251 0.07‰ for  $\delta^{238}$ U and 0.9996 ± 0.0025 for  $(^{234}$ U/ $(^{238}$ U) (Bura-Nakić et al. 2018), in agreement with previously reported values (Andersen et al. 2016; Stirling et al. 2007). Uranium isotopic analysis 253 of five seawater samples gave a  $\delta^{238}U = -0.39 \pm 0.04\%$  and  $(^{234}U/^{238}U)_{act} = 1.147 \pm 0.003$  (Bura-Nakić et al. 2018), again in very good agreement with reported data for seawater (Andersen et al.

 2010, 2014; Tissot and Dauphas, 2015; Weyer et al. 2008). See Bura-Nakić et al. 2018 and Andersen et al. (2016) for further details on methods and performance.

## 4.5. Other analysies

259 The total carbonate content was determined volumetrically with a Scheibler's apparatus (Allison and Moodie, 1965) at the Ruđer Bošković Institute in Zagreb, Croatia. Sulfide concentrations were analysed by linear sweep voltammetry (LSV) within 8 hours of sampling according to established procedures (Bura-Nakić et al. 2009; Ciglenečki et al. 2005; Ciglenečki et al. 2015) at the Ruđer Bošković Institute marine station situated near Šibenik. Electrochemical measurements were performed with μAutolab Electrochemical Instruments (EcoChemie) connected with 663VA Stand Metrohm electrode.

**5. Results**

## **5.1. Elemental concentrations**

## **5.1.1. Sediments and pore-waters**

 The mineral composition of sediment cores taken from the lake sediments of the Malo Jezero at cores C1, C2 and C3 (Figure 1) showed a high carbonate content, dominated by authigenic aragonite (comprising ca. 70-90% of the carbonate content) with minor calcite, magnesium calcite and dolomite. The calcite and dolomite originated from surrounding source rocks of Jurassic- Cretaceous limestones and dolomites. Besides the carbonate phases the sediments also contained small amount of mica, quartz, kaolinite and framboidal pyrite. The total carbonate content within the investigated sediments is high, comprising up to ~70% in cores C1 and C2 and 280 reaching up to ~90% in core C3. The [Sr]<sub>bulk</sub> is also relatively high, ranging from 3078 to 4158 µg  $g^{-1}$ . Concentrations of sedimentary organic carbon in core C1 were measured in Lojen et al. 282 (2010) with  $C_{\text{org}}$  (wt%) varying between 1.4 to 2.1%, with the lowest  $C_{\text{org}}$  at 13 cm depth. Using 283 marine and terrestrial  $\delta^{13}$ C endmembers signatures, Lojen et al. (2010) estimated the fraction of terrestrial organic C ranging from 53 to 76%.

 The distributions of typical lithogenic elements (e.g. Al, Ti and Li) display similar behavior with sediment depth. The concentrations of Al (0.38 to 6.4%), Ti (0.03 to 0.20%) and Li (0.31 to 0.46%) are generally low, but show a gradual increase with depth in each core, while the two deepest cores (C1 and C2) have higher lithogenic element concentrations than the shallower core C3 (Figure 2 and Supplementary Information Table 2).

292 The bulk concentrations of the investigated redox sensitive elements ( $[Mo]_{\text{bulk}}$ ,  $[U]_{\text{bulk}}$ ) and  $[V]_{\text{bulk}}$ 293 display somewhat contrasting behavior. The behavior of  $|V|_{\text{bulk}}$  is similar to that of the lithogenic 294 elements, with lower  $[V]_{bulk}$  in core C3 compared to cores C1 and C2. Similarly, a moderate 295 increase in  $[U]_{bulk}$  (from 2.2 to 8.4 µg g<sup>-1</sup>) can be observed in all three cores, this  $[U]_{bulk}$  increase 296 is intensified below the  $\sim$ 20 cm sediment horizon for the two deepest cores. These sediment 297 horizons also have high  $[Mo]_{bulk}$  (25 to 78 µg g<sup>-1</sup>), with a sharp contrast to lower  $[Mo]_{bulk}$  (2.1 to 20

298  $\mu$ g g<sup>-1</sup>) above the ~20 cm sediment horizon in core C1 (Figure 2 and Supplementary Information 299 Table 2).

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301 Pore-water  $[H_2S]$  for core C1 were determined as part of the present study and combined with already published pore-water data on [Mo], [U], [Fe] and [Mn] (Sondi et al. 2017; Supplementary 303 Information Table 4). The  $[H_2S]$  increase with increasing core depth (0 to 755 µmol  $I^{\text{-}1}$ ) with a sharp increase between the 11 to 17 cm sediment horizons, which likely correspond to the zone where reactive Fe diminishes and reduced Fe is predominating in the form of pyrite (Sondi et al. 2017). The pore-water concentration of dissolved Fe and Mn are highest in the subsurface sediment layer at 1 cm depth, reflecting the position of the oxic-anoxic boundary. The pore-water 308 [Mo] in the top sediment layer (107 nmol  $I<sup>1</sup>$ ) closely match overlying bottom water [Mo] (~110 309 nmol  $\lceil \cdot^1 \rceil$ , while it generally decreases and remains lower through the core (~50-90 nmol  $\lceil \cdot^1 \rceil$ ) indicating removal of Mo from the pore-waters. The [U] in the pore-waters is exhibiting significant decrease (more than 50% within the subsurface layer) in comparison to the [U] in the overlying bottom waters, implying substantial U removal from the pore-waters.

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# 314 **5.1.2 Catchment area and lake water**

 Major and trace element concentrations were determined in the soil (*terra rossa* and humus) as well as karst host rock samples (limestone or dolomite) collected close to the lake (Supplementary Information Table 3). The carbonate host rock samples are characterised by low [Sr] (92 to 392  $\mu$ g g<sup>-1</sup>) as well as low [Al] (153 to 970  $\mu$ g g<sup>-1</sup>), [Li] (0.24 to 1.4  $\mu$ g g<sup>-1</sup>) and [Ti] (7 to 49  $\mu$ g g<sup>-1</sup>). The 319 [Mo] within the host rock samples is low (0.14 to 0.73  $\mu$ g g<sup>-1</sup>), similarly to what is typically observed for carbonates (Vogelin et al. 2009) while the [U] and [V] are higher (1.3 to 3.6 and 3.5 to 23 µg  $g^{-1}$ , respectively). Soil samples are characterised by [Al], [Ti] and [Li] approximatelly three orders of magnitude higher than the host rock samples. Compared to the host rock samples, the soil samples have similar [Sr] (61 to 102  $\mu$ g/g<sup>-1</sup>) while [Mo] (1.4 to 4.0  $\mu$ g g<sup>-1</sup>), [U] (2.4 to 4.0  $\mu$ g g<sup>-1</sup>) 324 and [V] (72 to 168  $\mu$ g g<sup>-1</sup>) are all higher.

325

326 The [Mo] and [U] were measured in filtered waters sampled at 0, 12 and 25 m depth. These show 327 a narrow concetration range from 103.3 to 110.2 nmol  $I^{-1}$  and 13.0 to 13.8 nmol  $I^{-1}$  for [Mo] and 328 [U], respectively, close to average open ocean compositions (Supplementary Information Table 329 5).

330

# 331 **5.2 Mo and U isotopic composition**

#### 332 333 **5.2.1. Sediments**

334 The bulk sedimentary Mo isotope compositions ( $\delta^{98}$ Mo<sub>bulk</sub>) generally increase with depth within 335 the three sediment cores (Figure 2 and Table 1). The  $\delta^{98}$ Mo<sub>bulk</sub> compositions in core C1 varies 336 between +1.8 to +1.4‰, apart from the top sample at 1 cm depth with a lower  $\delta^{98}$ Mo<sub>bulk</sub> (+1.0‰). 337 Core C2 is characterized by  $\delta^{98}$ Mo<sub>bulk</sub> in the range from +2.4 to +1.7‰ while core C3 shows 338 variable Mo isotope compositions, including the lowest,  $δ^{98}$ Mo<sub>bulk</sub> (+1.6 to +0.7‰).

339

340 The cores exhibit limited variation in the bulk sedimentary  $\delta^{238}$ U compositions ( $\delta^{238}$ U<sub>bulk</sub>), ranging 341 from +0.03 to -0.29‰. Core C1 show the least variable  $\delta^{238}U_{\text{bulk}}$  (-0.05 to -0.15‰), followed by

342 core C2 (+0.02 to -0.13‰), while core C3 have  $\delta^{238}U_{\text{bulk}}$  of -0.29‰ for the top 1 cm and +0.03‰

343 at 23 cm depth. The  $(^{234}U/^{238}U)_{act}$  for the bulk sediments increase with depth in all three cores 344 (Figure 2 and Table 1). Cores C2 and C1 have the highest  $(^{234}U/^{238}U)_{act}$  (1.065 to 1.033) while 345 core C3 show lower  $(^{234}U/^{238}U)_{act}$  (from 1.023 to 1.000 for the 1 cm depth sample).

# **5.2.2. Soil and water samples**

348 The measured  $\delta^{98}$ Mo (average +2.41±0.02‰), and  $\delta^{238}$ U (average -0.36±0.02‰) in the Malo Jezero lake waters at 0, 12 and 25 m depth (Supplementary Information Table 5) are close to the 350 open ocean ( $\delta^{98}$ Mo<sub>Sw</sub>=+2.37‰ and  $\delta^{238}$ U<sub>Sw</sub>=-0.39‰; Andersen et al. 2014; Nakagawa et al. 2012). 351 The  $(^{234}U/^{238}U)_{act}$  for the same lake water samples (1.136 to 1.138) are slightly lower than the open ocean seawater (1.147; Andersen et al. 2010). The *terra rossa* and the humus soil samples 353 were characterized by variable  $(^{234}U/^{238}U)_{act}$  (1.102 and 0.977), but similar  $\delta^{238}U$  (-0.28‰ and -354  $0.22\%$ ) and  $\delta^{98}$ Mo (+0.53‰ and +0.62‰), see Table 1.

## **6. Discussion**

358 The main aim is to broaden our understanding of  $\delta^{98}$ Mo and  $\delta^{238}$ U signatures within carbonate 359 rich sediments under changing redox conditions. The discussion on the sedimentary  $\delta^{98}$ Mo and 360  $\delta^{238}$ U budget therefor primarily focusses on the deep Malo Jezero sediments (core C1), due to 361 the available literature data (Sondi et al. 2017) and the most complete  $\delta^{98}$ Mo and  $\delta^{238}$ U data-set. 

# **6.1. The sources of Mo and U in the sediments of Malo Jezero**

 The uptake of the redox-sensitive U, Mo and V in the Malo Jezero sediments will be impacted by various biogeochemical and physical processes. Three main sources may be considered important for the sediments in this particular system; *(1)* the addition from detrital material, e.g. siliciclastic and carbonate, *(2)* the formation of authigenic carbonate phases in the lake and *(3)*  other transport uptake mechanisms, either from authigenic processes directly within the waters and/or pore-water. Below, these three potential sources are discussed in turn.

 *(1)* Sondi et al. (2017) concluded that soils were an important source of material deposited in the Veliko and Malo Jezero based on the similar geochemistry of soil and sediment from sampled cores. However, the soils surrounding the lake are heterogenous and in order to estimate the possible influence of this detrital material on the sediment geochemistry, two general detrital end- members can be defined; limestone/dolomite vs. siliciclastic (clay) dominated soils (Figure 3). These two end-members have distinct geochemical signatures. The Malo Jezero sediments shows a near-linear relationship between typical lithogenic elements (e.g. [Al] vs. [Ti] and [Al] vs. [Li]), suggesting that the Malo Jezero lithogenic fraction in the sediments are a physical mixture of the two generalised detrital end-members (Figure 3). Vanadium concentrations in the sediments also show similar mixing relationship with [Al], while both [U] and [Mo] show generally higher concentrations than can be explained by simple mixing of these two end-member sources (Figure 3). This suggests that, while the redox sensitive V are likely dominated by the detrital fraction, additional source(s) of Mo and U uptake is/are required to explain the sediment data. 

 *(2)* The mineralogy of the sediments within Malo Jezero revealed that aragonite is the dominating carbonate fraction. This source of authigenic-formed aragonite in the lake is manifested in high

sediment [Sr] (3000-5000 µg g<sup>-1</sup>) compared to the soil and limestone host rocks (61 to 392 µg g<sup>-</sup> 387 388 <sup>1</sup>) a feature also observed in previous studies (Sondi and Juračić, 2010; Sondi et al. 2017). The authigenic-formed aragonite could provide an additional source of U and Mo to the sediments. However, marine aragonite is typically characterized by low [Mo] (Voegelin et al. 2009; Romaniello et al. 2016) and no correlation is observed between [Mo] and [Sr] (Figure 3), suggesting limited Mo addition from this source. While [U] is normally relatively high in marine aragonite (2-4 µg  $g^{-1}$  range; Dunk et al. 2002), there is no obvious correlation between [U] and [Sr] (Figure 3), suggesting that the U incorporation into this lake aragonite phase is minor. This suggests that the relatively high [Mo] and [U] observed in the deeper parts of cores C2 and C1 must be dominated by other uptake mechanism(s).

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 *(3)* Authigenic uptake of Mo and U in sediments often occur under reducing conditions. For both U and Mo, early *in-situ* diagenetic precipitation is observed in a range of environments with euxinic pore-waters (e.g. Chen et al. 2018; McManus et al. 2006; Morford et al. 2005; Romaniello et al. 2013, 2016; Zheng et al. 2002). High Mo accumulation also occur in strongly euxinic waters, presumably via the formation of sulfidic molybdate species followed by particulate organic 403 scavenging or formation and scavenging of authigenic FeMoS<sub>4</sub> minerals (e.g. Algeo and Lyons, 2006; Erickson and Helz, 2000; Helz et al. 1996; Helz et al. 2011; Helz and Vorlicek, 2019). Other potential important sources of addition of Mo and U in anoxic sediments may be associated directly with deposited organic matter (e.g. Kowalski et al. 2013; King et al. 2018; Zheng et al. 407 2002) and (particularly for Mo) Fe-Mn shuttle processes (e.g. Algeo & Lyons 2006; Scholz et al. 2017). All three cores show an increase in [U] with depth (Figure 2) suggesting increasing U addition to the sediments with decreasing redox potential at depth in the pore waters. The highest 410 [U] ( $>5 \mu g g^{-1}$ ) is observed at depths below 20 cm in cores C1 and C2, similar to the zones of the 411 highest [Mo] (>40 µg g<sup>-1</sup>). In contrast, the [Mo] is much lower in the horizons above (<5 µg g<sup>-1</sup>) and there is little evidence for any Mo addition in core C3 (Figure 2).

413

## 414 **6.2. Quantification of the authigenic U and Mo fractions in the Malo Jezero sediments**

415 With a general separation of the different sources of U and Mo in the sediments, it is possible to 416 further quantify the authigenic U and Mo fractions, and associated isotope signatures, to provide 417 better constraints on the processes governing the mechanisms of U and Mo enrichment.

418

419 The daughter <sup>234</sup>U to parent <sup>238</sup>U activity ratio, (<sup>234</sup>U/<sup>238</sup>U)<sub>act</sub> can provide an effective method to 420 distinguish between U from an authigenic or detrital source in Holocene-aged sediments (e.g. 421 Holmden et al. 2015; Andersen et al. 2016). This stem from modern seawater  $(^{234}U/^{238}U)_{act}$  is 422  $\sim$  15% enriched in <sup>234</sup>U due to α-recoil processes, while older detrital lithogenic material is normally 423 characterized with a  $(^{234}U/^{238}U)_{act}$  near secular equilibrium (Andersen et al. 2010). The 424  $(^{234}U/^{238}U)_{act}$  in the surface sample of core C3 is in secular equilibrium, implying negligible 425 authigenic U and can therefore be considered a detrital end-member. This sample is 426 characterized by [U]<sub>bulk</sub> of 2.2 µg g<sup>-1</sup>, U/AI ratio of 5.8  $\times$  10<sup>-4</sup> g/g and  $\delta^{238}$ U of -0.29‰. This 427 sediment sample is also characterized by high  $CaCO<sub>3</sub>$  and [Sr] suggesting that, as already 428 discussed, the contribution of U from authigenic aragonite has a minor impact on the sedimentary 429 U budget. Sediments with higher [U] show increasing  $(^{234}U/^{238}U)_{act}$  consistent with increasing 430 authigenic U accumulation. The sediments show a broadly linear correlation between  $(^{234}U/^{238}U)_{act}$  431 vs. 1/[U], suggesting a mixture between detrital and authigenic U (Figure 4). However, the inferred 432 authigenic end-member of this linear mixing line has a  $(^{234}U/^{238}U)_{act}$  of ~1.09, a value significantly 433 Iower than the measured  $(^{234}U/^{238}U)_{act}$  in the lake waters (~1.14). The origin of this relatively low 434 authigenic  $(^{234}U/^{238}U)_{act}$  estimation is not clear. One possibility is incongruent calcite dissolution, 435 as suggested to occur within the reducing Malo Jezero sediments (Lojen et al. 2010), releasing 436 U with low  $(^{234}U/^{238}U)_{act}$  which then mixes with seawater-derived U in the pore-waters. Alternative 437 suggestions may be related to additional groundwater input or dissolution of karst during seawater 438 flow-through in cthe ave systems, both adding U with low  $(^{234}U/^{238}U)_{act}$ . Lower than seawater 439  $(234U/238U)_{act}$  was also observed in the seawater lake Rogoznica, Croatia, which is also fed through 440 karst cave systems (Bura-Nakić et al. 2018). Whichever the mechanism, the uncertainties in the 441  $(234U/238U)_{\text{act}}$  for the both the detrital and authigenic U end-members, limits the usage of 442  $(2^{34}U/2^{38}U)_{\text{act}}$  to quantify the authigenic U contribution in the Mjet lake sediments (see also section 443 6.4).

444

445 Instead, the authigenic to detrital components for both U and Mo can be estimated comparing 446 these to Al, typically used for estimating authigenic enrichment factors (EF; e.g. Tribovilliard and 447 Algeo, 2009). However, the detrital fraction of U and Mo in each sediment horizon is a mixture 448 between the (i) limestone/dolomite and (ii) their weathering products in the form of *terra rossa* and 449 organic richs soils (as shown by the Al vs. Ti correlation, Figure 3). Thus, using only one average 450 [U]/[Al] or [Mo]/[Al] ratio for the detrital fraction would lead to under- and/or overestimation of the 451 authigenic U and Mo component. Instead, to estimate the U and Mo detrital vs. authigenic 452 contribution, a linear regression line can be estimated between the two generalized detrital end-453 members in the [U] or [Mo] vs. [Al] space, which reflect the degree of weathering of the detrital 454 source (Figure 5). Using this approach, the  $[U]_{\text{auth}}$  and  $[Mol_{\text{auth}}$  fractions are estimated to range 455 between 14 to 68%, and 4 to 98% of the bulk, respectively (Table 2). Both [Mo]<sub>auth</sub> and [U]<sub>auth</sub> show 456 progressive increases with depth in all the cores. Furthermore, both cores C2 and C1 show a 457 large increase in [Mo]<sub>auth</sub> between 18 to 25 cm depth, while C3 core show much lower overall 458 [Mo]<sub>auth</sub> (Figure 6). The two upper part of core C3 (1 and 9 cm depth) are characterized by very 459 low [Mo]<sub>bulk</sub> (0.18 and 0.13 µg g<sup>-1</sup>, respectively) close to the average carbonate catchment [Mo] 460 (0.29 µg g<sup>-1</sup>, n=5), showing the [Mo]<sub>auth</sub> is negligible if at all present in these sediments. The 461 estimated [U]<sub>auth</sub> for the top 1 cm in core C1 is the lowest within the investigated sediments (0.33 462  $\mu$ g g<sup>-1</sup>), in agreement with the minimal authigenic U uptake based on the (<sup>234</sup>U/<sup>238</sup>U)<sub>act</sub> composition. 463

### 464 **6.3. Estimating the authigenic U and Mo isotope signatures in the Malo Jezero sediments**

465 The estimation of  $\delta^{238}U_{\text{auth}}$  and  $\delta^{98}M_{\text{Oauth}}$  build on [U]<sub>auth</sub> and [Mo]<sub>auth</sub> calculations (Figure 6). 466 Combining the relative size of the detrital component with an estimated isotopic composition can 467 provide an estimated isotope composition of the authigenic fraction. As the detrital fraction for 468 each sediment sample is a mixture between the two detrital end-members, the isotope 469 composition also needs to be estimated for each sediment sample. For U, both end-member have 470 very similar  $\delta^{238}$ U (the clay fraction of -0.28‰ and carbonate/dolomite fraction of -0.29‰, based 471 on the 1 cm sediment sample in core C3) and both are similar to average 'bulk Earth' and 472 continental crust  $\delta^{238}$ U estimates (Andersen et al. 2017; Tissot et al. 2015). For the detrital Mo, 473 the  $\delta^{98}$ Mo of the clay fraction were estimated to be +0.58‰ (average of the measured soils) while 474 the carbonate/dolomite fraction were estimated to be +1.03‰ (based on the 1 cm sediment

- 475 sample in core C3), both slightly higher than typically used for detrital  $\delta^{98}$ Mo compositions (~0.0) 476 to +0.3‰; Kendall et al. 2017). The  $\delta^{238}$ U<sub>auth</sub> and  $\delta^{98}$ Mo<sub>auth</sub> estimates (Figures 6 and 7 and Table 477 2), have uncertainties estimated based on propagated measurement errors scaled to the relative 478 size of the detrital component (see Andersen et al. 2014 for details). This error propagation leads 479 to increasing uncertainty estimates with increasing detrital contribution. Therefore, detrital-480 dominated sediments with uncertainties >0.4‰ (±2σ) on the  $\delta^{238}$ U<sub>auth</sub> or  $\delta^{98}$ Mo<sub>auth</sub> (Table 2) are 481 deemed unreliable and omitted from figures and further discussion.
- 482

483 Core C1 show  $\delta^{238}U_{\text{auth}}$  from +0.25‰ to -0.07‰, all significantly higher than the seawater  $\delta^{238}U$ 484 composition of -0.39‰ and with a systematic decrease in  $\delta^{238}$ U<sub>auth</sub> with depth and higher [U]<sub>auth</sub> 485 (Figures 6 and 7). The measured sediments in core C2 and at depth in core C3 show high  $\delta^{238}$ U<sub>auth</sub> 486 similar to core C1 (Figures 6 and 7). Similarly to U, [Mo]<sub>auth</sub> show a progressive increase with 487 depth for core C1, with  $\delta^{98}$ Mo<sub>auth</sub> compositions between +1.09‰ and +2.02‰. Below ~20 cm the 488  $\delta^{98}$ Mo<sub>auth</sub> is relatively homogenous at ~+1.7‰, while above, the  $\delta^{98}$ Mo<sub>auth</sub> varies, including the 489 Iowest  $\delta^{98}$ Mo<sub>auth</sub> composition at 1 cm depth (+1.09‰). The  $\delta^{98}$ Mo<sub>auth</sub> varies from +1.85‰ to 490  $+2.45\%$  in core C2, with samples at 37 cm and 17 cm close to the seawater  $\delta^{98}$ Mo composition. 491 In core C3 the  $\delta^{98}$ Mo<sub>auth</sub> can only be estimated for the two deepest samples (17cm and 24 cm) 492 giving +1.18 and +1.72‰ (Figures 6 and 7).

493

### 494 **6.4. Evaluating the controls on the authigenic Mo and U in the Malo Jezero sediments**

495 The three water samples taken at different lake depths show both [U], [Mo],  $\delta^{238}$ U and  $\delta^{98}$ Mo very 496 close to open-ocean seawater compositions (Supplementary Information Table 5). This shows 497 that the lake is dominated by seawater-derived U and Mo, the authigenic Mo and U in sediments 498 can be directly compared to the seawater. The progressive increase in  $[U]_{\text{auth}}$  and  $[Mol_{\text{auth}}$  with 499 depth in cores C1 and C2 is similar to observations in Bahamas bank carbonate sediments, where 500 [U]<sub>auth</sub> and [Mo]<sub>auth</sub> increases have been interpreted to be from increasing organic matter 501 respiration and buld-up of pore-water H2S (Romaniello et al. 2013; 2016). Cores C1 and C2, 502 however, show a clear transition around  $\sim$ 20 cm depth with a sharp increase in sedimentary 503 [Mo]<sub>auth</sub> and [U]<sub>auth</sub> as well as pore-water [H<sub>2</sub>S]. This increase in [Mo]<sub>auth</sub> and [U]<sub>auth</sub> at depth could 504 be directly related *in situ* diagenetic respiration of organic matter or it may reflect an euxinic redox 505 regime in the lake with other pathways of U and Mo uptake at the time of sediment deposition 506 below  $\sim$ 20 cm depth. Correlations between Mo]<sub>auth</sub> and [U]<sub>auth</sub> vs. C<sub>org</sub> may potentially help to 507 distinguish between these two scenarios.

508

509 Positive correlations between both U vs.  $C_{org}$  and Mo vs.  $C_{org}$  have been observed in many 510 anoxic/euxinic sediments (e.g. McManus et al. 2005; Algeo and Lyons, 2006; Wagner et al. 2017). 511 The positive Mo vs. C<sub>org</sub> correlation have been attributed to processes of Mo scavenging from 512 seawater by organic shuttle processes (e.g. Algeo and Lyons, 2006; Wagner et al. 2017), or, 513 alternatively, a lowering of Mo solubility in waters from microbial sulfate reduction fueling on C<sub>org</sub> 514 (e.g. Helz and Vorlicek, 2019). A similar debate exist for positive U vs. C<sub>org</sub> correlations, suggested 515 to either occur from direct U scavenging with organic material (e.g. Zheng et al. 2002) or C<sub>org</sub> 516 fueling microbial mediated metal (and U) reduction within sediments (e.g. McManus et al. 2005; 517 Lovley et al. 1991). Available C<sub>org</sub> vs. bulk [Mo] and [U] data for core C1, show no positive 518 correlations and little variability in  $C_{org}$  with depth (Figure 8). Similarly, no U vs.  $C_{org}$  correlation 519 have been observed in Bahamas bank carbonate sediments (Tissot et al. 2018). In contrast, the 520 four upper samples from core C1 (<20 cm) show weak negative correlations between [Mo] vs. 521 C<sub>org</sub> and [U] vs. C<sub>org</sub> (Figure 8), in opposite direction to general observations in euxinic basins 522 (Algeo and Lyons, 2006). If these observations are considered significant, then increased 523 microbial metal reduction leading to higher [H<sub>2</sub>S] and more efficient Mo and U uptake, while using 524 up available  $C_{org}$  in the process, could be an explanation. The lack of correlation between the [Mo] 525 vs.  $C_{\text{org}}$  in the deeper part of the core (>20cm) is consistent with the model of Helz and Vorlicek 526 (2019) suggesting [Mo] vs. C<sub>org</sub> correlations degrade where the flux of Mo is independent of the 527 sulfate reduction process in strongly euxinic waters.

528

529 Yet in euxinic sediments, even minor U or Mo accumulation associated directly with organic 530 matter and/or Fe-Mn oxides, may have a significant impact on the Mo and U isotope systematics 531 as these uptake processes commonly show large isotopic fractionation (e.g. Abshire et al. 2020; 532 Barling et al. 2004; King et al. 2018; Kowalski et al. 2013). Impact of such processes in the lake 533 sediments can be investigated comparing  $\delta^{98}$ Mo<sub>auth</sub> and  $\delta^{238}$ U<sub>auth</sub> compositions to inverse [Mo]<sub>auth</sub> 534 and [U]<sub>auth</sub> (Figure 9), where correlations can be attributed to mixing of different sources of 535 authigenic Mo and U. The  $\delta^{238}U_{\text{auth}}$  show little variability over a range of [U]<sub>auth</sub>, with a general 536 trend of slightly increasing  $\delta^{238}U_{\text{auth}}$  with lower [U]<sub>auth</sub>. This suggest little contribution, if any, of 537 authigenic U associated directly organic matter carrying low  $\delta^{238}U_{\text{auth}}$  as observed in other anoxic 538 settings (e.g. Abshire et al., 2020; Holmden et al., 2015). Sources of authigenic U may be further 539 explored cross-plotting  $δ^{238}U_{\text{auth}}$  vs. [U]<sub>auth</sub>/C<sub>org</sub>. Here the lake sediments show a negative 540 correlation for samples with low [U]<sub>auth</sub>/C<sub>org</sub> (<2) and  $\delta^{238}U_{\text{auth}}$  decreasing from +0.2 to -0.05‰, 541 while at higher [U]<sub>auth</sub>/C<sub>org</sub> (>2), the δ<sup>238</sup>U<sub>auth</sub> are similar, centered around -0.05‰ (Figure 9). This 542 relationship suggests that at high [U]auth/Corg (>2) the U uptake is dominated by *in situ* U reduction, 543 while at lower [U]<sub>auth</sub>/C<sub>org</sub> (<2), U associated directly with organic matter constitute a significant U 544 fraction when compared to the *in situ* U reduction. This interpretation is consistent with the 545 sediment data from the Namibian shelf in Abshire et al. (2020), except that in this study the U 546 associated with organic matter has a high δ<sup>238</sup>U<sub>auth</sub> (~+0.2-0.4‰; Figure 9) in contrast to low δ<sup>238</sup>U 547  $(-0.6\%)$  in Abshire et al. (2020). The potential U source with such high δ<sup>238</sup>U, could be associated 548 with detrital, rather than authigenic, organic matter and adsorption of isotopically heavy  $U^{+4}$  before 549 transportation to the lake and sediments. Such a detrital organic U source would not be accounted 550 for in the estimated detrital U fraction and it would provide U with low  $(^{234}U/^{238}U)_{act}$  and a 551 mechanism to lower estimated authigenic  $(^{234}U/^{238}U)_{act}$  end-member (see section 6.2). For the Mo ss 552 systematics, the sediments with the lowest  $\delta^{98}$ Mo<sub>auth</sub> compositions also correspond to the lowest 553 [Mo]<sub>auth</sub> and could have significant Mo addition from organic matter and/or Fe-Mn oxides. The 554 samples with the high authigenic [Mo]<sub>auth</sub> below 20 cm depth in cores C1 and C2, on the other 555 hand, suggest one dominating authigenic [Mo]<sub>auth</sub> source (Figure 9). Cross-plotting  $\delta^{98}$ Mo<sub>auth</sub> vs. 556 [Mo]<sub>auth</sub>/C<sub>org</sub> show no correlation to indicate mixing between Mo associated organic matter and/or 557 Fe-Mn oxides with low  $\delta^{98}$ Mo<sub>auth</sub> compared to [Mo]<sub>auth</sub> associated with formation and scavenging 558 of sulfidic molybdate species or FeMoS<sub>4</sub> minerals with high  $\delta^{98}$ Mo<sub>auth</sub> (Figure 9). 559

560 In summary, these observations suggest that the both the U and Mo uptake is primarily related to 561 microbial sulfate and metal reduction processes (and [H2S] formation for Mo) within the deeper 562 parts of cores C1 and C2. In shallower sediments with lower authigenic U and Mo uptake from  these processes, the role of organic matter scavenging and/or Fe-Mn adsorption for U and Mo may be significant contributors to the authigenic U and Mo fractions.

## **6.5. Coupled authigenic Mo and U and isotope systematics in the Malo Jezero sediments**

 With a general understanding of the dominant uptake mechanisms for U and Mo, it offers the possibility to investigate the combined behavior of the two metals in more detail (Figure 10). The modest authigenic U and Mo enrichments in sediments above ~20 cm (Figure 10 and Supplementary Information Table 6) is consistent with anoxia/euxinia predominantly confined to 571 the pore-water of the sediments (Tribovilliard and Algeo, 2009), while the significant increase in 572 [Mo]<sub>auth</sub> below  $\sim$ 20 cm depth (cores C1 and C2) reflect Mo and U accumulation under euxinic water conditions during sediment deposition. These interpretations align well with the observed changing redox regime from euxinic to oxic bottom waters in the lake over time. Based on the geochemical systematics, the sediments can be divided into three zones with increasing depth: *i*) a zone of Mo addition from particulate transport; *ii*) a zone of pore-water Mo and U uptake and *iii*) a deeper zone of previously water column/pore-water Mo and U uptake.

 Zone *i)* shows high [Fe] and [Mn] in the pore-water profiles (Figure 11) suggesting a zone of Fe- Mn oxide addition and dissolution in the shallowest sediment (<1 cm depth). Pore-water [Mo] surrounding these sediments are close or just below seawater [Mo] (Figure 11). Molybdenum transport from an Fe-Mn particulate shuttle process (e.g. Scholz et al. 2017) is therefore a likely 583 explanation for the relatively low  $\delta^{98}$ Mo<sub>auth</sub> (1.02%) in the top 1 cm sediment. However, this 584 sediment sample has relatively low  $[Mo]_{bulk}$  (2.1 µg g<sup>-1</sup>) and any Mo addition from Fe-Mn oxides can therefor only provide a small contribution to the total accumulated Mo in the sediments as also discussed in section 6.4).

 Zone *ii)* the shallower <20 cm sediments in cores C2 and C1 are characterized with relatively low 589 Mo<sub>EF</sub> (0.3 to 11) and  $U_{EF}$  (1.1 to 2.3) suggesting euxinic conditions localized within the pore-waters 590 (Figure 8). The  $\delta^{238}U_{\text{auth}}(\sim +0.1 \text{ to } +0.25\%)$  is consistent with the observed  $\delta^{238}U$  range in modern sediments with variable anoxic pore-water levels (e.g. Andersen et al. 2016). The data would be consistent with permanent anoxia and a shallow oxic-anoxia boundary (low oxygen penetration) 593 in the pore-waters with authigenic U accumulation and a  $\Delta^{238}$ U ~+0.6‰ (Andersen et al. 2014; Figure 7), however, as discussed in section 6.4, U addition associated with organic matter may 595 also be significant. The  $[Mol<sub>auth</sub>$  varies between +1.0 to +2.3‰ in the same samples. Such 596  $\delta^{98}$ Mo<sub>auth</sub>, lower than the seawater  $\delta^{98}$ Mo composition, have been observed in reducing organic 597 carbon-rich sediments, attributed to a range of processes including low  $[H<sub>2</sub>S]$  and incomplete 598 thiomolybdate or FeMoS<sub>4</sub> transformation and, as discussed in section 6.4, addition of isotopically light Mo associated with organic matter deposition or a Fe-Mn shuttle process (e.g. Azrieli-Tal et al. 2013; Poulson Brucker et al. 2009; Kowalski et al. 2013; Kendall et al. 2017; Scholz et al. 2017; 601 King et al. 2018; Helz & Vorlicek, 2019). These  $\delta^{98}$ Mo<sub>auth</sub> compositions are similar to observations 602 in Bahamas bank carbonate sediments where the samples with the lowest pore-water [H<sub>2</sub>S] have 603 low sedimentary [Mo] and  $\delta^{98}$ Mo in the range from ~1.05‰ to ~1.25‰ (Romaniello et al. 2016). 604 The negative [Mo] and [U] vs. C<sub>org</sub> correlations suggest increased microbial sulfate reduction using 605 up available C<sub>org</sub> (Figure 8), a process that fits with the increasing gradient in [H<sub>2</sub>S] with depth and 606 the observed  $\delta^{238}U_{\text{auth}}$  and  $\delta^{98}M_{\text{Output}}$  systematics. In the upper part (5 to 9 cm) with low pore-water

607 [H<sub>2</sub>S] (0.6 to 2.1 µmol  $I^{-1}$ ), the authigenic [Mo] is low, but the  $\delta^{98}$ Mo<sub>auth</sub> close to the seawater  $\delta^{98}$ Mo 608 composition. The  $[H_2S]$  is well below the threshold (~11 µmol  $I^1$ ) needed for the complete 609 transformation of MoO<sub>4</sub><sup>2</sup> to MoS<sub>4</sub><sup>2</sup> (Erickson and Helz, 2000). Consequently, for this type of 610 authigenic Mo uptake, Mo is expected to be incorporated into the solid phase mostly in the form 611 of MoO<sub>4</sub><sup>2-</sup> and this process appears to be accompanied by a low levels of Mo enrichment, but with 612 little net Mo isotope fractionation. The deeper part (13 to 17 cm) is characterized by higher 613 porewater  $[H_2S]$  (29 to 200 µmol  $I^1$ ) and the sediments have higher authigenic [Mo], but  $\delta^{98}$ Mo<sub>auth</sub> 614 lower than the seawater  $\delta^{98}$ Mo composition. These observations are in accordance with previous 615 findings (Bura-Nakić et al., 2018) and could relate to the Mo isotope fractionation in the earliest 616 stages of Mo removal is significant and driven by  $[H_2S]$  near the action point of switch for complete 617 transformation of MoO<sub>4</sub><sup>2</sup> to MoS<sub>4</sub><sup>2</sup> or FeMoS<sub>4</sub> formation (Helz et al., 1996; Helz and Vorlicek, 618 2019; Kerl et al., 2017; Nägler et al., 2011; Tossel et al. 2005). The data within this study is 619 generally in agreement with the work of Romaniello et al. (2016) highlighting the important role of 620 [H<sub>2</sub>S] regulating the [Mo] within the pore-waters and sedimenst. However, in contrast to the 621 observations in Romaniello et al. (2016) there is no clear trend between [Mo]<sub>auth</sub> vs.  $\delta^{98}$ Mo<sub>auth</sub>, with 622 the highest  $\delta^{98}$ Mo<sub>auth</sub> recorded at a very low [Mo]<sub>auth</sub> and almost in absence of pore-water [H<sub>2</sub>S].

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624 Zone *iii)* the high [Mo]auth and [U]auth below 20 cm depth in both cores C2 and C1 suggest highly 625 euxinic conditions in the lake bottom- and pore-waters during sediment deposition. Bura-Nakić et 626 al. (2018) investigated combined  $\delta^{98}$ Mo<sub>auth</sub> vs.  $\delta^{238}$ U<sub>auth</sub> in another euxinic Croatian sea-water lake 627 (Rogoznica) and compared the results to organic-rich sediments from other restricted euxinic 628 basins. The deep sediments in the lake show combined  $\delta^{98}$ Mo<sub>auth</sub> vs.  $\delta^{238}U_{\text{auth}}$  systematics that are 629 broadly similar to observations from Saanich Inlet and Cariaco Basin (Figure 10). The  $\delta^{98}$ Mo<sub>auth</sub> 630 vs.  $\delta^{238}U_{\text{auth}}$  systematics of the modern euxinic basins revealed patterns that are likely to be mainly 631 controlled by interlinked factors of basin size, [H<sub>2</sub>S] and deep-water renewal rates (Bura-Nakić et 632 al. 2018). In this framework, strongly euxinic basins with slow deep-water renewal rates, lead to 633 near quantitative Mo uptake and nearly unfractionated Mo isotope compositions in the sediments. 634 This may, for instance, be accomplished by near complete transition from molybdate to tetra-635 thiomolybdate and effective scavenging to the sediments. In contrast, in basins with fast deep-636 water renewal rates, less-effective Mo scavenging lead to non-quantitative removal of Mo from 637 the water column and expressed Mo isotopes fractionations in the sediments. If the transformation 638 to tetra-thiomolybdate is incomplete, Mo scavenged to the sediments will express a  $\Delta^{98}$ Mo that 639 are ~0.7‰ lower than seawater (e.g. Bura-Nakić et al., 2018; Dahl et al., 2010; Dickson; 2017). 640 Such a scenario for the Mo accumulation could be consistent with the average  $\delta^{98}$ Mo<sub>auth</sub> of ~1.6‰ 641 in the sediments below 20 cm in core C1 (Figure 7 and 10). Core C2, below 20 cm, show similar 642 behavior as core C1 for two samples, while two other samples have higher  $\delta^{98}$ Mo<sub>auth</sub> near the 643 seawater composition (Figure 7). The more variable  $\delta^{98}$ Mo<sub>auth</sub> for core C2 may be related to a 644 position closer to the lake's chemocline and variable  $[H_2S]$  in the water column. The U uptake into 645 the organic carbon-rich sediments of euxinic basins are normally less quantitative than that of Mo, 646 expressing a  $\Delta^{238}$ U of ~+0.6‰ between the reduced U in the sediments and the overlaying bottom-647 waters (Andersen et al., 2014; Andersen et al., 2018; Bura-Nakić et al., 2018). Given a 648 predominantly U uptake from diffusion and reduction within pore-waters of the sediments, slow 649 deep-water renewal compared to U diffusion rates, may deplete U in the water column above the 650 sediments (e.g. Anderson et al., 1989; Rolison et al., 2017) and drive the waters towards lower

651  $\delta^{238}$ U. The effect of slow deep-water renewal rates is that U removal becomes more quantitative 652 and the  $\delta^{238}U_{\text{auth}}$  in the sediments move towards the seawater  $\delta^{238}U$  composition (e.g. Andersen 653 et al., 2014; Andersen et al. 2018; Bura-Nakić et al., 2018). Thus, as with  $\delta^{98}$ Mo<sub>auth</sub>, the  $\delta^{238}$ U<sub>auth</sub> 654 data is fully consistent with bottom water euxinia in the deeper part of the lake, but with a relatively 655 fast deep-water overturning time compared to the basin size, leading to  $\delta^{238}U_{\text{auth}}$  of ~0‰ in 656 sediments, ~0.4‰ higher than open ocean seawater  $\delta^{238}$ U composition (Figure 7 and 10).

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 Similar interpretations as based on the Mo and U isotope systematics are, in principle, possible to obtain from Mo vs. U enrichment factors (using Al normalization, see Tribovilliard et al. 2012). In this study, the Al normalization was performed using the calculated detrital [U]/[Al] and [Mo]/[Al] for each specific sample, based on the two detrital end-members (Supplementary Information Table 6). It must be noted that, the large carbonate fraction in these sediments, have to be considered when comparing these results to other sediments, richer in detrital siliciclastic material 664 (Figure 10). The observed  $M_{\text{OEF}}/U_{\text{EF}}$  for the deep part of cores C2 and C1 are about twice as high as the seawater Mo/U molar ratio, a trend often interpreted as evidence for Mo addition to sediments via Fe-Mn particulate shuttle processes (e.g. Algeo and Tribovillard, 2009). However, 667 similar high Mo<sub>EF</sub>/U<sub>EF</sub> have also been observed for the Saanich Inlet and Cariaco Basin and the 668 combined Mo<sub>EF</sub>/U<sub>EF</sub> and  $\delta^{98}$ Mo<sub>auth</sub>/ $\delta^{238}$ U<sub>auth</sub> systematics (Figure 10) are fully consistent with different Mo and U removal rates within euxinic basins with relatively fast deep-water overturning rates (see Bura-Nakić et al. 2018 for discussion).

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### **6.6. Implications for the interpretation of δ <sup>238</sup>U and δ** 672 **<sup>98</sup>Mo in carbonate-rich sediments**

 It is clear that the process of early diagenesis from reducing pore-water conditions within carbonate-rich sediments may lead to the accumulation of authigenic U and Mo (e.g. Romaniello et al. 2013, 2016). Based on Bahamas carbonate-rich sediments, Chen et al. (2018) and Tissot 676 et al. (2018) suggested a  $\Delta^{238}$ U of +0.27 ± 0.14‰ and +0.24 ± 0.06‰, respectively, to be 677 applicable to derive seawater  $\delta^{238}$ U estimates from measured bulk (anoxic) carbonate  $\delta^{238}$ U. The  $\Delta^{238}$ U of the carbonate sediments in this study were on average +0.29 ± 0.03‰ higher than seawater. The carbonate sediments from 1 to 17 cm depth in core C1 are interpreted to have been deposited under reducing pore-waters and these samples are therefore more comparable with the conditions of the Bahamas bank carbonate-rich sediments. These core C1 sediments 682 have a mean  $\Delta^{238}$ U of +0.31 ± 0.01‰ compared to seawater, slightly higher but within uncertainty of the Bahamas carbonate-rich sediments (Cheng et al. 2018; Romaniello et al. 2013; Tissot et 684 al. 2018). Despite the similar  $\Delta^{238}$ U at these different settings, it has to be noted that the observed  $\delta^{238}$ U in carbonate-rich sediments are a combination both detrital and authigenic U which may 686 have variable  $\delta^{238}U$  and isotope fractionation factors (e.g. dependent on oxygen penetration depth; Andersen et al. 2014). Another uncertainty is where the authigenic U is bound in the sediments, as already discussed, the majority of the U is not associated with the authigenic- formed aragonite in the Malo Jezero sediments. Instead it appears that the authigenic U is added 690 as  $U^{\vee}$  with associated U isotope fractionation. Authigenic U addition in the Bahamas carbonate- rich sediments has been suggested to be either in the form of reduced U mineral phases or 692 incorporated into the carbonate crystal lattice under anoxic conditions (e.g. Chen et al. 2018), but 693 further work is required to understand where this  $U^{\vee}$  is hosted and its exact associated U isotope fractionation in anoxic carbonate-rich settings.

- 695 Core C1 with high-resolution pore-water  $[H_2S]$  data allows to discuss the relationship between the 696 sediment [Mo] concentration and pore-water [H<sub>2</sub>S]. The [H<sub>2</sub>S] within the upper 17 cm is increasing from 0.6 to 200 µmol  $I<sup>-1</sup>$  while the bulk sediment [Mo] increases (from 2.1 to 9.6 µg g<sup>-1</sup>). Romaniello 698 et al. (2016) showed that Mo-enriched Bahamas carbonates ( $[Mo]$  1-10  $\mu$ g g<sup>-1</sup>) deposited under 699 high pore-water  $[H_2S]_{aa}$  can record  $\delta^{98}$ Mo values close to seawater. However, as observed in this 700 study and in the case of Romaniello et al. (2016), there may be an  $\Delta^{98}$ Mo offset of ~-0.5‰ between 701 the seawater and bulk sediments even at high pore-water  $[H_2S]$ . In contrast, in Malo Jezero the  $702 - \delta^{98}$ Mo composition is approaching seawater in the anoxic carbonate-rich sediments with low pore-703 waters  $[H_2S]$  (<< 11 µmol  $I^1$ ) situated close to the sediment surface (Figure 10 and 11). A few 704 samples from core C2 are characterized with the highest  $\delta^{98}$ Mo near the seawater composition. The more variable  $\delta^{98}$ Mo<sub>auth</sub> in core C2, compared to C1, may be a consequence of the closer 706 vicinity to the chemocline in the lake and more variable redox conditions within the pore-waters. 707 However, neither pore-water  $[H_2S]$  nor other trace metal data for core C2 is available, limiting a 708 more detailed discussion. Core C3 is characterized with the lowest [Mo]<sub>bulk</sub> and  $\delta^{98}$ Mo<sub>bulk</sub> and the 709 Mo is dominated by the (detrital) carbonate/dolomite fraction with a  $\delta^{98}$ Mo near ~+1.1‰. Overall, 710 the data suggest that pore-water  $[H_2S]$  during early diagenetic processes is critical for authigenic 711 Mo uptake, as also suggested for Bahamas bank carbonate-rich sediments (Romaniello et al. 712 2016), while Mo uptake associated directly with organic matter or Fe-Mn shuttle processes appear 713 secondary in this setting. Clearly, further work is warranted to identify the link between the pore-714 water environment, the host of authigenic Mo and isotope fractionation mechanisms in anoxic 715 carbonate-rich sediments.
- 716

### 717 **7. Conclusions**

 The geochemistry of the carbonate-dominated sediment samples in Malo Jezero are recording the significant documented changes in water column redox conditions. In the deepest part of core C1 (>20 cm depth), high authigenic Mo and U accumulation are consistent with euxinic bottom-721 water conditions during sediment deposition. Both the  $\delta^{98}$ Mo<sub>auth</sub> (~+1.6‰) and  $\delta^{238}$ U<sub>auth</sub> (~-0‰) are different from the seawater composition, showing similar systematics as observed in Saanich Inlet and Cariaco Basin organic carbon-rich sediments. These observations are consistent with U and Mo uptake from euxinic deep water and pore-water, in a basin with relatively fast deep-water renewal rates. In the shallower part of the same core, both Mo and U uptake likely occurred from a reducing pore-water environment leading to lower authigenic Mo and U accumulation. While 727 the  $\delta^{238}U_{\text{auth}}$  is relatively similar to the deeper section, the  $\delta^{98}Mo_{\text{auth}}(1.1 \text{ to } 2.3\%)$  is more variable,

728 primarily linked to variable pore-water  $[H_2S]$ .

729

730 The results from Malo Jezero show similar  $\delta^{98}$ Mo and  $\delta^{238}$ U as recorded within the Bahamas bank 731 carbonate-rich sediments (Chen et al. 2018; Romaniello et al. 2013; Tissot et al. 2018). The 732 results here are generally in accordance with reported shift in bulk sedimentary  $\delta^{238}$ U towards 733 higher values due to the diagenetic processes of U uptake from reducing pore-waters. The  $\Delta^{238}$ U 734 between the bulk  $\delta^{238}$ U carbonate-rich sediments and seawater (+0.31 ± 0.01‰) is slightly higher, 735 but within uncertainty of suggestions based on Bahamas samples  $(+0.27 \pm 0.14)$ % (Chen et al. 736 2018) and +0.24 ± 0.06‰, (Tissot et al. 2018)). If such a  $\Delta^{238}$ U is used as a 'correction factor' for 737 obtaining seawater compositions in paleo-redox studies, it is important to consider the exact

738 deposition environment of the samples (oxic vs. anoxic pore-waters) and that the bulk  $\delta^{238}$ U is a 739 mixture of authigenic and detrital derived U. The results also suggest that  $\delta^{98}$ Mo in carbonate-rich

sediments can, under certain conditions, be similar to seawater, however, further work is needed

741 to understand the processes controlling the  $\delta^{98}$ Mo in carbonate-rich sediments and its potential

- as a paleo-redox proxy.
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### **Table captions (within text)**

1012 **Table 1.** Bulk sedimentary (cores C1, C2 and C3) Mo and U isotopic composition (δ<sup>98</sup>Mo<sub>bulk</sub> and 1013  $\delta^{238}$ U<sub>bulk</sub>) as well and (<sup>234</sup>U/<sup>238</sup>U)<sub>act</sub> in Malo Jezero sediments.

1014 **Table 2.** Sedimentary (cores C1, C2 and C3) authigenic Mo and U compositions (δ<sup>98</sup>Mo<sub>auth</sub> and 1015  $\delta^{238}$ U<sub>auth</sub>) and concentration ([Mo]<sub>auth</sub> and [U]<sub>auth</sub>) in Malo Jezero sediments.

## **Figure captions**

 **Figure 1.** Location of Mljet Island, Croatia, in the Adriatic Sea and the Malo and Veliko Jezero at the western tip of the Island. The sediment sampling locations are indicated at lake depths of 30 meter (core C1), 22 meter (core C2), and 13 meter (core C3).

 **Figure 2.** Vertical distribution of trace elements (Al, Ti, Sr, V, Mo and U) in Malo Jezero sediments for cores C1 (blue triangles), C2 (green circles) and C3 (red squares). The vertical distribution of 1024 CaCO<sub>3</sub> content (wt%) (data for C1 core are taken from Sondi et al. (2017)) and of bulk sedimentary  $\delta^{98}$ Mo,  $\delta^{238}$ U and  $(^{234}$ U/ $^{238}$ U)<sub>act</sub> is also shown.

 **Figure 3.** [Al]bulk vs [Ti]bulk, [Li]bulk, [Sr]bulk, [Mo]bulk, [U]bulk, [V]bulk as well as [Sr]bulk vs [Mo]bulk, [U]bulk, 1028 [V]<sub>bulk</sub> covariation in Malo Jezero sediments for cores C1 (blue triangles), C2 (green circles) and C3 (red squares). The detrital limestone/dolomite (grey) and soil siliciclastic (yellow) end- members are marked as rhombs. The limestone/dolomite end-member represents an average of five samples while the soil-endmember is based on the *terra rossa* sample.

**Figure 4**.  $(^{234}U/^{238}U)_{act}$  covariation with  $[Sr]_{bulk}$  (A) and with  $1/[U]_{bulk}$  (B) for Malo Jezero sediments in cores C1 (blue triangles), C2 (green circles) and C3 (red squares). The *terra rossa* and dolomite/limestone (top sample in core C3) detrital end-members are shown as yellow and grey 1036 rhomb, respectively. The green cross is the measured  $(^{234}U/^{238}U)_{act}$  in the lake waters (average of 1.137). Arrow represent a two-component mixing estimate.

 **Figure 5**. Uranium (A) and Mo (B) to Al cross-plots for cores C1 (blue triangles), C2 (green circles) and C3 (red squares) and detrital dolomite/limestone 'rock' (grey rhombs) and soil '*terra rossa'* (yellow rhombs) end-members. Mixing lines between the detrital end-members are also shown. The mixing lines are used to calculate the relative authigenic vs. detrital fraction of each sample, visualized by the vertical vector line for one sample in A). Plot C) is a zoom in from B) in the lower [Mo] range.

1046 **Figure 6**. Depth profiles of authigenic [Mo] and [U] as well as the authigenic  $\delta^{98}$ Mo and  $\delta^{238}$ U for Malo Jezero sediments for cores C1 (blue triangles), C2 (green circles) and C3 (red squares).

**1049 Figure 7.** The authigenic  $\delta^{238}$ U vs. [U] and authigenic  $\delta^{98}$ Mo vs. [Mo] for cores C1 (blue triangles), 1050 C2 (green circles) and C3 (red squares). The sediment samples from below and above 20 cm 1051 depth in cores C1 and C2, are shown as open and filled symbols, respectively. Average δ<sup>238</sup>U off-1052 sets from the seawater  $\delta^{238}U$  composition for these two groups of data, are shown in the 1053 authigenic  $\delta^{238}$ U vs. [U] plot.

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**Figure 8.** Bulk [Mo] (A) and [U] (B) vs. C<sub>org</sub> (wt %) correlation within core 1 (C<sub>org</sub> data from Lojen et al. (2010)). The data shown no clear correlations, however the data from <20 cm show a weak 1057 negative correlation between both  $[Mo] (A)$  and  $[U] (B)$  vs.  $C_{org}$  (see text for further discussion). A similar trend in the data is observed plotting authigenic, instead of bulk, Mo and U.

**Figure 9.** Plots of δ<sup>238</sup>U<sub>auth</sub> vs. 1/[U]<sub>auth</sub>(ppm), δ<sup>98</sup>Mo<sub>auth</sub> vs. 1/[Mo]<sub>auth</sub>(ppm), δ<sup>238</sup>U<sub>auth</sub> vs. 1060 [U]<sub>auth</sub>(ppm)/C<sub>org</sub> (Wt.%) and  $\delta^{98}$ Mo<sub>auth</sub> vs. [Mo]<sub>auth</sub>(ppm)/ C<sub>org</sub> (Wt.%). Mixing relationships between different sources of authigenic Mo and U accumulations may be expressed as linear correlations in the diagrams (see main text for further discussion).

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**Figure 10.** U vs Mo enrichment factors (A) and authigenic  $\delta^{98}$ Mo vs  $\delta^{238}$ U (B) for Malo Jezero sediments compared to the average estimates of other modern euxinic basins (from Bura-Nakić et al. 2018 and references therein). The sediment samples from below and above 20 cm depth in cores C1 and C2 are shown as open and filled symbols, respectively. The grey squares represent average estimates for core C1 below (grey square with red cross) and above (grey square with blue cross) 20 cm depth, in (A) and (B). The colored circles represent sediment data from other restricted basins as labelled in diagram B. The arrows within diagram B indicates low Mo/U versus high Mo/U uptake (from Bura-Nakić et al. 2018).

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1073 **Figure 11.** Depth profiles of pore-water H2S (orange), Mo (blue) and Fe (blue) and Mn (orange) 1074 concentrations for core C1. Depth profiles of sedimentary  $\delta^{98}$ Mo<sub>auth</sub> and [Mo]<sub>auth</sub> are also shown. 1075 In parallel is a simplified model for  $\delta^{98}$ Mo<sub>auth</sub> accumulation divided into the three zones: *i*) zone of 1076 Mn-oxide and Fe-oxyhydroxides dissolution; *ii*) zone of sedimentary Mo uptake from pore-waters 1077 and *iii*) zone of previous Mo uptake from within euxinic bottom-waters. Within zone *i*) the mixing 1078 of Mo sources with the different isotopic compositions may lead to low  $\Delta^{98}$ Mo. Zone ii) is 1079 characterized with shallow zone of sedimentary Mo uptake in the form of MoO<sub>4</sub><sup>2-</sup> accompanied 1080 with minor  $\Delta^{98}$ Mo and a deeper zone of the sedimentary Mo uptake accompanied with the more 1081 expressed  $\Delta^{98}$ Mo, likely due to the non-quantitative transformation of MoO<sub>4</sub><sup>2-</sup> to the MoS<sub>4</sub><sup>2-</sup>. Zone 1082 *iii*) is characterized with Mo uptake from within a euxinic water column at the time of deposition, 1083 accompanied with  $\Delta^{98}$ Mo of ~-0.7‰ compared to seawater (see text for further discussion).

























 















1113 Table 2. Sedimentary Mo and U authigenic isotopic composition (δ<sup>98</sup>Mo<sub>auth</sub> and δ<sup>238</sup>U<sub>auth</sub>) and 1114 authigenic concentration ([Mo]<sub>auth</sub> and [U]<sub>auth</sub>) in Malo Jezero sediments within cores C1, C2 and C3.



1118 \*The detrital fraction of [Mo]<sub>bulk</sub> too dominant to calculate an authigenic fraction.

1120 **Supplementary Table 1.** H2S concentrations in Malo Jezero water column from 1951 to 1961 1121 (Data taken from Buljan and Špan, 1976).





1125 **Table 2.** Vertical distribution of trace elements (Al, Ti, Li, Sr, Mo, U and V) and CaCO<sub>3</sub> content in 1126 Malo Jezero sediments within cores C1, C2 and C3.



1128 Repeated measurements of BCR–2 gave reproducibilities better than ± 10% (1 S.D.) and mean values

1129 within ± 10% of certified concentrations

1132 **Supplementary Table 3.** Al, Ti, Li, Sr, Mo, V and U concentrations in the catchment Malo Jezero 1133 samples.

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1136 Repeated measurements of BCR–2 gave reproducibilities better than  $\pm$  10% (1 S.D.) and mean values within  $\pm$  10%<br>1137 of certified concentrations of certified concentrations

1138

 **Supplementary Table 4.** Pore-water Mo, U, Fe and Mn concentrations were taken from Sondi et 1141 al. (2017) while the porewater  $[H_2S]$  was obtained within this study as described within the 'methods' section.



1146 **Supplementary Table 5.** Dissolved Mo and U concentrations and isotope compositinos in the 1147 Malo Jezero water column (0, 12 and 25 m depth).

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1156 **Supplementary Table 6**. Sedimentary lithogenic Mo and U fractions ([Mo]<sub>lith</sub> and [U]<sub>lith</sub> – estimated 1157 using linear regression equations presented in Figure 3, Molith/Al and Ulith/Al ratios and estimated 1158 Mo and U enrichment factors (Mo<sub>EF</sub> and U<sub>EF</sub>).

