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1	Prepared for Geochim. Cosmochim. Acta
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3	Zn stable isotope fractionation during adsorption onto todorokite: A molecular perspective from
4	X-ray absorption spectroscopy and density functional theory
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Abstract

Mineral-solution interface reactions control the mobility and fate of trace metals (e.g., Zn) and may 30 31 drive their associated isotopic fractionation in the natural environment, but understanding the coupling between interfacial reactions and isotopic behavior requires a molecular-level understanding of these 32 33 processes. In this study, we investigate Zn stable isotope fractionation during adsorption to todorokite 34 as a function of reaction time, pH and Zn concentrations. We show that Zn stable isotope fractionation reaches equilibrium at ~12 h reaction time at pH 6 and Zn concentrations of 0.05 mM. For Zn 35 36 concentrations of 0.05 and 0.2 mM, Zn isotopic fractionation between adsorbed and aqueous Zn $(\Delta^{66}Zn_{adsorbed-aqueous})$ is approximately $-0.1 \pm 0.04\%$ at pH 3-5, and $\Delta^{66}Zn_{adsorbed-aqueous}$ gradually 37 increases from $-0.1 \pm 0.04\%$ to $0.05 \pm 0.05\%$ at higher pH 6-8. Extended X-ray absorption fine 38 structure (EXAFS) spectroscopy shows that Zn adsorbs to the todorokite surface as an outer-sphere 39 octahedral complex with an average Zn–O interatomic distance of 2.06 Å at pH 3. In contrast, Zn is 40 41 predominantly present as a tetrahedral coordinated structure with a shorter average Zn–O interatomic distance of ~2.00-2.05 Å at pH 6 and 8, suggesting the presence of a mixture of octahedral outer-sphere 42 and tetrahedral inner-sphere surface complexes. Density functional theory calculations suggest that 43 44 outer-sphere surface complexes exist in the center of the structural tunnels of todorokite, yielding a theoretical Zn isotopic fractionation (Δ^{66} Zn_{adsorbed-aqueous}) of -0.2‰ to -0.3‰, whereas the tetrahedral 45 inner-sphere surface complex results in a large Δ^{66} Zn_{adsorbed-aqueous} of +0.5‰ to +0.8‰. Combined 46 47 laboratory experiments and theoretical calculations demonstrate that different magnitudes of Zn isotopic fractionation are controlled by structural changes (e.g., coordination and bond distance) in the 48 Zn surface complexes formed on todorokite relative to its aqueous form (i.e., aqua $Zn(H_2O)_6^{2+}$). These 49 results provide important new constraints for understanding Zn isotope signatures in natural Mn-rich 50 51 sediments and lead to a more complete understanding of Zn isotopes in the ocean.

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Keywords: Mn oxide; zinc; adsorption; sorption; stable isotope; fractionation; surface precipitation;
EXAFS; DFT

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1. Introduction

Metals such as zinc (Zn) play a dual role in the Earth system. On the one hand, they are important 58 59 micronutrients required by organisms for their life processes (Sinoir et al., 2012; Moore et al., 2013). In the oceans, for example, metals support photosynthesis and thus the drawdown of carbon dioxide, so 60 their concentrations in seawater are critical for regulating climate (Morel et al., 1994). On the other 61 hand, at elevated concentrations, these same metals are toxic, and their release through human activities 62 causes severe pollution (Sandstead, 2014). Studying the biogeochemical cycling of metals and how 63 64 these cycles control metal concentrations is therefore critically important to understand the role and fate 65 of metals in the Earth system.

Excitingly stable metal isotopes provide an original perspective for tracking metal cycling because 66 stable metal isotope signatures can be used as a "fingerprint" to pinpoint specific metal sources and 67 processes that control metal mobility and fate (Wiederhold, 2015). In particular, Zn isotopes have the 68 potential to shed light on the oceanic biogeochemical cycling of Zn (Conway and John, 2014; Little et 69 al., 2014; Conway and John, 2015; John et al., 2014; Little et al., 2016; Vance et al., 2016; Sweere et 70 al., 2018). In the Zn oceanic cycle, the Zn isotopic composition of the dissolved pool (~0.5% in δ^{66} Zn_{JMC}) 71 72 is substantially heavier than the best estimates of the Zn isotopic composition of the input fluxes to the ocean ($\leq 0.3\%$ in δ^{66} Zn_{JMC}), such as rivers, dust, hydrothermal fluids and benthic fluxes (Conway and 73 John, 2014; Little et al., 2014; Little et al., 2016; Liu et al., 2019). Marine oxic sediments (e.g., 74 manganese oxides and carbonates) scavenge isotopically heavy Zn ($\sim 0.9\%$ in δ^{66} Zn) (Pichat et al., 2003; 75 76 Little et al., 2014; Bryan et al., 2015; Dong et al., 2016), but organic-rich continental margin sediments bury light Zn (Little et al., 2016). The processes that control the mobility and fate of Zn and how these 77 78 processes determine its associated isotopic behavior and isotopic composition in the oceans, however, are poorly understood (Little et al., 2016; Vance et al., 2016; Isson et al., 2018). The lack of knowledge 79 80 on Zn uptake, retention and biogeochemical behavior in marine sediments and how this impacts Zn 81 isotopic signatures means that the application of Zn isotopes to constrain Zn mass balance, required for an improved understanding of Zn cycling, is speculative. Investigating Zn uptake, retention and isotope 82 fractionation during adsorption to marine minerals can shed light on these knowledge gaps. 83

84 Significant isotopic fractionations are observed during mineral-solution interface reactions in a 85 variety of aqueous environments (including marine sediments and terrestrial soils, rivers, aquifers and

wetlands) (Wasylenki et al., 2014; Wiederhold, 2015; Vance et al., 2016). Detailed information on Zn 86 isotopic fractionation during its adsorption to a variety of mineral surfaces is summarized in Table 1 87 88 (Pokrovsky et al., 2005; Balistrieri et al., 2008; Juillot et al., 2008; Bryan et al., 2015; Dong and Wasylenky, 2016; Guinoiseau et al., 2016; Nelson et al., 2017; Gou et al., 2018). The first investigation 89 of Zn isotope fractionation during adsorption to minerals reports that heavy Zn isotopes are enriched on 90 91 pyrolusite, hematite, gibbsite and corundum surfaces, while light Zn isotopes preferentially adsorb to birnessite and goethite (Pokrovsky et al., 2005). Since this study, however, Zn isotopic fractionation 92 93 during adsorption to birnessite has been found to be significantly influenced by solution ionic strength and Zn surface loading (Bryan et al., 2015). At low ionic strength, there is a small kinetic effect 94 $(\Delta^{66}Zn_{adsorbed-aqueous} \sim -0.2\%)$ for experimental durations up to 48 hours, but after 100 hours, 95 fractionations are indistinguishable from zero (Δ^{66} Zn_{adsorbed-aqueous} = 0.05 ± 0.08‰). At high ionic 96 97 strength, heavy isotopes are always preferentially adsorbed, but there is a strong dependence on surface loading, with Δ^{66} Zn_{adsorbed-aqueous} = 2.74‰ at low surface loadings, which is reduced to 0.16‰ for high 98 99 surface loadings. In addition, an enrichment of heavy isotopes is also reported at the surface of goethite $(\Delta^{66}Zn_{adsorbed-aqueous} = 0.29 \pm 0.07\%)$ (Juillot et al., 2008), which contrasts with the first reported results 100 $(\Delta^{66}Zn_{adsorbed-aqueous} = -0.18$ to -0.28%) (Pokrovsky et al., 2005). More investigations now report 101 distinct Zn isotopic fractionation behaviors in different mineral systems (e.g., calcite, kaolinite, silicon 102 oxides and aluminum oxide) (Dong and Wasylenky, 2016; Guinoiseau et al., 2016; Nelson et al., 2017; 103 104 Gou et al., 2018). This complicated fractionation behavior means that successful application of the stable isotope fingerprinting technique to track the source and fate of Zn in the oceanic environment 105 relies on a robust understanding of its isotopic fractionation during adsorption under different chemical 106 107 conditions and to different types of mineral surfaces.

To understand Zn isotopic fractionation, it is first important to distinguish which type of fractionation dominates during interface reactions occurring in the oceanic environment. Generally, isotopic fractionations can be divided into mass-dependent (MDF) and mass-independent (MIF) fractionation, where MDF includes equilibrium fractionation, Rayleigh fractionation and kinetic fractionation (Rosman and Tatlor, 1998; Moynier et al., 2017). Zn isotopic fractionation during interface reactions has been reported to obey the MDF trend (Moynier et al., 2017). According to classical stable isotope theory, several physiochemical factors (including temperature, pressure, chemical and solution 115 composition, and chemical bond stiffness) should then affect this isotopic fractionation (Bigeleisen and Mayer, 1947; Schauble, 2004). For interface reactions occurring at low temperature, temperature and 116 117 pressure are well constrained, and chemical composition and solution properties can be experimentally controlled and explored. This means that chemical bond stiffness should be the dominant factor 118 controlling Zn isotopic fractionation at equilibrium. Chemical bond stiffness is directly related to the 119 120 local chemical structure and thus the adsorption mechanism. Previous work with Zn (Gou et al., 2018), thallium (Peacock and Moon, 2012) and cerium (Nadaka et al., 2017) demonstrates that the application 121 of extended X-ray absorption fine structure (EXAFS) spectroscopy can clearly determine the local 122 chemical structure of metals associated with mineral surfaces and thus determine metal adsorption 123 mechanisms, and these studies then relate these adsorption mechanisms to isotopic behavior to constrain 124 the relationship between interface reactions and isotopic fractionation. In turn, this approach can be 125 used to provide a good understanding of the origin and evolution of metal isotope signatures in the 126 127 environment.

Studies reporting the application of EXAFS spectroscopy to elucidate Zn isotopic fractionation 128 mechanisms, however, are scarce. This is due to the complexity of isotopic and EXAFS measurements 129 130 and the high elemental concentration required for high-quality EXAFS spectra, which generally exceeds the workable concentration necessary for isotopic analysis. Researchers have therefore mostly measured 131 EXAFS spectra and isotopic fractionation separately under different experimental conditions, which 132 133 can lead to a mismatch between EXAFS spectra and isotopic data that obscures the relationship between adsorption mechanisms and isotopic fractionation behavior (Manceau et al., 2002; Bryan et al., 2015; 134 Wang et al., 2018). To date, only a limited number of studies have conducted EXAFS characterization 135 and Zn isotope measurements under the same chemical conditions (Juillot et al., 2008; Nelson et al., 136 2017; Gou et al., 2018). According to these studies, for Zn adsorption onto 2-line ferrihydrite, the 137 138 change in R_{Zn-O} values between aqueous (~2.06 Å) and adsorbed Zn (1.96 Å) explains the large isotopic fractionation (Δ^{66} Zn_{adsorbed-aqueous} = 0.53‰) (Juillot et al., 2008). In the case of goethite, no significant 139 R_{Zn-O} differences between aqueous Zn and adsorbed Zn are consistent with the small isotopic 140 fractionation ($\Delta^{66/64}$ Zn_{adsorbed-aducous} = 0.29‰) (Juillot et al., 2008). In the Zn- γ -Al₂O₃ system, both the 141 R_{Zn-Q} and the coordination number are proposed to control isotopic fractionation (Gou et al., 2018). In 142 silica systems, despite different coordination numbers, the isotopic fractionation factors for octahedral 143

and tetrahedral Zn complexes versus aqueous Zn are not distinguishable beyond uncertainties, which is attributed to the low energetic difference between octahedral and tetrahedral Zn on quartz and amorphous silica surfaces (Nelson et al., 2017). Moreover, the combined application of EXAFS spectroscopy and isotopic fractionation measurements under the same chemical conditions has not been used to constrain the relationship between interface reactions and isotopic fractionation for Zn adsorption on todorokite.

Todorokite is one of the main Mn-bearing phases in oxic marine sediments (Burns and Burns, 150 1977). It is commonly accepted that todorokite only forms from the transformation of birnessite 151 precursors (Burns and Burns, 1977, Feng et al., 2004; Bodei et al., 2007; Atkins et al., 2014). Birnessite 152 often has trace metals (e.g., Zn) adsorbed on its surface and isomorphically substituted into its structure, 153 which can reach 10⁶ times over their concentrations in seawater (Arrhenius, 1963; Koschinsky and Hein, 154 2003). Specifically, with regard to Zn, natural ferromanganese oxides in oxic marine sediments provide 155 the dominant sinks for Zn in the modern marine Zn cycle (Little et al., 2013; Vance et al., 2016). These 156 minerals typically contain between $\sim 531 \mu g \cdot g^{-1} - 1845 \mu g \cdot g^{-1}$ Zn, exclusively sequestered by the Mn-157 rich part (Koschinsky and Halbach, 1995; Koschinsky and Hein, 2003), as a result of an adsorption 158 159 equilibrium between Zn and birnessite. When todorokite forms from birnessite, however, the fate of Zn adsorbed by birnessite is likely to be ultimately controlled by its adsorption to todorokite (Atkins et al., 160 2016; Little et al., 2020). This interaction between Zn and todorokite thus potentially plays an important 161 162 role in controlling Zn isotope compositions in seawater. Todorokite has a 3D tunnel structure consisting of triple chains of edge-sharing Mn⁴⁺-O₆ octahedra that share corners, and Mn⁴⁺ substitution by cations 163 of lower valence (e.g., Mn³⁺, Ni²⁺, Co³⁺, or Cu²⁺; Post, 1999; Atkins et al., 2014; Wu et al., 2019; 164 Wegorzewski et al., 2020) creates a deficit of charge. The deficit of surface charge may be balanced by 165 exchangeable cations, such as Tl, forming outer-sphere complexes in the todorokite tunnel structure 166 (Wick et al., 2019) or by the formation of inner-sphere complexes on the todorokite surface. Therefore, 167 todorokite is likely to adsorb Zn via different mechanisms and thus fractionate Zn isotopes in a variety 168 of different ways. 169

170 In this study, we focus on the adsorption and isotopic fractionation of Zn with todorokite. 171 Specifically, we investigate the behavior of Zn isotopes during adsorption to todorokite under various 172 pH and Zn loading conditions and employ EXAFS spectroscopy to reveal the molecular-level

adsorption mechanism of Zn (i.e., coordination number, bond distances). Importantly, the structural 173 information provided by EXAFS sheds light on the nature of the Zn-mineral bond stiffness that we then 174 175 use to interpret the metal stable isotope fractionation signatures at equilibrium. To augment the experimental results, we also use a computational approach based on density functional theory (DFT) 176 to simulate metal adsorption mechanisms and calculate the values of reduced partition function ratios 177 (as 1000ln\beta) for Zn bound to todorokite. Our results provide a new in-depth understanding of the 178 coupled adsorption-fractionation behavior of Zn, which is necessary to better constrain the application 179 180 of Zn isotope signatures to track the oceanic and global Zn cycle.

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2. Materials and Methods

184 **2.1. Todorokite synthesis and characterization**

185 Todorokite was synthesized following the procedures described by Feng et al. (2004). First, 250 ml of 5.5 M NaOH solution (refrigerated for 5 h at 4 °C) was added quickly to 200 ml of 0.5 M MnCl₂ 186 solution. The mixed solution was then stirred vigorously for 5 h and aerated with O₂ at a rate of 2 L 187 188 min⁻¹. The resulting precipitate was washed with deionized (DI) water until the conductivity reached below 2 μ S·cm⁻¹. Then, the synthetic birnessite, which remained as a wet paste, was dispersed in 1 M 189 MgCl₂ solution, stirred for 12 h at room temperature, and then washed three times with DI water. The 190 191 resulting Mg-buserite was resuspended in a triangle flask connected to a condensation device and heated to reflux at 100 °C under stirring. After being refluxed for 24 h, the suspension was cooled to room 192 temperature, washed until the conductivity of the supernatant reached below $2 \mu S \cdot cm^{-1}$, and then freeze-193 dried for later use. The Zn impurity in synthetic todorokite was negligible, as determined by inductively 194 coupled plasma-mass spectrometry (ICP-MS) analysis (Agilent 7700). 195

196The synthetic todorokite was characterized using a powder X-ray diffractometer (XRD) (Bruker,197D8 ADVANCE) with Cu Kα radiation (λ = 0.15418 nm). The X-ray diffractometer was operated at 40198kV and 40 mA, with a 2θ range from 5° to 60° with a step size of 0.02° and a scan speed of 1 s at each199step. Similar to 10 Å phyllomanganate (i.e., Mg-buserite), todorokite has a basal d-spacing of 1 nm200(Burns et al., 1983, 1985; Bodei et al., 2007; Feng et al., 2010; Atkins et al., 2014); however, Mg-buserite201is not stable and can transform to 0.7 nm birnessite when dehydrated. For the identification of todorokite,

the synthetic sample was heated for 12 h at 140 °C before XRD analysis to exclude the presence of Mgbuserite.

For transmission electron microscopy (TEM) imaging, the sample was gently ground to powder and dispersed in absolute alcohol via sonication. Then, the sample was loaded on a holey carbon TEM grid. Imaging was performed at an accelerating voltage of 200 kV on an FEI Tecnai (F20) microscope. The specific surface area of the synthetic todorokite is 209 ± 4 (2SD, n = 5) m² g⁻¹, as measured by the N₂ Brunauer-Emmet-Teller (BET) method.

209

210 2.2. Adsorption experiments

All reagents used in this study were of reagent grade. Distilled DI water (>18.2 M Ω) was used for all experiments. The Zn stock solution (35 mM) used in this study was prepared from Zn(NO₃)₂·6H₂O (CAS: 10196-18-6, Sigma–Aldrich) and DI water. The acids used in all experiments were purified via double subboiling distillation. The labware used in the adsorption and isotopic experiments was made of Teflon material to minimize Zn contamination.

Two types of Zn adsorption experiments were carried out under ambient environmental conditions: 216 217 (i) kinetic adsorption experiments (duration of 0.5-120 h) and (ii) adsorption edge experiments from pH 3 to 9. For the kinetic adsorption experiments, suspensions of 0.4 g L⁻¹ todorokite were equilibrated 218 at pH 6 for 24 h in 0.1 M NaNO₃ solution while constantly stirring. After this pre-equilibration step, the 219 220 necessary amount of Zn from the $Zn(NO_3)_2$ stock solution was added to achieve an initial Zn concentration of 0.05 mM. After a reaction time of 0.5 to 120 h, the solid and liquid phases were 221 separated by centrifugation. The supernatants were passed through a methacrylate butadiene styrene 222 filtration assembly with a polyethersulfone (PES) membrane and 0.22 µm pores (Sartorius, Germany) 223 224 and analyzed for Zn using inductively coupled plasma-optical emission spectroscopy (Thermo ICP 225 6000 series ICP-OES).

For the adsorption edge experiments, prior to Zn addition, todorokite powder was suspended for 227 24 h at a fixed pH (3 to 9) at three different ionic strengths (I = 0.001, 0.01 and 0.1 M NaNO₃). Then, 228 small amounts of $Zn(NO_3)_2$ stock solution were added dropwise to todorokite suspensions to achieve 229 initial Zn concentrations of 0.05 and 0.2 mM. These initial Zn concentrations were selected because the 230 concentrations of Zn adsorbed on todorokite (0.002 < Zn/Mn atomic ratio < 0.052) covered the range

of Zn contents found in marine Mn-rich sediments such as ferromanganese crusts and nodules (0.001 <231 Zn/Mn atomic ratio < 0.007) (Manheim and Lane-Bostwick, 1991; Little et al., 2014). While adding Zn, 232 the suspension was stirred vigorously to avoid the formation of Zn precipitates due to local 233 oversaturation of the suspension. The solid/liquid ratio was set equal to 0.4 g L^{-1} , and the pH of all 234 adsorption experiments was adjusted to the desired value by the addition of 0.01 M NaOH/HNO₃. Based 235 236 on the kinetic experiments, the reaction time for adsorption edge experiments was set to 48 h to ensure equilibrium. After 48 h, the solid and liquid phases were separated by centrifugation, and the supernatant 237 238 was further filtered through a methacrylate butadiene styrene filtration assembly with a polyethersulfone (PES) membrane and 0.22 µm pores (Sartorius, Germany). The Zn concentration in 239 the supernatants was measured by inductively coupled plasma-optical emission spectroscopy (Thermo 240 ICP 6000 series ICP-OES). The dominant Zn species in the adsorption experiments was calculated 241 using Visual MINTEQ.3.1 and found to be Zn^{2+} across the pH range 3 – 9 (Fig. S1). The Zn surface 242 coverage was calculated from the difference in concentrations between the initial and equilibrium 243 solutions by taking into account the surface area of todorokite and the solid/liquid ratio. Blank 244 experiments with no Zn added to the reacting suspension of todorokite were performed to monitor the 245 246 external Zn contamination during adsorption experiments. All the experimental data are averages of duplicates, and the relative errors are less than 5%. 247

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2.3. Zn stable isotope measurements

Isotopic samples were obtained after the batch adsorption experiments. Solid samples separated 250 by centrifugation were dissolved in 11 M HCl and heated at 100 °C in sealed Teflon vials for 3 h to 251 extract the adsorbed Zn. The amount of aqueous Zn remaining in the solids after centrifugation was less 252 than 1‰ of the adsorbed Zn, according to the estimation of the Na content extracted by this step and 253 254 the Na/Zn ratio of the supernatant (Gou et al., 2018). The digested solutions from the solids and the 255 supernatants obtained after various adsorption experiments were evaporated to dryness and redissolved three times in 11 M HCl to convert Zn to a chloride form. After that, each residue was dissolved in a 2 256 M HCl solution and loaded on columns filled with AG MP-1 anion-exchange resin (100–200 mesh; 257 258 Bio-Rad, USA). Samples were purified following the procedure described by Gou et al. (2018). Column yields were measured after Zn purification and determined to be $100 \pm 6\%$. Total procedural blanks of 259

260 Zn were analyzed for each experimental run and were found to be less than 20 ng, which was negligible 261 (< 0.1%) compared to the ~20 μ g Zn in each sample. After purification, the samples were evaporated 262 to dryness and dissolved in 0.05 M HNO₃. Aliquots of varying volumes of this Zn solution were sampled 263 to obtain a final 0.05 M HNO₃ solution with 0.5 ppm Zn and 0.2 ppm Cu NIST 647. Cu NIST 647 and 264 Zn (high purity in-house standard) mixed standard solutions with equivalent Cu/Zn ratios of samples 265 were also prepared to run alternatively with the samples to correct the instrumental mass bias (Marechal 266 et al., 1999).

267 The Zn isotopic ratio was measured using a Neptune Plus (Thermo Fisher Scientific) MC-ICP-MS at the Isotope Geochemistry Lab of Nanjing University. The instrument was run in "wet-plasma" 268 mode at low mass resolution (M/ Δ M = 400) using a free aspirating glass expansion nebulizer. Prior to 269 isotopic composition measurement, a 0.05 M HNO₃ solution was introduced before each isotope ratio 270 measurement to make the baseline decrease to a constant value. Then, a 40 s on-peak blank of 0.05 M 271 HNO₃ was measured for background subtraction. Each measurement consisted of three blocks of 60 272 cycles (60 s for each cycle). The signal of ⁶²Ni was collected to correct for isobaric interference on ⁶⁴Zn 273 from ⁶⁴Ni, while the impact on the quality of the ⁶⁴Zn MC-ICP-MS signal was found to be negligible 274 275 due to effective chromatography.

276 The Zn isotopic compositions of all samples were reported relative to the in-house standard as:

277
$$\delta^{66} Zn = (({}^{66} Zn / {}^{64} Zn)_{sample} / ({}^{66} Zn / {}^{64} Zn)_{HPS}) - 1) \times 1000 (\%)$$
(1)

The in-house standard was calibrated with respect to IRMM 3702 and JMC Lyon international standards, 278 giving δ^{66} Zn_{HPS}=-0.36 ± 0.03‰ and δ^{66} Zn_{HPS}=-0.09 ± 0.03‰, respectively (Gou et al., 2018). The 279 long-term external reproducibility of the ^{66/64}Zn ratio measurement by MC-ICP-MS was better than 280 0.03‰ (2 SD) based on repeat analysis of a standard solution and geological standards over a period of 281 3 years. The measured δ^{66} Zn values of two reference materials (i.e., Nod-P-1, BCR) are consistent with 282 those reported in the literature, attesting to the accuracy of the analytical procedure (Marechal et al., 283 1999; Bryan et al., 2015; Gou et al., 2018). The isotopic fractionation of Zn (Δ^{66} Zn_{adsorbed-aqueous}) between 284 the solid phase and aqueous phase is defined as: 285

286
$$\Delta^{66} Zn_{adsorbed-aqueous} = \delta^{66} Zn_{adsorbed} - \delta^{66} Zn_{aqueous}$$
(2)

287 The mass balance offset was calculated as follows:

288 Offset = $\delta^{66}Zn_{stock} - (\delta^{66}Zn_{adsorbed} \times \% \text{ adsorbed} + \delta^{66}Zn_{aqueous}(100 - \% \text{ adsorbed}))$ (3)

290 **2.4. EXAFS sample preparation and data analysis**

EXAFS samples were prepared in the same manner as the isotopic experiments, with a large vessel to yield at least 20 mg of solid for analysis. The reaction conditions, including the solid/liquid ratio, initial Zn concentration, ionic strength, and pH, were kept the same as those of the corresponding isotopic samples. The samples for EXAFS analysis were prepared as fresh wet pastes and stored in a refrigerator prior to EXAFS analysis.

296 The Zn K-edge EXAFS data for adsorption and standard samples were collected at the 1W1B beamline of Beijing Synchrotron Radiation Facility (BSRF, China) and BL12C in Photon Factory (PF) 297 synchrotron-radiation facility within the National Laboratory for High Energy Physics (KEK), Tsukuba, 298 Japan. The electron storage ring operated at 2.5 GeV with an average beam current of 200 mA in BSRF 299 300 and at 2.5 GeV with a maximum beam current of 450 mA in PF. A pair of Si(111) crystals were employed as a monochromator, which was detuned by 30% to reject X-ray harmonics. The spectra of 301 all samples were measured by both transmission and fluorescence mode simultaneously, and the 302 fluorescence data were recorded using a multichannel Ge detector. The X-ray energy was calibrated 303 304 with a Zn metal foil, and the first inflection point in the Zn K-edge was set to 9659 eV. An Fe metal filter was used to dampen the fluorescence of Mn. Because of the fluorescence of Mn, a minimum of 305 three scans were collected for each sample to obtain a good signal/noise ratio. Concentrated samples 306 307 and standards were analyzed in transmission mode.

308 The Zn EXAFS spectra were processed and analyzed using an IFEFFIT 1.2.11 program package for data normalization and Fourier transformation (Ravel and Kelly, 2007). The $\chi(k)$ spectra were k^3 -309 weighted and Fourier transformed (FT) over the k range $3-12 \text{ Å}^{-1}$. Shell-by-shell fitting was performed 310 in R-space to obtain detailed local structural information. Theoretical backscattering-amplitude and 311 312 phase-shift values were calculated based on the crystal structures of Zn(NO₃)₂ and todorokite model clusters. During sample spectra processing, the coordination number N, the distance R, and the Debye-313 Waller factor σ^2 were allowed to float as adjustable parameters, and a single energy shift ΔE_0 was used 314 for all shells. The number of adjustable parameters in the fitting procedures did not exceed the maximum 315 number of independent parameters. The quality of each fit was evaluated quantitatively with the reduced 316 317 chi square and R-factor parameters. The amplitude reduction factor (S_0^2) was estimated to be 0.97 based

318 on the fitting of a $Zn(NO_3)_2$ solution standard and was then applied to the adsorption samples.

319

320 **2.5. Density functional theory calculations**

321 Initial Zn-todorokite models were created by expansion of the experimental unit-cell structure of todorokite mineral (Post et al., 2003) to a $1 \times 2 \times 1$ supercell, which contained one or two Zn cations 322 323 per eight H₂O molecules in each tunnel space (ZnMn₁₂O₂₄·8H₂O or Zn₂Mn₁₂O₂₄·8H₂O). Water molecules were initially placed at the corner sites in the tunnel according to X-ray structure refinement 324 results (Post et al., 2003), while Zn cations were placed at the center or corner sites in the tunnel. Mn(III) 325 cations are believed to occupy the larger corner-sharing octahedra of todorokite (Post et al., 2003). In 326 the current models, two or four corner-sharing Mn octahedra sites were imposed by the electron 327 configuration of Mn(III), with the other sites imposed by Mn(IV). After relaxation of the structures, the 328 magnetic moments of Mn(IV) and Mn(III) sites were 2.9–3.1 and 3.7–3.8 $\mu_{\rm B}$. For aqueous Zn²⁺, we 329 constructed a cluster model of Zn^{2+} surrounded by one shell or two shells of H₂O molecules [$Zn^{2+}.6(H_2O)$] 330 or $Zn^{2+} \cdot 18(H_2O)$] in a 30 × 30 × 30 Å periodic box based on the symmetry reported in Ducher et al. 331 (2018). 332

333 All DFT calculations were performed using the CASTEP code (Clark et al., 2005) under the spinpolarized generalized gradient approximation (GGA) with the Perdew-Burke-Ernzerhof (PBE) 334 functional (Perdew et al., 1996). The valence electron states of the Mn, Zn, O, and H ultrasoft 335 pseudopotentials (Vanderbilt, 1990) were $3s^23p^63d^54s^2$, $3d^{10}4s^2$, $2s^22p^4$, and $1s^1$, respectively. The 336 kinetic energy cutoff was 500 eV. The k-point grid for the first Brillouin zone (Monkhorst and Pack, 337 1976) was $1 \times 1 \times 1$ for aqueous Zn^{2+} models and $4 \times 10 \times 4$ for Zn-todorokite models. The calculated 338 atomic force converged to less than 0.01 eV/Å. The magnetic ordering among edge-sharing Mn 339 octahedra was ferromagnetic, but the ordering between the corner-sharing of the triple chains of edge-340 sharing octahedra was antiferromagnetic (Kaltak et al., 2019). Geometry optimizations were performed 341 with no symmetry imposed until the following conditions were met: total energy ≤ 0.000005 eV, atomic 342 forces $\leq 0.05 \text{ eV/Å}$ ($\leq 0.005 \text{ eV/Å}$ for aqueous Zn^{2+} models), stress $\leq 0.03 \text{ GPa}$ and atomic 343 displacement ≤ 0.0005 Å. All atomic coordinates were fully relaxed, while $\alpha = \gamma = 90^{\circ}$ was fixed in 344 todorokite models, and the supercell size was fixed in aqueous Zn²⁺ models. The crystal structures were 345 rendered using VESTA software (Momma and Izumi, 2011). 346

347 The 66 Zn/ 64 Zn reduced partition function ratios (β -factors) of Zn-bearing todorokite were 348 calculated with harmonic vibrational frequencies based on the high-temperature product rule 349 (Bigeleisen and Mayer, 1947; Schauble, 2004; Blanchard et al., 2017):

350
$$\beta = \left[\prod_{i=1}^{3N_{at}} \prod_{q} \frac{\mathbf{v}_{q,i}^{*}}{\mathbf{v}_{q,i}} \frac{e^{-h\mathbf{v}_{q,i}^{*}/(2kT)}}{1 - e^{-h\mathbf{v}_{q,i}^{*}/(kT)}} \frac{1 - e^{-h\mathbf{v}_{q,i}/(kT)}}{e^{-h\mathbf{v}_{q,i}/(2kT)}}\right]^{1/(N_{q}N)}$$
(4)

where v is the harmonic vibrational frequency of the i^{th} vibrational mode at a phonon wave vector q; h, 351 *k*_{*B*}, and *T* are the Planck constant, Boltzmann constant, and absolute temperature in Kelvin, respectively; 352 $N_{\rm at}$, $N_{\rm q}$, and N represent the number of atoms in a calculated model, the number of phonon wave vectors, 353 and the number of sites of isotopes, respectively; and * represents a frequency for a heavier isotope (⁶⁶Zn 354 in this study). The harmonic vibrational frequencies were obtained only at the gamma point by phonon 355 calculations based on a finite displacement method (Ackland et al. 1997; Parlinski et al. 1997), which 356 is implemented in CASTEP; in Eq. (4), N_q was 1, and the ith vibrational mode started from 4. The 357 ⁶⁶Zn/⁶⁴Zn equilibrium isotopic fractionation factor between Zn-adsorbed todorokite and aqueous Zn²⁺, 358 Δ^{66} Zn_{adsorb-aq}, was obtained by combining the corresponding β -factors of the solid phase and aqueous 359 Zn^{2+} . 360

361

362

363 **3. Results**

364 **3.1 Characterization of synthetic todorokite**

The XRD pattern of the synthetic todorokite sample is shown in Fig. 1a. It shows well-resolved 365 diffraction peaks at ~9.6 Å (001), ~4.8 Å (002), 3.2 Å (003), ~2.5 Å (210), ~2.4 Å (21-1), ~2.2 Å (21-366 2), ~1.9 Å (31-2), ~1.7 Å (21-4), ~1.5 Å (21-5) and ~1.4 Å (020). These peaks are consistent with 367 those of typical todorokite (JCPDS 38-0475) and naturally formed todorokites (Post et al., 2003; Feng 368 et al., 2004; Bodei et al., 2007; Manceau et al., 2007; Atkins et al., 2014; Wu et al., 2019). Fig. 1b and 369 c show the morphology of synthetic todorokite by TEM observation. The particles of synthetic 370 todorokite exhibit fibrous laths aligned with each other at 120° to form large aggregates (Fig. 1b). From 371 the lattice image of the synthetic todorokite (Fig. 1c), we observe that the todorokite fibers elongate 372 longitudinally along the (010) direction. Ideal todorokite exhibits lattice fringes with fringe widths 373 (along the (001) direction) of \sim 1 nm corresponding to a 3 × 3 MnO₆ octahedra tunnel size. However, 374

tunnels of different MnO_6 octahedral fringe widths along the (001) direction have been observed (Fig. 1c). This indicates that in the trilling intergrowths, the tunnel size along the (001) direction is variable. 376 377 Such morphologies (fibrous and trilling patterns) and intergrowths are the typical characteristics of natural todorokite, consistent with previously reported synthetic and natural todorokites (Feng et al., 378 2004; Bodei et al., 2007; Manceau et al., 2007; Atkins et al., 2014; Wu et al., 2019). 379

380

3.2 Zn adsorption onto todorokite 381

382 Fig. 2a shows Zn adsorption ($[Zn]_{initial} = 0.05 \text{ mM}$) as a function of pH for three different ionic strengths (0.005, 0.01, 0.1 M NaNO₃). A substantial increase in Zn adsorption from 0.11 µmol m⁻² to 383 0.60 µmol m⁻² occurs over the pH range from 3 to 9. Clearly, there is an ionic strength effect, with the 384 highest Zn adsorption in 0.005 M NaNO3 solution over most of the pH range. Similar evolution is 385 observed when studying Zn adsorption from solution as a function of pH for an initial Zn concentration 386 of 0.2 mM (Fig. 2b). A substantial increase in Zn adsorption from 0.36 µmol m⁻² to 2.39 µmol m⁻² occurs 387 over the pH range from 3 to 9. Zn adsorption significantly varies with ionic strength between 0.005 and 388 0.1 M NaNO₃, with the highest Zn adsorption in 0.005 M NaNO₃ solution. Such ionic strength effects 389 390 on Zn adsorption to other mineral surfaces have also been reported (Guinoiseau et al., 2016; Nelson et al., 2017). This ionic strength-dependent behavior of Zn adsorption suggests that the formation of outer-391 sphere surface complexes is the predominant adsorption mechanism, especially at pH 3-7 (Strawn and 392 Sparks, 1999; Elzinga et al., 2002). Since Na⁺ ions in the background electrolyte can compete with Zn 393 adsorption for these relatively weak adsorption sites, less Zn will be adsorbed when the ionic strength 394 395 increases. To confirm that outer-sphere Zn adsorption complexes formed on todorokite, EXAFS 396 analyses were performed (see section 3.3).

The results of time series experiments of Zn adsorption to todorokite are presented in Fig. 3a. Zn 397 398 adsorption is initially fast, with 30% of total Zn removed from solution in the first 15 min. Thereafter, 399 Zn adsorption slows slightly and remains almost constant from 24 to 120 h. After 120 h of reaction time, approximately 50% of total Zn is adsorbed, resulting in a surface coverage of 0.29 µmol m⁻². These 400 results suggest that the adsorption equilibrium between todorokite and the reacting solution is achieved 401 402 after 24 h of reaction time.

403

404 **3.3 Zn isotope fractionation during the adsorption process**

The Zn isotopic compositions of the adsorbed and aqueous phases in the time series experiments 405 are plotted in Fig. 3b and tabulated in Table 2. The initial isotopic composition (δ^{66} Zn) of the Zn stock 406 solution is $0.28 \pm 0.06\%$ (2SD, n = 6). In the time series experiments, light Zn isotopes are preferentially 407 partitioned to the solid phase, and heavy isotopes remain in solution. During the first 12 h, the δ^{66} Zn 408 values of the aqueous phase show a rapid decrease from $0.38\% \pm 0.06$ to $0.28 \pm 0.04\%$, while 409 simultaneously, the δ^{66} Zn values of adsorbed Zn increase from 0.23 ± 0.05‰ to 0.29 ± 0.01‰. From 410 24 h to 120 h, no significant changes in the δ^{66} Zn values of the aqueous and adsorbed phases are 411 observed. The Zn isotope fractionation between adsorbed and aqueous phases (Δ^{66} Zn_{adsorbed-aqueous}) 412 evolves during the first 12 h from $-0.15 \pm 0.08\%$ to $-0.06 \pm 0.04\%$ and remains nearly unchanged for 413 longer reaction times (Fig. 3c and Table 2). This suggests that isotopic equilibrium between adsorbed 414 and aqueous Zn was achieved after ~ 12 h of reaction. 415

416 The pH series experiments were conducted at different initial Zn concentrations of 0.05 and 0.2 mM, and the results are presented in Fig. 4 and Table 2. In the experiments of [Zn]_{initial} = 0.05 mM, light 417 Zn isotopes are preferentially partitioned to the solid phase relative to the aqueous phase (Fig. 4a). The 418 419 Zn isotopic compositions of the adsorbed phase remain almost constant at approximately $0.2 \pm 0.02\%$ at pH 3 to 5 and increase from $0.26 \pm 0.01\%$ to $0.29 \pm 0.01\%$ when pH values increase from 6 to 9. 420 Aqueous compositions are almost identical within analytical uncertainties over the pH range from 3 to 421 422 8. It should be noted that the Zn isotope composition of the aqueous phase at pH 9 is absent because aqueous Zn remaining in solution is below the detection limit owing to the nearly 100% Zn uptake 423 under this condition. Overall, Zn isotope fractionation between the adsorbed and aqueous phases is 424 slightly negative for the initial Zn concentration of 0.05 mM, and the average Δ^{66} Zn_{adsorbed-aqueous} for all 425 12 samples is $-0.07 \pm 0.04\%$ (Fig. 4b). 426

In the experiments of $[Zn]_{initial} = 0.2 \text{ mM}$, Zn isotope fractionation behavior can be divided into two pH groups (Fig. 4c). From pH 3 to 6, the Zn isotope compositions and evolution of the adsorbed and aqueous phases are similar to those in the experiments with $[Zn]_{initial} = 0.05 \text{ mM}$. A continual enrichment in light isotopes in the adsorbed phase and heavy isotopes in the solution is observed. At pH 7 to 8, heavy Zn isotopes are preferentially partitioned onto the adsorbed phase relative to the aqueous solution. The enrichment of heavy isotopes on todorokite, which is not observed in the

experiments of $[Zn]_{initial} = 0.05$ mM, seems to imply that relatively high Zn concentrations may induce 433 different Zn isotope fractionation. The Zn isotopic fractionation value (Δ^{66} Zn_{adsorbed-aqueous}) is calculated 434 to be $-0.08 \pm 0.03\%$ in the pH 3 to 6 experiments, while the $\Delta^{66}Zn_{adsorbed-aqueous}$ value increases to 0.05 435 $\pm 0.05\%$ in the pH 7 to 8 experiments (Fig. 4d). Mass balance calculations are applied for almost all 436 samples to assess the success of the experiments and exclude the possibility of external contamination. 437 The mass balance offset is calculated according to the equation (δ^{66} Zn_{aqueous} (1-% adsorbed) 438 $+\delta^{66}$ Zn_{adsorbed} % adsorbed) $-\delta^{66}$ Zn_{stock}. These values are within $\pm 0.05\%$ of the Zn isotope composition 439 440 of the starting stock solution (Table 2), indicating that the experimental data are reliable.

441

442 **3.4 Zn surface speciation as revealed from EXAFS**

Fig. 5a shows the background subtracted k^3 -weighted EXAFS spectra (χ function) for the Zn-443 todorokite samples and three Zn reference compounds (e.g., Zn solution, Zn hydroxide (Zn(OH)₂), and 444 hydrozincite $(Zn_3(OH)_6(CO_3)_2)$). The EXAFS spectra of the Zn-adsorbed todorokite samples show a 445 distinct change with pH. The samples prepared at pH 3 with Zn concentrations of 0.05 and 0.2 mM 446 yield similar EXAFS oscillations, which are almost identical to the EXAFS of the Zn(NO₃)₂ solution. 447 448 Correspondingly, Fig. 5b shows the Fourier transformed spectra (uncorrected for phase shift) for the Zn-todorokite samples, where only a single peak is observed. This peak results from O backscattering 449 in the first coordination shell of the Zn atoms. The fit results for the Zn-todorokite samples and the 450 451 reference compounds are presented in Table 3, resulting in bond distances of ~2.06 Å and coordination numbers of 5.8–6.2. Fitting of the Zn(NO₃)₂ solution EXAFS data results in a bond distance of 2.06 Å 452 and coordination number of 6, which are in good agreement with many previous studies (Bochatay and 453 Persson, 2000; Roberts et al., 2003; Nelson et al., 2017; Gou et al., 2018). At pH 3, the similarity 454 between the Zn adsorption structure and aqueous Zn aqua cation provides strong evidence for the 455 formation of an outer-sphere surface complex, in agreement with the marked ionic-strength dependent 456 adsorption behavior observed at pH 3 (Fig. 2). 457

458 Compared to the pH 3 samples, the samples prepared at pH 6 and 8 with initial Zn concentrations 459 of 0.05 and 0.2 mM exhibit slightly different features at higher k values between 7 and 8.5 Å⁻¹, 460 suggesting a change in the mechanism of surface complexation (Fig. 5a). In the Fourier transform, the 461 first shell O atoms are dominated by a major peak that shifts to lower values. This indicates that the 462 bond length between Zn and O (R_{Zn-O}) in these samples is shorter than that of Zn in solution. Further 463 shells at higher R are observed, indicating the presence of backscatterer second shell atoms residing at 464 longer distances, such as Zn and/or Mn. The presence of backscatterer second shell atoms suggests that 465 Zn either forms covalent bonds with todorokite surface sites or precipitates. Shell-by-shell fits for these samples exhibit EXAFS contributions from the first shell Zn-O scattering as well as second shell 466 467 Zn–Zn/Mn scatterers. The fit of the first shell yields $R_{Zn-O} = 2.00-2.05$ Å and coordination numbers (CN) ranging from 4.8–5.5. The bond distances for the O shell at $R_{Zn-O} = 2.00-2.05$ Å are similar to 468 those found for the hydrozincite $(Zn_5(CO_3)_2(OH)_6)$ sample $(R_{Zn-O} \sim 2.00 \text{ Å}, \text{ see Table 4})$, where Zn 469 forms a mixture of octahedrally and tetrahedrally coordinated complexes. Bochatay and Persson (2000) 470 found that Zn surface complexes adsorbed on manganite had Zn–O bond distances of approximately 471 472 1.98 to 2.04 Å and that these distances are consistent with the formation of a mixture of octahedral and 473 tetrahedral complexes. Similar observations for Zn adsorption on other mineral surfaces have also been 474 reported (Waychunas et al., 2002; Cismasu et al., 2013; Nelson et al., 2017; Gou et al., 2018). Therefore, for the Zn adsorbed todorokite samples prepared at pH 6 and 8, the Zn–O bond distances between 2.00-475 476 2.05 Å are indicative of a mixture of octahedrally and tetrahedrally coordinated structures.

477 To fit the second shell for the pH 6 and 8 samples, we tested both Zn–Zn and Zn–Mn paths. Because of the similar backscatter power between Zn and Mn, either Zn–Zn or Zn–Mn paths can fit the 478 second shell, yielding an interatomic distance of 3.10–3.14 Å and CN of 1.5–3.0. Due to the high 479 480 uncertainty in the higher shell coordination number in fitting EXAFS spectra (Ravel and Kelly, 2007), bond distance was used to assess the local structure of Zn on todorokite. If a Zn-Zn path is assumed, 481 the Zn–Zn distance is 3.10-3.14 Å, which is similar to that of hydrozincite (Zn₅(CO₃)₂(OH)₆), where a 482 distance of 3.12 Å is fitted, and much shorter than the Zn–Zn bond distance (3.23 Å) in Zn hydroxide 483 (Zn(OH)₂) (Table 3). However, precipitated or polymerized Zn as either hydrozincite or Zn hydroxide 484 at pH 6 is highly unlikely (Fig. S1) because of the low pH (below neutral) and low Zn concentration 485 (Bochatay and Persson, 2000). Thus, Mn backscattering seems more reasonable, and a good fit with a 486 Zn–Mn distance of 3.10–3.14 Å (Table 3) suggests the formation of an inner-sphere surface complex 487 488 structure.

489

490 **3.5 Theoretical calculation of the stable isotope fractionation**

In the DFT geometry optimizations, the coordination structure of Zn cations in the tunnel strongly 491 depends on the cation content and position (Table 4), while the lattice parameters of Zn todorokite 492 493 models show minimal changes depending on the cation content. The calculated lattice parameters (a =10.1, b = 2.88, c = 9.67 Å, $\beta = 95.6^{\circ}$) differ by less than 1.2–3.5% from the experimental results (a =494 9.76, b = 2.84, c = 9.56 Å, $\beta = 94.2^{\circ}$; Feng et al., 2004). The atomic coordinates of the geometry-495 optimized structures are provided in Table S1. In ZnMn₁₂O₂₄·8H₂O models, Zn forms octohedral 496 coordination with six H₂O molecules (i.e., outer-sphere surface complexes) at the center of the tunnel 497 (model 1); at the corners, Zn forms tetrahedral coordination with two H₂O molecules and two surface 498 O molecules (i.e., inner-sphere surface complexes; model 2). The d < Zn-O > of octahedral Zn is larger499 than that of tetrahedral Zn (2.09 vs. 1.96 Å). In Zn₂Mn₁₂O₂₄·8H₂O models, two hydrated Zn cations 500 501 cannot occupy the center sites at the same time because of limited space. If one Zn occupies the center site, the other Zn should occupy the corner site (model 3); otherwise, both Zn cations occupy the corner 502 503 sites (model 4) to minimize repulsion among tunnel cations (Fig. 6). In model 3, we find octahedral Zn at the corner site, coordinated with one H2O and five surface O (i.e., octahedral inner-sphere surface 504 complexes), at d < Zn-O = 2.17 Å (Table 4). The presence of octahedral Zn at the corner site also 505 506 increases the d<Zn-O> of the outer-sphere complex of Zn at the center from 2.09 to 2.14 Å. In model 4, both Zn cations form tetrahedral coordination at the corners, with a slight increase in d < Zn-O > from 507 1.96 to 1.98 Å. Based on the current DFT calculations, octohedrally coordinated Zn shows a wider 508 range of d<Zn-O> from 2.09 to 2.14 Å than tetrahedrally coordinated Zn (1.96–1.98 Å). Based on the 509 geometry-optimized structures, the beta factors $(10^3 \ln\beta^{66} Zn^{/64} Zn)$ calculated at 300 K are 2.8–3.3‰ 510 for octahedral Zn and 3.5–3.9‰ for tetrahedral Zn (Fig. 6). 511

The d < Zn-O> of $Zn^{2+}.6(H_2O)$, a simple model for aqueous Zn^{2+} , is calculated to be 2.12 Å (Table 512 4). This distance is slightly overestimated by 2.9% compared to the distance of aqueous Zn^{2+} obtained 513 by the current EXAFS analysis (2.06 Å; Table 3). On the other hand, when a larger cluster model 514 including double hydration shells, $Zn^{2+} \cdot 18(H_2O)$, is geometry optimized, d < Zn-O > is calculated to be 515 2.17 Å. This distance is consistent with the result of a recent DFT study, in which DFT-based MD 516 simulations of a periodic $Zn^{2+}.64(H_2O)$ model predict 2.17 Å for d < Zn-O > for aqueous Zn^{2+} (Ducher 517 et al., 2018). However, the DFT method with more realistic aqueous Zn^{2+} models used in the current 518 study and Ducher et al. (2018) predict d < Zn-O > to be even longer than the distance of the simple 519

520 $Zn^{2+} \cdot 6(H_2O)$ cluster model (~5% error compared to the experimental value). Because the level of 521 accuracy of DFT in prediction of the bond distances of cations differs between aqueous and solid phases, 522 careful interpretation is required for direct comparison of the *d*<Zn-O> for the solid and aqueous phases 523 and hence the beta factors between them. When the beta factor of $Zn^{2+} \cdot 6(H_2O)$ is used for aqueous Zn^{2+} , 524 the ⁶⁶Zn/⁶⁴Zn equilibrium isotopic fractionation factor between adsorbed Zn and aqueous Zn^{2+} ($\Delta^{66}Zn$) 525 is -0.2 to -0.3‰ for octahedral Zn cations and +0.5 to +0.8‰ for tetrahedral Zn in todorokite.

- 526
- 527 528

4. Discussion

529 4.1 Constraining Zn isotope fractionation factors

In the experiments at pH values of 3.0 to 6.0 and Zn concentrations of 0.05-0.02 mM, both δ^{66} Zn_{adsorbed} and δ^{66} Zn_{aqueous} showed a linear increase as the proportion of adsorbed Zn increased (Fig. 7a). Equilibrium and Rayleigh fractionation models are used to fit our experimental data. In the equilibrium fractionation model, adsorbed Zn can still isotopically exchange with aqueous Zn irrespective of the degree of adsorption. In the Rayleigh fractionation model, Zn is separated from the aqueous solution to the solid phase during adsorption, and adsorbed and aqueous Zn do not isotopically equilibrate with one another. The two models are described based on the following equations:

537 Equilibrium model:
$$\delta^{66}Zn_{aqueous} = \frac{\delta^{66}Zn_{stock} - 1000 \times f \times (\alpha_{adsorbed-aqueous} - 1)}{1 - f + (f \times \alpha_{adsorbed-aqueous})}$$
 (5)

538 Rayleigh model:
$$\delta^{66}Zn_{aqueous} = (1000 + \delta^{66}Zn_{stock})(1-f)^{(\alpha_{adsorbed-aqueous}-1)} - 1000$$
 (6)

where f is the fraction of Zn sorbed, and $\alpha_{adsorbed-aqueous}$ is the estimated isotopic fractionation factor between adsorbed and aqueous Zn. The δ^{66} Zn_{stock} is the measured value (0.28 ± 0.06‰) of the initial aqueous Zn. An $\alpha_{adsorbed-aqueous}$ of 0.9999 is obtained from linear regression of the experimental data. The δ^{66} Zn_{aqueous} and $\alpha_{adsorbed-aqueous}$ values are used to predict the corresponding δ^{66} Zn_{adsorbed} on the todorokite surface according to Eq. (3).

544
$$\delta^{66}Zn_{adsorbed} = \alpha_{adsorbed-aqueous} \left(\delta^{66}Zn_{aqueous} + 1000 \right) - 1000$$
 (7)

Fig. 7a shows that the equilibrium fractionation model fits our experimental data better than the Rayleigh model, indicating adequate isotope exchange between the aqueous solution and the todorokite surface throughout the adsorption reactions. This equilibrium fractionation during adsorption reactions is in full agreement with that reported in studies for Zn and other metal adsorption (e.g., Cd, Li, Cu,

- Mo) on mineral surfaces (Barling and Anbar, 2004; Wasylenky et al., 2008; Wasylenky et al., 2014;
 Gou et al., 2018).
- 551 Finally, the isotopic fractionation between adsorbed and aqueous Zn (Δ^{66} Zn_{adsorbed-aqueous}) is 552 obtained from isotopic fractionation factors ($\alpha_{adsorbed-aqueous}$) following Eq. (4).

553
$$\Delta^{66}Zn_{adsorbed-aqueous} \cong 1000 \times ln\alpha_{adsorbed-aqueous}$$
 (8)

554 This calculation yields a Δ^{66} Zn_{adsorbed-aqueous} value equal to -0.10 ± 0.04 %, consistent with the average

555 $(-0.11 \pm 0.04\%)$ from the measured $\Delta^{66}Zn_{adsorbed-aqueous}$ value.

556 In the adsorption experiments at pH values of 7.0 to 8.0 and Zn concentrations of 0.05–0.2 mM, Zn isotope compositions of solid and aqueous phases do not change within analytical error with respect 557 to the fraction of adsorbed Zn (Fig. 7b). This relatively constant fractionation between solid and aqueous 558 phases indicates achievement of isotopic equilibrium. This result is in agreement with our time series 559 experiments, which show that Zn isotope fractionation during adsorption on todorokite reaches 560 equilibrium at approximately 12 h (see section 3.3). Previous studies also suggest that equilibrium of 561 Zn isotopes is achieved within a few hours of adsorption to mineral surfaces of 2-line ferrihydrite, 562 calcite, quartz and amorphous silica (Juillot et al., 2008; Dong et al., 2016; Nelson et al., 2017). 563

To summarize, different magnitudes for Zn isotope fractionation during adsorption to todorokite under different pH values are observed. The Δ^{66} Zn_{adsorbed-aqueous} values of -0.10 ± 0.04 ‰ and 0.05 ± 0.05 ‰ are calculated for pH 3.0 to 6.0 and pH 7.0 to 8.0, respectively. Such differences under different pH values are related to different Zn surface speciation, as detailed in the following section.

568

569 **4.2 Effect of Zn speciation on Zn isotope fractionation**

570 The distinct Zn isotope fractionation behaviors under different pH values (Fig. 7) might be 571 explained by two main processes: i) isotopic fractionation arises during the exchange of isotopes among 572 different Zn species in solution; ii) fractionation occurs between dissolved and adsorbed Zn species at 573 the mineral-solution interface.

Regarding the first process, aqueous Zn speciation for the adsorption edge experiments with Zn concentrations of 0.05 and 0.2 mM was calculated using Visual MINTEQ 3.1 (Fig. S1). Calculations show that $Zn(H_2O)_6^{2+}$ is the main species (91.5%) with only a minor amount (8.5%) of $ZnNO_3(H_2O)_5^+$ at pH < 8 (the major pH range used in the adsorption experiments). At low ionic strength (e.g., 0.001

M NaNO₃), the proportion of $ZnNO_3(H_2O)_5^+$ is even smaller. A number of studies based on ab initio 578 calculations report the values of reduced partition function ratios (as 1000lnß) for various Zn species in 579 solution (Pye et al., 2006; Fujii et al., 2010; Fujii et al., 2014). Zn-Cl complexes, for example, are 580 reported to incorporate lighter isotopes, with 1000 lnß values ranging from 2.293-3.136 as the number 581 of Cl⁻ ligands in the complex increases, compared with 3.263 for $Zn(H_2O)_6^{2+}$ (Fujii et al., 2010). The 582 magnitudes of fractionation from $Zn(H_2O)_6^{2+}$ for chloro complexes translate to values of $\Delta^{66}Zn_{hexaquo}$ 583 _{chloro} ranging from 0.13‰ for ZnCl(H₂O)₅⁺ to 1.40‰ for ZnCl₄²⁻. Fujii et al. (2014) also estimated the 584 $\ln\beta$ of carbonate complexes of Zn to be 3.877 for ZnHCO₃⁻ and 4.199 for ZnCO₃⁰ and predicted that 585 carbonate complexes should be associated with a relatively heavy pool of Zn in aqueous solution. 586 Similarly, $ZnSO_4^{0}$, which comprises a heavy pool of Zn by 0.3% relative to $Zn(H_2O)_6^{2+}$, has been 587 reported (Fujii et al., 2014). Thus, the formation of ZnNO₃(H₂O)₅⁺ may cause isotopic fractionation 588 from $Zn(H_2O)_6^{2+}$. While the 1000 lnß value for $ZnNO_3(H_2O)_5^+$ is currently unknown, the fractionation 589 factors for Zn-ligand complexes may be predicted according to a positive correlation between the 590 complexation constant (logK) and isotopic fractionation (Ryan et al., 2014; Markovic et al., 2017). 591 Based on the small log K (0.4) and low proportion (8.5%) of $ZnNO_3(H_2O)_5^+$ under the experimental 592 593 conditions (Allison et al., 1991), we surmise that the formation of $ZnNO_3(H_2O)_5^+$ does not significantly 594 contribute to the measured Zn isotope fractionation. Further studies on the calculation of this species 595 would be useful to test this hypothesis.

596 Regarding the second process, adsorption of aqueous species to mineral surfaces is often accompanied by a change in the chemical bond stiffness of the element of interest, which stems from a 597 different molecular structure (surface speciation) of Zn and would thus cause isotopic fractionation. As 598 599 Zn is naturally present only as divalent Zn and is insensitive to redox, the coordination number is proposed to be an important factor in controlling the vibrational frequencies (i.e., stiffness) of Zn-600 oxygen bonds (Schauble, 2004). Thus, coordination number changes have been used to explain Zn 601 602 isotope fractionations during adsorption, where essentially heavy Zn isotopes are found to be more likely enriched in structures with low coordination numbers in which Zn has stiffer bonds (Schauble, 603 2004). In this study, outer-sphere Zn complexes at low pH values (e.g., pH 3) on todorokite concentrate 604 slightly lighter Zn compared to the aqueous phase (Δ^{66} Zn_{adsorbed-aqueous} = $\sim -0.1 \pm 0.04\%$) (Fig. 8), 605 although the structure of Zn in aqueous solution and Zn in an outer-sphere complex on todorokite are 606

very similar, with Zn surrounded by approximately six water molecules. According to DFT calculations, 607 outer-sphere octahedral Zn complexes show a range of Zn–O bond distances and 10³lnβ ⁶⁶Zn/⁶⁴Zn 608 depending on the tunnel composition, which could lead to the slightly negative isotopic fractionation 609 observed (Table 4; Fig. 6). We note that while Zn^{2+} structures in aqueous solution and todorokite tunnels 610 are dynamic and highly disordered, the fractionation factors reported in the current study are based on 611 certain end-member structures; more accurate values could be obtained by exhaustive sampling for 612 disordered Zn²⁺ configurations via MD simulations. At higher pH values from 6 to 8, a mixture of outer-613 614 sphere octahedral and inner-sphere tetrahedral Zn occurs, and Zn isotopic compositions of the adsorbed phase become heavier than the aqueous phase as pH increases (Fig. 4c). EXAFS is unable to discern a 615 change in coordination numbers when there is a mix of two similarly coordinated complexes (i.e., four 616 617 coordination and six coordination), and while there is likely to be a small decrease in the average coordination number of the combined complexes (Table 3), this cannot be resolved with EXAFS 618 because the uncertainty in the coordination numbers is at least ± 1 (Ravel and Kelly, 2007). However, a 619 decrease in the coordination number does occur at high pH values due to the formation of tetrahedral 620 Zn with lower coordination numbers, and this decrease is likely responsible for the slightly positive 621 fractionation (Δ^{66} Zn_{adsorbed-aqueous} ~ 0.05 ± 0.05‰) (Fig. 8). On the other hand, interatomic distances have 622 also often been used to rationalize isotopic fractionation trends (Guinoiseau et al., 2016; Nelson et al., 623 2017), where essentially heavy Zn isotopes are found to be more likely enriched in structures with low 624 625 interatomic distances in which Zn has stiffer bonds (Schauble, 2004). However, it should be noted that coordination is often correlated to bond distances that a structure with lower coordination numbers 626 would be associated with shorter bond length (Juillot et al., 2008). According to a comparison of our 627 calculated vibrational frequencies (Fig. S3), todorokite with tetrahedral Zn tends to exhibit higher 628 vibrational frequencies related to Zn-O bonds than todorokite with octahedral Zn, although H₂O 629 630 vibrational modes are also involved in the frequencies. The current DFT calculation predicts that the equilibrium fractionation factor between inner-sphere tetrahedral Zn on todorokite and aqueous Zn 631 ranges from 0.5-0.8 ‰ at 300 K (Table 4), similar to the experimental fractionation values for 632 633 tetrahedral Zn adsorbed on the surface of silica (Nelson et al., 2017), ferrihydrite (Juillot et al., 2008), 634 Al oxide (Gou et al., 2018), and kaolinite (Guinoiseau et al., 2016). This does indicate that tetrahedral Zn on todorokite concentrates heavy isotopes, and thus, the presence of tetrahedral Zn at high pH values 635

(e.g., pH 8) could explain our experimental observations that Zn isotopic fractionation (Δ^{66} Zn_{adsorbed}-636 _{aqueous}) evolves from $-0.1 \pm 0.04\%$ at pH 3 with only outer-sphere octahedral complexes to $0.05 \pm 0.05\%$ 637 at pH 8 with a mixture of outer-sphere octahedral complexes and inner-sphere tetrahedral complexes. 638 Overall, the isotopic fractionation of Zn during adsorption to todorokite is driven by Zn speciation 639 changes after adsorption versus its aqueous form before adsorption. The formation of outer-sphere 640 octahedral surface complexes that have a long Zn-O distance close to the structure of the aqueous 641 $Zn(H_2O)_6^{2+}$ cluster results in insignificant Zn isotope fractionation, whereas the formation of inner-642 sphere tetrahedral surface complexes yields a shorter Zn-O distance than the aqueous $Zn(H_2O)_6^{2+}$ cluster 643 enriches heavier isotopes (Fig. 6). 644

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646 **4.3 Implications for understanding Zn isotope signatures in Mn-rich marine sediments**

To date, previous work has measured the Zn isotope signatures of natural Fe-Mn crusts and nodules, 647 which contain birnessite-type Mn oxides, and the relationship between Zn adsorption and isotopic 648 fractionation for birnessite in controlled experiments. Work shows that natural Fe-Mn crusts and 649 nodules are enriched in heavy Zn isotopes relative to seawater (~0.5‰) (Marechal et al., 2000; Bermin 650 651 et al., 2006; Conway et al., 2013; Little et al., 2014a; Zhao et al., 2014), while birnessite at high ionic strength (i.e., synthetic seawater solutions), which is analogous to birnessite in Fe-Mn deposits, is also 652 enriched in heavy Zn isotopes (Bryan et al., 2015). The surface layers of Fe-Mn nodules from different 653 654 oceanic areas are also enriched in heavy Zn isotopes relative to seawater but have variable Zn isotope compositions, ranging from 0.53% to 1.16% (Marechal et al., 2000), while the Zn isotope compositions 655 of a depth profile from three Fe-Mn crusts also vary between 0.8‰ and 1.2‰ (Little et al., 2014a). In 656 the experimental work above (Bryan et al., 2015), a Zn isotopic fractionation (Δ^{66} Zn_{adsorbed-solution}) of 657 2.74‰ is reported at low surface loadings, which is reduced to 0.16‰ for high surface loadings. The 658 659 observations that Fe-Mn deposits concentrate heavy Zn isotopes with a range of isotopic compositions 660 (Marechal et al., 2000; Little et al., 2014a) therefore appear to be explained by the presence of birnessite in natural Fe-Mn crusts and nodules with variable surface loadings of adsorbed Zn. To understand how 661 Zn adsorption drives Zn isotopic fractionation, previous work also employs X-ray fluorescence 662 mapping of Fe-Mn crusts to confirm that Zn is predominantly associated with birnessite and EXAFS 663 spectroscopy to show that Zn exists in tetrahedral coordination in these deposits (Little et al., 2014b). 664

The enrichment of Fe-Mn crusts in heavier Zn isotopes and of oxic marine sediments that contain birnessite-type Mn oxides is thus attributed to the adsorption of Zn via tetrahedral complexes at the birnessite surface, which enrich heavy Zn isotopes (Little et al., 2014b; Bryan et al., 2015).

Todorokite is one of the main Mn-bearing phases in oxic marine sediments, where it is typically 668 intimately associated with birnessite-type minerals (Burns and Burns, 1977; Post, 1999; Peacock and 669 670 Moon, 2012). It is thought to only form as a result of the aging and associated mineralogical transformation of birnessite (Feng et al., 2004; Atkins et al., 2014), and todorokite is especially present 671 in diagenetic settings and low-temperature hydrothermal settings, where the transformation of 672 birnessite-type precursor phases is favored (Burns and Burns, 1977). For example, todorokite is rarely 673 found in hydrogenetic ferromanganese crusts precipitated directly from ambient seawater (authigenic) 674 675 where birnessite-type minerals are dominant but is found in variable proportions in diagenetic ferromanganese nodules formed at the sediment-seawater interface, reflecting the influence of sediment 676 677 pore waters on individual nodules, and it is prevalent below the sediment-seawater interface in close proximity to hydrothermal fluids (Burns and Burns, 1977; Post, 1999). In the present study, we quantify 678 679 Zn isotope fractionation during adsorption to todorokite for the first time. The results show that the 680 magnitude of Zn isotopic fractionation during adsorption to todorokite is much smaller than that measured above for Zn isotope signatures in Fe-Mn crusts and nodules (Table 2). It is true that the 681 isotopic composition of Zn adsorbed to todorokite becomes heavier from pH 3 to 8 (Fig. 4), as the Zn 682 683 complexes evolve from outer-sphere octahedral coordination to a mixture of outer-sphere octahedral and inner-sphere tetrahedral coordination, but the isotopic fractionation (Δ^{66} Zn_{adsorbed-solution} = 0.05 ± 684 0.05%) at circumneutral pH is still very small compared to the average Δ^{66} Zn_{Mn oxides-seawater} value of 685 ~0.5‰ (Marechal et al., 2000; Bermin et al., 2006; Little et al., 2014a; Zhao et al., 2014). Furthermore, 686 while our DFT calculations predict equilibrium fractionation between pure inner-sphere tetrahedral Zn 687 on todorokite and aqueous Zn to range from 0.5–0.8‰, it is unlikely that pure tetrahedral Zn exists at 688 the Zn/Mn atomic ratio (~0.004) of natural Mn-rich marine sediments (Manheim and Lane-Bostwick, 689 1991) (Fig. S2). Instead, isotopic fractionation between todorokite and solution close to natural 690 691 conditions should approach $-0.10 \pm 0.04\%$ (Fig. S2).

692 Further work is needed to determine the extent to which the Zn isotopic fractionation values 693 determined in this work are directly applicable to natural marine sediments. Our experimental

conditions of circumneutral pH are close to the pH of marine sediment porewaters (Reimers et al., 1996), 694 695 and while our experiments differ in terms of temperature and ionic strength compared to marine 696 sediment porewaters, this is not expected to substantially alter Zn adsorption or isotopic fractionation behavior. Specifically, the slightly elevated temperature used here compared to typical marine sediment 697 porewaters (~10 °C) is unlikely to substantially change Zn adsorption or isotopic fractionation behavior 698 (Fujii et al., 2010; Ducher et al., 2016). Additionally, the elevated temperature used here to more rapidly 699 induce the transformation of birnessite to todorokite than occurs in marine sediments is shown not to 700 701 result in a different transformation mechanism or pathway for birnessite to todorokite and is thus not 702 expected to alter the mechanism of Zn adsorption or isotopic fractionation (Atkins et al., 2014, 2016). The ionic strength of our experimental system is lower than that in marine sediment porewaters, but for 703 Zn inorganic speciation, the dominant species of Zn in porewaters is expected to be $Zn^{2+}(aq)$ (Byrne, 704 705 2002) and thus the same as that in our experimental system. Our experimental conditions of ionic speciation potentially differ compared to marine sediment porewaters, however, in terms of Zn organic 706 speciation. Organic ligands in marine sediments can chelate trace metals such as Zn and may induce 707 708 associated isotopic fractionation (Jouvin et al., 2009). Moreover, although the fate of Zn adsorbed by 709 birnessite is likely to be ultimately controlled by its adsorption to todorokite (Atkins et al., 2016; Little et al., 2020), the isotopic behavior of Zn during its migration from birnessite as it transforms into 710 todorokite and its subsequent adsorption to todorokite may not be completely captured in our 711 712 experiments that do not include the transformation process. Therefore, the exact isotopic compositions of Zn in marine todorokite might be hard to predict. Overall, however, we suggest that Zn isotopic 713 fractionation in natural marine todorokite can be inferred from structural differences that occur between 714 715 (higher coordination numbers, longer average Zn-O distance) outer-sphere octahedral complexes (thus light) and (lower coordination numbers, shorter average Zn-O distance) inner-sphere tetrahedral 716 717 complexes (thus heavy). As such, we can reasonably infer that relatively light Zn isotopes are enriched in natural todorokite compared to those in natural birnessite, according to the dominant outer-sphere 718 octahedral Zn complexes on todorokite versus inner-sphere tetrahedral Zn complexes on natural 719 birnessite. This means that diagenetic sediments that contain todorokite are likely to be lighter than the 720 721 Fe-Mn deposits and oxic sediments measured above, which are all either largely hydrogenetic (Fe-Mn crusts) or with limited diagenetic influence (surfaces of Fe-Mn nodules) or oxic/authigenic samples. 722

723 It is commonly accepted that todorokite only forms from the transformation of birnessite precursors (Burns and Burns, 1977, Feng et al., 2004; Bodei et al., 2007; Atkins et al., 2014). During 724 725 this transformation process, previous work shows that Ni adsorbed by birnessite is released to solution (Atkins et al., 2016), while Co is retained in the newly formed todorokite phase (Wu et al., 2019). More 726 generally, these studies propose that non-Jahn-Teller distorted cations, such as Ni, are released during 727 transformation, while Jahn-Teller distorted cations, such as Co, are structurally incorporated during 728 transformation (Atkins et al., 2014, 2016; Wu et al., 2019), and the presence of Jahn-Teller distorted 729 730 trace metals could facilitate the formation of todorokite (Bodei et al., 2007; Cui et al., 2008; Atkins et al., 2014; Zhao et al., 2015). With relevance to our work, Zn, similar to Ni, does not have Jahn-Teller 731 distortion and is thus predicted to be released to solution during the birnessite to todorokite 732 transformation (Atkins et al., 2014, 2016). Diagenetic/suboxic sediments are thus expected to be a 733 734 source of Zn to porewaters, which might potentially diffuse into the overlying water column (Aktins et al., 2014, 2016). As such, because Zn is potentially released to the water column during the 735 transformation of birnessite to todorokite and because light Zn isotopes are enriched with todorokite 736 compared to Zn with birnessite, it is therefore likely that these processes act in combination to result in 737 738 Zn isotope compositions of diagenetic/suboxic sediments that are lighter than oxic sediments that 739 contain authigenic Mn oxides or those with limited diagenetic influence.

Future work should investigate the Zn isotopic composition of natural sediments to help support the experimental implication that diagenetic/suboxic sediments are enriched in light Zn isotopes. The possibility of further Zn isotopic fractionation during mineralogical transformation and Zn release, which affects the isotopic signal of diagenetic/suboxic sediments, should also be investigated in controlled laboratory experiments. Finally, follow-up work should constrain the effect of organic ligands on Zn isotopic fractionation during adsorption to todorokite to ensure that experiments capture both the inorganic and organic speciation of Zn complexes in seawater.

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5. Conclusion

We report the first study to quantify Zn isotope fractionation during adsorption to synthetic todorokite and constrain the fractionation mechanisms for different experimental conditions. The results

show that the molecular-level characteristics of Zn adsorption complexes can explain distinct Zn isotope 752 signatures. Zn adsorbs to the todorokite surface at low pH as an outer-sphere octahedral complex (R_{Zn}-753 $_{\rm O}$ = 2.06 Å) with slightly negative isotopic fractionation from aqueous Zn (Δ^{66} Zn_{adsorbed-aqueous} = -0.1 ± 754 0.04‰). At higher pH values, Zn begins to form a mixture of outer-sphere octahedral and inner-sphere 755 tetrahedral complexes ($R_{Zn-O} = 2.00 - 2.05$ Å) at the todorokite surface with a measured equilibrium 756 isotope fractionation (Δ^{66} Zn_{adsorbed-aqueous}) ranging from -0.1 ± 0.04‰ to 0.05 ± 0.05‰. We assert that 757 structural changes (e.g., coordination and bond distance) are the cause of the differences in Zn isotope 758 partitioning to the todorokite surface because inner-sphere tetrahedral Zn formation on the todorokite 759 surface with a lower coordination number and shorter Zn-O bond distance relative to outer-sphere 760 octahedral Zn corresponds to a higher vibrational frequency and enriches heavy isotopes. Furthermore, 761 we can constrain two distinct Zn isotopic fractionation magnitudes from DFT calculations: Δ^{66} Zn_{adsorbed}-762 $_{aqueous} = -0.10 \pm 0.04\%$ for the octahedral surface complex and $\Delta^{66}Zn_{adsorbed-aqueous} = 0.5-0.8\%$ for the 763 764 tetrahedral inner-sphere corner-sharing surface complex on todorokite.

The results reported here provide important new constraints for understanding Zn isotope 765 signatures in Mn-rich marine sediments. We propose that because light Zn isotopes are enriched in 766 767 todorokite compared to birnessite, diagenetic/suboxic sediments that contain todorokite as a result of the diagenetic transformation of birnessite will have lighter Zn isotope signatures than hydrogenetic Fe-768 Mn crusts and oxic sediments. In addition, previous work shows that Zn is expected to be released from 769 770 newly formed todorokite to sediment porewater and potentially the overlying water column (Atkins et al., 2014, 2016); thus, the enrichment of light isotopes on todorokite and the release of Zn from this 771 mineral phase might act in combination to constrain the isotope signatures of diagenetic/suboxic 772 773 sediments to relatively light values. Overall, we predict that diagenetic/suboxic sediments that contain todorokite are likely enriched in lighter Zn isotopes than authigenic/oxic sediments that contain 774 birnessite (Marechal et al., 2000; Little et al., 2014a; Bryan et al. 2015). Field studies are needed to 775 support this experimentally based prediction, leading to a more complete understanding of Zn isotope 776 systematics in the ocean. 777

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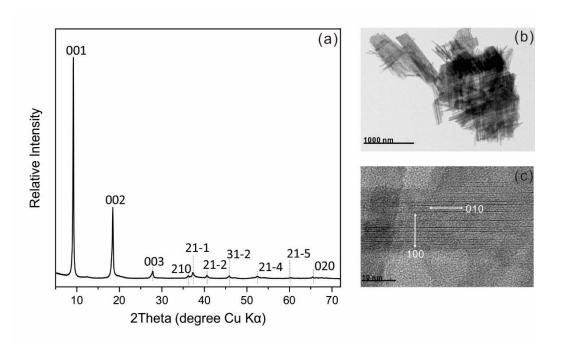
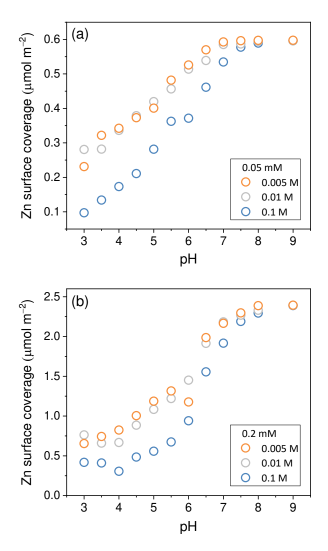


Fig. 1. Characterization of synthetic todorokite using powder X-ray diffraction (a) and transmission
electron microscopy (TEM) (b, c). X-ray diffraction peaks are labeled based on JCPDS-38-0475 for
todorokite. 010 and 100 in (c) indicate lattice plane directions.



1019Fig. 2. Effect of pH on Zn adsorption to todorokite at different ionic strengths. Adsorption experiments1020were carried out at two Zn concentrations of 0.05 mM (a) and 0.2 mM (b). The solid/solution ratio was1021set to 0.4 g/L suspension, the equilibrium time was 48 h, and room temperature was 25 ± 1 °C.

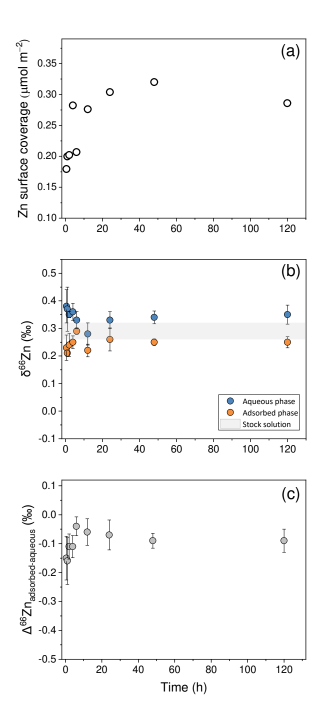
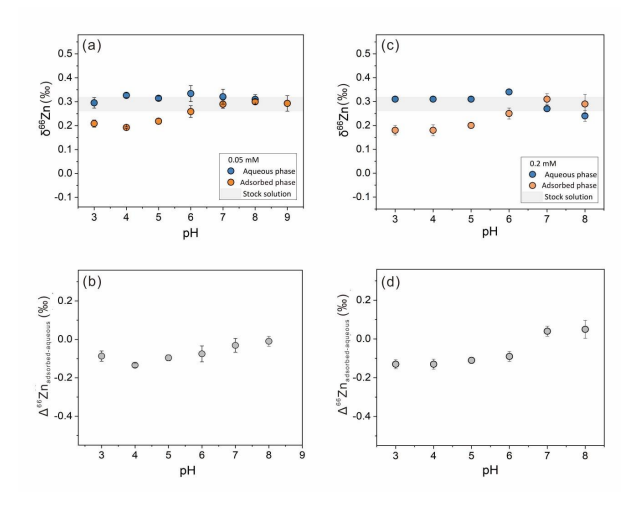


Fig. 3. Time-series experiments of (a) Zn adsorption kinetics, (b) δ^{66} Zn values for adsorbed Zn and aqueous Zn; (c) Zn stable isotopic fractionation between adsorbed Zn and aqueous Zn phases. The experiments were carried out at a Zn concentration of 0.05 mM at pH 6 in a background electrolyte of 0.1 M NaNO₃ at a solid/solution ratio of 0.4 g/L and room temperature (25 ± 1 °C). The gray shaded area indicates the initial Zn isotope composition of the stock solution used in all experiments. Error bars represent 2SD on 3 replicate analyses of each sample.

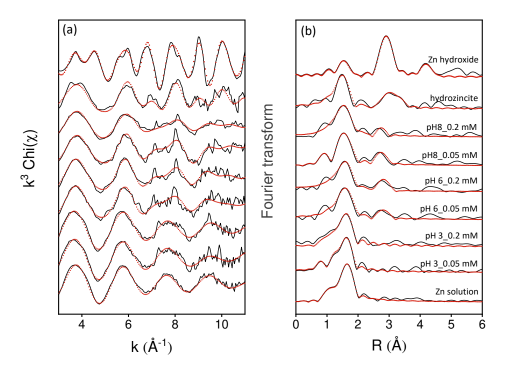
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Fig. 4. Plots of Zn isotope compositions of adsorbed Zn and aqueous Zn and Zn isotopic fractionation between the two phases in adsorption edge experiments at 0.05 mM Zn loading (a, b) and adsorption edge experiments at 0.2 mM Zn loading (c, d). The gray shaded area indicates the initial Zn isotope composition of the stock solution used in all experiments. Error bars represent 2SD on 3 replicate analyses of each sample.

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1048 Fig. 5. (a) k^3 -weighted Zn K-edge EXAFS data of Zn adsorption samples and reference compounds; (b) corresponding Fourier transforms in R space. Experimental and shell-by-shell fitting data are shown as black solid lines and red dots, respectively.

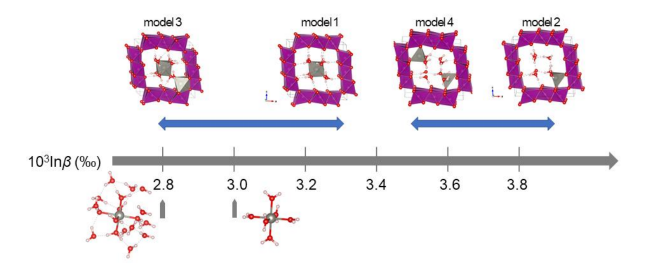




Fig. 6. A range of beta factors $(10^3 \ln\beta^{66} Zn/^{64} Zn)$ DFT-calculated at 300 K for aqueous Zn^{2+} clusters and 2056 Zn-todorokite adsorption models (models 1–4). Purple octahedron = Mn octahedron; gray polyhedra = 1057 Zn octahedra or tetrahedra; red = O; pink = H; dashed lines = H-bonds.

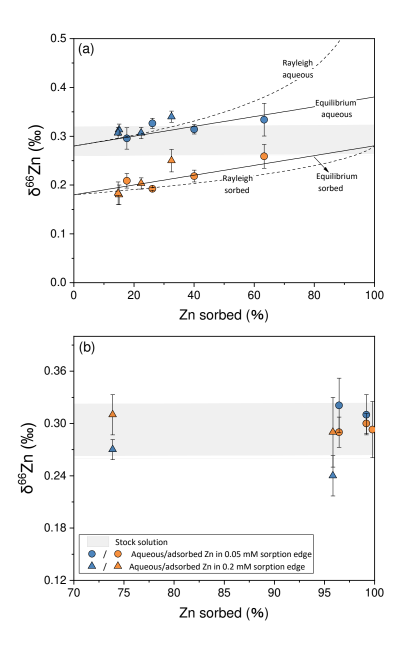
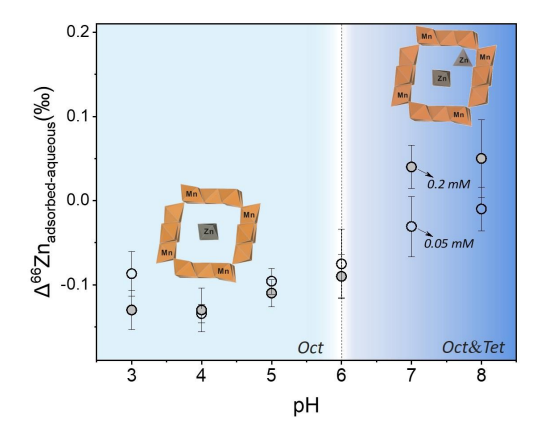


Fig. 7. Plots of Zn isotope compositions of adsorbed Zn and aqueous Zn as a function of the 1061 fraction of adsorbed Zn in experiments performed at pH values of 3.0-6.0 and Zn 1062 concentrations of 0.05-0.02 mM (a) and in experiments performed at pH values of 7.0-8.0 and 1063 1064 Zn concentrations of 0.05–0.02 mM (b). The solid and dashed lines in figure (a) represent δ^{66} Zn 1065 values for adsorbed Zn and aqueous Zn in the batch equilibrium model and the Rayleigh 1066 fractionation model, respectively. An isotopic fractionation factor, $\alpha_{adsorbed-aqueous}$, of 0.9999 is 1067 obtained based on the best-fit batch equilibrium model. The gray shaded area indicates the initial Zn isotope composition of the stock solution used in all experiments. Error bars represent 1068 1069 2SD of three replicate analyses of each sample.



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Fig. 8 Schematic diagram of the speciation of adsorbed Zn and associated isotopic fractionation
 on todorokite with respect to different pH values. The dashed line and shaded area are assigned
 based on approximate locations obtained from Zn isotope and EXAFS spectroscopic analyses.
 Oct refers to surface complexes of Zn coordinated with six oxygen atoms, and Tet refers to

1075 inner-sphere corner-sharing Zn complexes in tetrahedral coordination with four oxygen atoms.

Minerals	pН	I (M)	Surface coverage $\Gamma(\mu mol m^{-2})$	αadsorbed- aqueous	$\Delta^{66} Zn_{adorbed-aqueous}$ (‰)	Reference
Goethite	5.28-6.14	0.01	0.02- 0.03	nd	-0.20±0.03	Pokrovsky et al., 2005
	4-8	0.1	0.01-1.30	1.00029	~ 0.29	Juillot et al., 20081080
Hematite	5.50-6.61	0.01	0.02-0.07	nd	0.04 to 0.61	Pokrovsky et al., 20051
Ferrihydrite	4-8	0.1	0.17-3.21	1.00053	~ 0.53	Juillot et al., 2008 ¹⁰⁸²
Birnessite	4.10-5.65	0.01	0.03-0.08	nd	-0.17 ± 0.06	Pokrovsky et al., 2005
	~8.2	0	> 0.185	nd	$0.05{\pm}0.08$	Bryan et al., 2015 ¹⁰⁸⁴
	~8.2	0.7 g/kg	nd	nd	0.16-2.7	1085
		(synthetic seawater)				1086 1087
Pyrolusite	6.06-8.34	0.01	0.58-3.95	nd	0.10±0.03	Pokrovsky et al., 2009
Calcite	8.2-8.4	0	nd	nd	0.41±0.18	Dong and Wasylenk20161090
	8.2-8.4	0.7 g/kg (synthetic seawater)	nd	nd	0.73±0.08	1091 1092 1093
Kaolinite	<5	0.01	0.13-0.46	1.00018	$0.18{\pm}0.06$	Guinoiseau et al., 2096
	>5	0.1	0.02-0.47	1.00049	$0.49{\pm}0.06$	1095
Corundum	6.87-7.55	0.01	0.22-0.51	nd	0.10±0.09	Pokrovsky et al., 2005
Gibbsite	6.37-6.98	0.01	0.37-0.67	nd	0.13±0.12	Pokrovsky et al., 2005
γ-Al ₂ O ₃	6.0-7.5	0.1	< 0.8	nd	$0.02{\pm}0.07$	Gou et al., 2018 1098
	6.5-8.0	0.1	1.5-3.2	1.00047	0.47 ± 0.03	1099
Quartz	5.28-5.91	0.004	< 0.6	nd	-0.01 ± 0.06	Nelson et al., 20181100
	6.53-7.15	0.1	0.6-1.4	nd	0.60±0.11	1101
Amorphous silica	6.23-6.99	0.004-0.100	< 1.4	nd	0.94±0.11	Nelson et al., 20181102

Table 1. Summary of Zn isotope fractionation during adsorption onto typical environmental minerals

1104 Note. Italic surface loading values were calculated based on the data reported in related references. All uncertainties are reported using 2 sd. nd

1105 means there are not specific descriptions in references.

106		Table 2. C	Chemical and	isotopic results	s of zinc adso	rption	experime	ents in this s	tudy				
Experiments	Time (h)	[Zn] _{initial} (mM)	Adsorbed Zn(%)	Coverage (µmol m ⁻²)	Zn/Mn (mol:mol)	pН	$\delta^{66}Zn_{aq}$	ueous 2sd	$\delta^{66}Zn_{adso}$	rbed 2sd	$\Delta^{66} Zn_{adsorbed}$	2sd	Mass balance
	0.5	0.05	30.0	0.18	0.004	6	0.38	0.06	0.23	0.05	-0.15	0.08	0.05
	1	0.05	33.4	0.20	0.005	6	0.37	0.08	0.21	0.01	-0.16	0.08	0.04
	2	0.05	33.8	0.20	0.005	6	0.35	0.01	0.24	0.04	-0.11	0.04	0.04
Todorokite	4	0.05	47.2	0.28	0.006	6	0.36	0.03	0.25	0.02	-0.11	0.04	0.03
adsorption	6	0.05	34.6	0.21	0.005	6	0.33	0.03	0.29	0.01	-0.04	0.03	0.04
kinetics	12	0.05	46.2	0.28	0.006	6	0.28	0.04	0.22	0.02	-0.06	0.04	-0.03
	24	0.05	52.1	0.30	0.007	6	0.33	0.03	0.26	0.04	-0.07	0.05	0.01
	48	0.05	55.3	0.32	0.007	6	0.34	0.02	0.25	0.01	-0.09	0.02	0.01
	120	0.05	45.7	0.29	0.006	6	0.35	0.03	0.26	0.02	-0.09	0.04	0.03
	48	0.05	17.6	0.11	0.002	3	0.30	0.02	0.21	0.04	-0.09	0.03	0.00
0.05 mM	48	0.05	26.1	0.16	0.003	4	0.33	0.01	0.19	0.01	-0.13	0.01	0.01
Todorokite	48	0.05	40.1	0.24	0.005	5	0.31	0.01	0.22	0.00	-0.10	0.02	0.00
adsorption	48	0.05	63.3	0.38	0.008	6	0.33	0.03	0.26	0.01	-0.08	0.04	0.01
edges	48	0.05	96.5	0.58	0.013	7	0.32	0.03	0.29	0.02	-0.03	0.04	0.01
_	48	0.05	99.2	0.59	0.014	8	0.31	0.02	0.30	0.02	-0.01	0.03	0.02
	48	0.05	100	0.60	0.014	9			0.29	0.01			
	48	0.2	15.1	0.36	0.008	3	0.31	0.01	0.18	0.02	-0.13	0.02	-0.01
0.2 mM	48	0.2	14.7	0.35	0.008	4	0.31	0.01	0.18	0.02	-0.13	0.03	-0.01
Todorokite	48	0.2	22.4	0.54	0.012	5	0.31	0.01	0.20	0.01	-0.11	0.02	-0.01
adsorption	48	0.2	32.5	0.78	0.018	6	0.34	0.01	0.25	0.02	-0.09	0.03	-0.03
edges	48	0.2	73.9	1.77	0.040	7	0.27	0.01	0.31	0.02	0.04	0.03	-0.02
_	48	0.2	95.8	2.29	0.052	8	0.24	0.02	0.29	0.04	0.05	0.05	-0.01

Table 2. Chemical and isotopic results of zinc adsorption experiments in this study

Note. Todorokite suspension was 0.4 g/L. The ionic strength is 0.1 66 Zn_{stock} is 0.28 ± 0.06‰ (2SD, n = 6). Each sample is an average of three 1107 measurements on a Neptune with the 2 sd values reported to reflect internal precision on replicate. Mass balance offset = (δ^{66} Zn_{aqueous} (1-adsorbed %) 1108 $+\delta^{66}Zn_{adsorbed}$ adsorbed %)- $\delta^{66}Zn_{stock}$. 1109

Samples	Shell no.	Path	CN	R (Å)	$\sigma^2(\text{\AA}^2)$	R-factor	$\Delta E_0 (eV)$
Zinc hydroxide	1	Zn–O	3.7	1.98	0.006	0.0009	5.16
	2	Zn–Zn	10.4	3.23	0.009		
	3	Zn–Zn	3.8	4.62	0.004		
hydrozincite	1	Zn–O	4.5	1.98	0.009	0.0082	-5.98
	2	Zn–Zn	1.4	3.12	0.012		
	2	Zn–Zn	2.2	3.52	0.006		
Zn(NO ₃) _{2 aq}	1	Zn–O	6 ^a	2.06	0.009	0.0001	2.91
Tod pH 3_0.05	1	Zn–O	6.2	2.06	0.010	0.0010	-1.28
Tod pH 3_0.2	1	Zn–O	5.8	2.06	0.009	0.0011	-1.86
Tod pH 6_0.05	1	Zn–O	5.5	2.05	0.011	0.003	-0.08
	2	Zn–Mn	3.0	3.14	0.016		
Tod pH 6_0.2	1	Zn–O	5.4	2.04	0.011	0.002	-0.70
	2	Zn–Mn	2.7	3.13	0.013		
Tod pH 8_0.05	1	Zn–O	4.8	2.01	0.009	0.0009	-4.61
	2	Zn–Mn	1.5	3.10	0.007		
Tod pH 8_0.2	1	Zn–O	5.3	2.00	0.013	0.0013	-3.16
	2	Zn–Mn	1.9	3.12	0.018		

Table 3. Fitting Results of the EXAFS Spectrum for Model Compounds and Adsorption

1111 Samples (${}^{c}S_{0}{}^{2} = 0.97$)

Sample name of Tod pH 3_0.2 means Zn surface speciation on todorokite at pH 3 and initial Zn concentration of 0.2 mM. ^aFixed value; ^bValue fixed identical to the first subshell; ${}^{c}S_{0}{}^{2}$ is the amplitude reduction factor, estimated by the fitting for Zn(NO₃)_{2 aq}, and set to 0.97 for all samples; CN is the coordination number; R is the interatomic distance in Å; σ^{2} in Å² is the Debye–Waller factor in Å²; ΔE_{0} is the energy shift in eV; R-factor is the absolute misfit between experimental data and theory.

Table 4. DFT-calculated bond distances and $10^{3} \ln \beta^{66} \text{Zn}^{/64} \text{Zn}$ at 300 K of geometry-optimized

model	Zn ²⁺ ·18(H ₂ O)	$Zn^{2+}.6(H_2O)$	ZnMn ₁₂	D ₂₄ ·8H ₂ O		Zn_2Mn_{12}	O ₂₄ ·8H ₂ O	
model	ZII ·18(H2O)	ZII [•] 0(H ₂ O)	model 1	model 2	moo	lel 3	moc	lel 4
position	—	—	center	corner	center	corner	corner	corner
d(Zn-O) (Å)	2.07	2.12	2.02	1.92	2.03	1.94	1.93	1.93
	2.08	2.12	2.05	1.92	2.07	2.07	1.94	1.97
	2.13	2.12	2.11	1.98	2.08	2.12	1.97	2.01
	2.13	2.12	2.11	2.04	2.15	2.19	1.99	2.03
	2.28	2.12	2.14	_	2.21	2.31	_	_
	2.31	2.12	2.14	_	2.27	2.39	_	_
	<2.17>	<2.12>	<2.09>	<1.96>	<2.14>	<2.17>	<1.96>	<1.98>
$10^{3} ln\beta$ (‰)	2.8	3.0	3.33	3.80	2.85	2.78	3.90	3.47

1115 aqueous Zn^{2+} clusters and Zn-todorokite adsorption models.

1117 Values in angle brackets are the average distances.

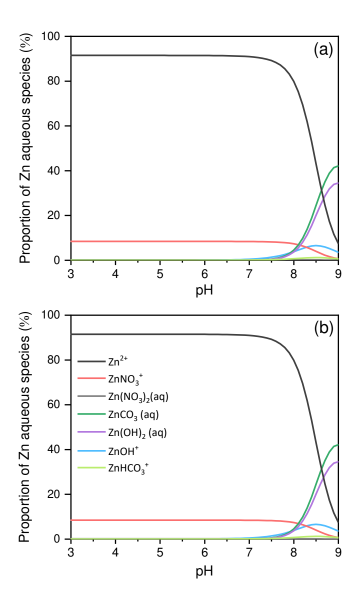


Figure S1. Zn speciation was calculated by Visual MINTEQ 3.1 as a function of pH for 0.05 mM
Zn(NO₃)₂ (a) and 0.2 mM Zn(NO₃)₂ (b). The background electrolyte of 0.1 M NaNO₃ is used.

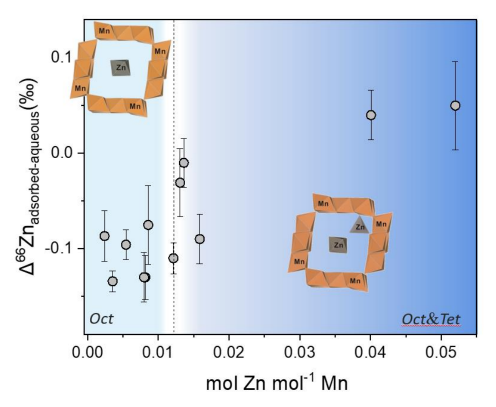


Fig. S2. Schematic diagram of the speciation of adsorbed Zn and associated isotopic fractionation on todorokite with respect to different Zn/Mn atomic ratios. The dashed line and shaded area are assigned based on approximate locations obtained from Zn isotope and EXAFS spectroscopic analyses. Oct refers to surface complexes of Zn coordinated with six oxygen atoms, and Tet refers to inner-sphere corner-sharing Zn complexes in tetrahedral coordination with four oxygen atoms.

- 1102
- 1163
- 1164

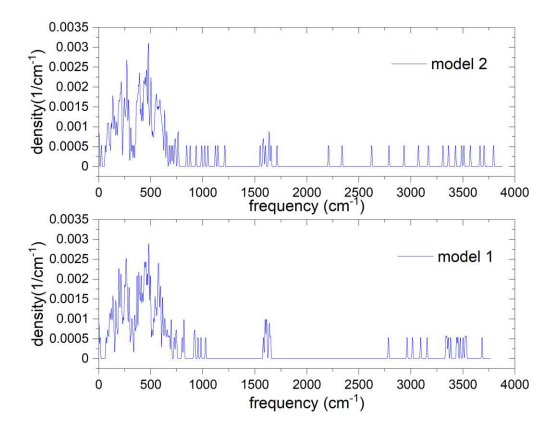


Fig. S3. Calculated vibrational frequency distribution for todorokite with octahedral Zn
complex (model 1) and tetrahedral Zn complex (model 2) in the tunnel (see Table 4; Fig. 6).

mode	frequency	mode	frequency	mode	frequency	mode	frequency	mode	frequency	mode	frequency
1	-0.063994	31	166.021714	61	389.267016	91	722.102712	121	1600.290088	151	3603.178472
2	-0.053651	32	174.522895	62	394.462786	92	734.478207	122	1604.379479	152	3730.175376
3	-0.035946	33	179.861923	63	399.932967	93	756.731442	123	1617.371621	153	3757.964777
4	54.046687	34	182.908279	64	415.769212	94	765.864994	124	1631.256126	154	3783.267821
5	69.797570	35	194.011509	65	419.661500	95	784.764501	125	1635.803142	155	3788.071655
6	78.993112	36	197.309681	66	434.158955	96	801.699106	126	1646.224439	156	3788.574558
7	86.487339	37	204.988166	67	445.213720	97	828.270354	127	1656.653680	157	3802.381993
8	97.031409	38	210.381849	68	462.738553	98	847.600924	128	1662.063232	158	3807.501854
9	97.390051	39	213.763057	69	464.180890	99	856.334263	129	1676.376898	159	3808.012953
10	102.550787	40	224.425616	70	468.179548	100	878.154549	130	2928.170043	160	3822.440595
11	102.874733	41	230.050622	71	475.404656	101	884.813513	131	3031.230875	161	3825.380287
12	109.914249	42	236.135761	72	482.847085	102	904.963291	132	3036.076586	162	3827.541174
13	111.763895	43	253.763173	73	484.457018	103	922.440093	133	3068.348404	163	3828.762801
14	114.372725	44	255.139537	74	491.859407	104	927.756921	134	3111.321918	164	3834.596816
15	119.066005	45	258.509348	75	497.134984	105	962.652455	135	3161.667959	165	3855.980483
16	120.929323	46	268.911889	76	499.677391	106	975.366068	136	3224.863060		
17	124.418906	47	275.481480	77	512.831368	107	978.034905	137	3235.519503		
18	126.292720	48	276.919269	78	521.755825	108	1022.119072	138	3261.452794		
19	130.273929	49	284.757014	79	538.742739	109	1023.475738	139	3317.950758		
20	131.946246	50	290.452096	80	547.683967	110	1053.490639	140	3327.957290		
21	137.870369	51	293.524121	81	554.792466	111	1093.525621	141	3363. 1 89707		
22	139.257778	52	312.262238	82	558.279331	112	1558.643755	142	3391.929227		
23	141.419605	53	325.094361	83	574.624535	113	1562.651430	143	3399.484043		
24	142.044549	54	331.011978	84	597.877899	114	1568.702901	144	3415.153964		
25	147.596160	55	336.721561	85	608.153959	115	1571.719201	145	3456.906280		
26	149.519126	56	350.872407	86	619.625586	116	1573.258393	146	3462.3 1 6505		
27	151.763994	57	368.459135	87	639.380829	117	1590.311060	147	3484.040549		
28	156.097570	58	370.172946	88	657.830940	118	1592.337638	148	3492.490737		
29	159.826666	59	377.709630	89	696.188202	119	1594.411632	149	3567.8 1 5999		
30	162.598666	60	380.678281	90	711.504624	120	1598.565532	150	3575.801221		

Fig. S4. Calculated vibrational frequencies of $Zn^{2+} \cdot 18(H_2O)$ in a $30 \times 30 \times 30$ Å cell.

#======================================					
# Aqua_Zn_g #	-	y Kideok Kwon			
data_VESTA	_phase_1				
_chemical_n		on	"		
_cell_length_			30.00000		
_cell_length_ cell_length	_		30.00000 30.00000		
_cell_angle_			90		
_cell_angle_			90		
_cell_angle_	gamma		90		
_space_group			'P 1'		
_space_group	p_IT_numb	er	1		
loop					
	p symop o	peration xyz			
'x, y, z'					
loop					
atom si	te label				
	te_occupan	cy			
_atom_si	te_fract_x				
	te_fract_y				
	te_fract_z				
	te_adp_typ te U iso o				
	te_type_syi				
– – H1	1.0	0.373995	0.647149	0.451344	Uiso 0.010000 H
H2	1.0	0.375172	0.594671	0.446471	Uiso 0.010000 H
H3	1.0	0.664050	0.647650	0.552901	Uiso 0.010000 H
H4 H5	1.0 1.0	0.651402 0.558167	0.690576 0.637646	$0.526740 \\ 0.445479$	Uiso 0.010000 H Uiso 0.010000 H
H6	1.0	0.527050	0.677275	0.443479	Uiso 0.010000 H
H7	1.0	0.625090	0.602347	0.447150	Uiso 0.010000 H
H8	1.0	0.618912	0.628871	0.492411	Uiso 0.010000 H
H9	1.0	0.507973	0.607608	0.391773	Uiso 0.010000 H
H10 H11	1.0 1.0	0.504458	0.576935	0.348737	Uiso 0.010000 H
H11 H12	1.0	0.354094 0.406154	0.521014 0.526773	0.412243 0.423296	Uiso 0.010000 H Uiso 0.010000 H
H12 H13	1.0	0.349501	0.575706	0.597548	Uiso 0.010000 H
H14	1.0	0.390836	0.545375	0.610490	Uiso 0.010000 H
H15	1.0	0.401863	0.468970	0.539260	Uiso 0.010000 H
H16	1.0	0.393092	0.418151	0.553423	Uiso 0.010000 H
H17 H18	1.0 1.0	0.598183 0.582986	0.442423 0.469730	0.534869 0.576217	Uiso 0.010000 H Uiso 0.010000 H
H18 H19	1.0	0.382980	0.433288	0.370217 0.452051	Uiso 0.010000 H
H2O	1.0	0.490770	0.416002	0.402561	Uiso 0.010000 H
H21	1.0	0.426994	0.614146	0.598370	Uiso 0.010000 H
H22	1.0	0.464543	0.650542	0.593967	Uiso 0.010000 H
H23 H24	1.0 1.0	0.433561 0.420664	0.491699 0.472512	0.656905 0.609211	Uiso 0.010000 H Uiso 0.010000 H
H24 H25	1.0	0.420004	0.563767	0.009211	Uiso 0.010000 H
H26	1.0	0.565631	0.516469	0.508776	Uiso 0.010000 H
H27	1.0	0.488913	0.620276	0.471052	Uiso 0.010000 H
H28	1.0	0.436863	0.611814	0.480478	Uiso 0.010000 H
H29	1.0	0.518362	0.457058	0.527602	Uiso 0.010000 H
H30 H31	1.0 1.0	0.465597 0.477170	0.448004 0.543382	0.529632 0.411290	Uiso 0.010000 H Uiso 0.010000 H
H31 H32	1.0	0.477492	0.490510	0.411290	Uiso 0.010000 H
H32	1.0	0.482223	0.576070	0.578451	Uiso 0.010000 H
H34	1.0	0.472560	0.525762	0.589638	Uiso 0.010000 H
H35	1.0	0.387268	0.525724	0.487917	Uiso 0.010000 H
H36	1.0	0.392819	0.542804	0.538905	Uiso 0.010000 H

1174 Table S1. Atomic coordinates (in CIF format) of geometry-optimized structures

1242	02	1.0	0.642630	0.659667	0.531629	Uiso 0.010000 O	
1243	O3	1.0	0.527811	0.645258	0.433840	Uiso 0.010000 O	
1244	O4	1.0	0.602671	0.610963	0.469072	Uiso 0.010000 O	
1245	05	1.0	0.491579	0.582465	0.377989	Uiso 0.010000 O	
1246	O6	1.0	0.375234	0.535274	0.432350	Uiso 0.010000 O	
1247	07	1.0	0.380237	0.568761	0.589555	Uiso 0.010000 O	
1248	08	1.0	0.413511	0.443174	0.556999	Uiso 0.010000 O	
1249	O9	1.0	0.576554	0.464658	0.544731	Uiso 0.010000 O	
1250	O10	1.0	0.499681	0.441578	0.420460	Uiso 0.010000 O	
1251	O11	1.0	0.459077	0.619708	0.602454	Uiso 0.010000 O	
1252	O12	1.0	0.428823	0.499963	0.625838	Uiso 0.010000 O	
1253	O13	1.0	0.467933	0.603201	0.490181	Uiso 0.010000 O	
1254	O14	1.0	0.549529	0.536836	0.488700	Uiso 0.010000 O	
1255	O15	1.0	0.489945	0.461273	0.511344	Uiso 0.010000 O	
1256	O16	1.0	0.463951	0.520599	0.432700	Uiso 0.010000 O	
1257	O17	1.0	0.489360	0.544616	0.568381	Uiso 0.010000 O	
1258	O18	1.0	0.405199	0.522080	0.515766	Uiso 0.010000 O	
1259	Zn1	1.0	0.479362	0.535865	0.500399	Uiso 0.010000 Zn	
1260							

1261	#					
1262	# todorokite	with center	Zn + corner Zn	generated by Kide	eok Kwon	
1263	#					
1264						
1265	data_VESTA	_phase_1				
1266 1267	ah ami a al m		~ **	"		
1267	chemical_n cell length	_	on	10.09752		
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1272	_cell_angle_			95.56499		
1273	_cell_angle_	-		90		
1274	_space_grou		_	'P 1'		
1275 1276	_space_grou	p_11_numb	er	1		
1277	loop					
1278		p symop of	peration xyz			
1279	'x, y, z'	,,				
1280						
1281	loop_					
1282	_atom_si	_				
1283 1284		te_occupano te fract x	cy			
1285		te fract y				
1286		te fract z				
1287		te_adp_type	e			
1288		te_U_iso_o				
1289		te_type_syn				
1290 1291	H1	1.0	0.329082	0.335333	0.746587	Uiso 0.010000 H
1291	H2 H3	1.0 1.0	0.329039 0.692644	0.584329 0.400854	0.669824 0.345317	Uiso 0.010000 H Uiso 0.010000 H
1293	H4	1.0	0.543313	0.468731	0.258119	Uiso 0.010000 H
1294	H5	1.0	0.261840	0.570551	0.388263	Uiso 0.010000 H
1295	H6	1.0	0.369294	0.776457	0.398045	Uiso 0.010000 H
1296	H7	1.0	0.709554	0.105743	0.640013	Uiso 0.010000 H
1297	H8	1.0	0.581035	0.056236	0.717826	Uiso 0.010000 H
1298	H9	1.0	0.275043	0.103937	0.378869	Uiso 0.010000 H
1299 1300	H10 H11	1.0 1.0	0.407806 0.702185	0.078529 0.565439	0.300580 0.646803	Uiso 0.010000 H Uiso 0.010000 H
1301	H12	1.0	0.583073	0.570155	0.744982	Uiso 0.010000 H
1302	H13	1.0	0.373726	0.940398	0.589971	Uiso 0.010000 H
1303	H14	1.0	0.400259	0.918803	0.752904	Uiso 0.010000 H
1304	H15	1.0	0.617226	0.723049	0.385099	Uiso 0.010000 H
1305	H16	1.0	0.618008	0.952325	0.485416	Uiso 0.010000 H
1306 1307	01	1.0	0.448240	0.014121	0.127484	Uiso 0.010000 O Uiso 0.010000 O
1307	O2 O3	1.0 1.0	0.917209 0.873600	0.011561 0.016294	0.157400 0.651025	Uiso 0.010000 O
1309	04	1.0	0.543243	0.006960	0.891854	Uiso 0.010000 O
1310	05	1.0	0.058278	0.013046	0.848859	Uiso 0.010000 O
1311	O6	1.0	0.103412	0.016556	0.353576	Uiso 0.010000 O
1312	07	1.0	0.145371	0.258689	0.113393	Uiso 0.010000 O
1313	08	1.0	0.685263	0.245323	0.092979	Uiso 0.010000 O
1314 1315	09	1.0	0.892520	0.257577	0.390115	Uiso 0.010000 O
1316	O10 O11	1.0 1.0	0.833373 0.312773	0.266525 0.264223	0.888525 0.930396	Uiso 0.010000 O Uiso 0.010000 O
1317	012	1.0	0.069519	0.264906	0.601388	Uiso 0.010000 O
1318	013	1.0	0.459285	0.504169	0.131681	Uiso 0.010000 O
1319	O14	1.0	0.925808	0.512994	0.150302	Uiso 0.010000 O
1320	015	1.0	0.872516	0.513463	0.649873	Uiso 0.010000 O
1321	016	1.0	0.556568	0.512140	0.903845	Uiso 0.010000 O
1322	017	1.0	0.059255	0.513687	0.847900	Uiso 0.010000 O
1323 1324	O18 O19	1.0 1.0	0.100569 0.145738	0.511510 0.763389	0.354063 0.113709	Uiso 0.010000 O Uiso 0.010000 O
1324	O19 O20	1.0	0.145/38 0.686095	0.769120	0.091608	Uiso 0.010000 O
1326	O20 O21	1.0	0.898750	0.767196	0.390571	Uiso 0.010000 O
1327	022	1.0	0.834210	0.756755	0.888264	Uiso 0.010000 O
1328	O23	1.0	0.314019	0.755205	0.937976	Uiso 0.010000 O
1329	O24	1.0	0.069562	0.760764	0.601342	Uiso 0.010000 O

1330 1331 1332 1333 1334 1335 1336 1337 1338 1339 1340 1341	O25 O26 O27 O28 O29 O30 O31 O32 Mn1 Mn2 Mn3 Mn4	$\begin{array}{c} 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\$	0.338194 0.601737 0.359014 0.609656 0.374168 0.605579 0.339709 0.622136 0.496473 -0.012346 0.496738 -0.012667	0.411334 0.452289 0.606472 0.109317 0.096745 0.600938 0.871634 0.898502 0.257776 0.013582 0.758511 0.513531	0.656275 0.356979 0.399259 0.626360 0.394720 0.646554 0.671660 0.387306 0.011920 0.501206 0.016606 0.500538	Uiso 0.010000 O Uiso 0.010000 Mn Uiso 0.010000 Mn Uiso 0.010000 Mn
1342 1343 1344 1345 1346 1347 1348 1349 1350 1351 1352	Mn5 Mn6 Mn7 Mn8 Mn9 Mn10 Mn11 Mn12 Zn1 Zn1 Zn2	$ \begin{array}{c} 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\ 1.0\\$	0.215415 0.019128 0.769427 0.965490 0.219340 0.017318 0.759733 0.966294 0.746451 0.480904	0.510814 0.760034 0.510264 0.762936 0.010009 0.264693 0.010306 0.263709 0.996133 0.382990	1.011237 0.238736 -0.000620 0.763229 1.011192 0.238158 -0.024409 0.763158 0.258490 0.523201	Uiso 0.010000 Mn Uiso 0.010000 Zn Uiso 0.010000 Zn

1354	#					
1355 1356	# todorokite #	with center	Zn generated b	y Kideok Kwon		
1357	π					
1358 1359	data_VESTA	_phase_1				
1360	chemical n	ame comm	on	"		
1361	cell_length_	_a		10.00397		
1362	_cell_length			5.76920		
1363 1364	_cell_length			9.67044 90		
1365	_cell_angle_ _cell_angle_			90 94.26335		
1366	cell angle			90		
1367	space_grou		M_alt	'P 1'		
1368	_space_grou	p_IT_numb	er	1		
1369 1370	loon					
1370	loopspace_group	n symon o	peration xyz			
1372	yrou 'x, y, z'	p_ojmop_o	peration_ity2			
1373						
1374	loop_					
1375 1376	_atom_si	te_label te_occupan	A			
1377		te_fract x	cy			
1378		te_fract_y				
1379	_atom_si	te_fract_z				
1380		te_adp_type				
1381 1382		te_U_iso_o te_type_syr				
1383	_atom_si H1	1.0	0.341869	0.262644	0.752013	Uiso 0.010000 H
1384	H2	1.0	0.328182	0.482901	0.658062	Uiso 0.010000 H
1385	H3	1.0	0.636490	0.501912	0.382005	Uiso 0.010000 H
1386	H4	1.0	0.643517	0.261186	0.299327	Uiso 0.010000 H
1387 1388	H5	1.0	0.272456	0.543852	0.388503	Uiso 0.010000 H
1389	Н6 Н7	1.0 1.0	0.402208 0.596428	0.698244 0.019193	0.392557 0.717616	Uiso 0.010000 H Uiso 0.010000 H
1390	H8	1.0	0.472684	-0.115748	0.641457	Uiso 0.010000 H
1391	Н9	1.0	0.277399	0.057560	0.369550	Uiso 0.010000 H
1392	H10	1.0	0.410352	0.049998	0.287362	Uiso 0.010000 H
1393 1394	H11	1.0	0.705206	0.505730	0.640341	Uiso 0.010000 H
1394	H12 H13	1.0 1.0	0.593612 0.244164	0.512594 0.814774	0.748463 0.634418	Uiso 0.010000 H Uiso 0.010000 H
1396	H14	1.0	0.329095	0.791888	0.778320	Uiso 0.010000 H
1397	H15	1.0	0.624653	0.834625	0.487160	Uiso 0.010000 H
1398	H16	1.0	0.727492	0.818948	0.372211	Uiso 0.010000 H
1399 1400	O1 O2	1.0 1.0	0.458837	0.073289	0.131959 0.144040	Uiso 0.010000 O Uiso 0.010000 O
1400	02 03	1.0	$0.924666 \\ 0.878288$	0.073811 0.074464	0.144040	Uiso 0.010000 O
1402	04	1.0	0.589726	0.071542	0.916688	Uiso 0.010000 O
1403	05	1.0	0.064936	0.073457	0.850459	Uiso 0.010000 O
1404	06	1.0	0.100884	0.074849	0.345843	Uiso 0.010000 O
1405 1406	O7 O8	1.0 1.0	$0.151020 \\ 0.694840$	0.322984 0.321775	0.111674 0.109038	Uiso 0.010000 O Uiso 0.010000 O
1407	08	1.0	0.903566	0.325263	0.395243	Uiso 0.010000 O
1408	O10	1.0	0.838023	0.323876	0.905146	Uiso 0.010000 O
1409	O11	1.0	0.328633	0.319560	0.935086	Uiso 0.010000 O
1410	012	1.0	0.071244	0.326480	0.602968	Uiso 0.010000 O
1411 1412	O13 O14	1.0 1.0	0.455824 0.924251	0.571511 0.574208	0.127981 0.144290	Uiso 0.010000 O Uiso 0.010000 O
1413	014	1.0	0.924251	0.574746	0.658008	Uiso 0.010000 O
1414	016	1.0	0.588620	0.572929	0.918092	Uiso 0.010000 O
1415	017	1.0	0.064715	0.574973	0.850285	Uiso 0.010000 O
1416	018	1.0	0.099898	0.574428	0.346235	Uiso 0.010000 O
1417 1418	O19 O20	1.0 1.0	0.151339 0.696500	0.822945 0.822864	0.112247 0.108673	Uiso 0.010000 O Uiso 0.010000 O
1410	O20 O21	1.0	0.896500	0.822864	0.108673 0.394743	Uiso 0.010000 O
1420	021	1.0	0.837438	0.822673	0.904065	Uiso 0.010000 O
1421	O23	1.0	0.329180	0.823735	0.938196	Uiso 0.010000 O
1422	O24	1.0	0.073996	0.825933	0.602535	Uiso 0.010000 O

1423	O25	1.0	0.338389	0.309438	0.653753	Uiso 0.010000 O	
1424	O26	1.0	0.632438	0.325916	0.391194	Uiso 0.010000 O	
1425	O27	1.0	0.370678	0.539251	0.408232	Uiso 0.010000 O	
1426	O28	1.0	0.556589	-0.025041	0.625059	Uiso 0.010000 O	
1427	O29	1.0	0.375376	0.027752	0.381467	Uiso 0.010000 O	
1428	O30	1.0	0.609388	0.474459	0.651008	Uiso 0.010000 O	
1429	O31	1.0	0.333419	0.764574	0.675731	Uiso 0.010000 O	
1430	O32	1.0	0.635632	0.775128	0.392308	Uiso 0.010000 O	
1431	Mn1	1.0	0.510591	0.323515	0.025841	Uiso 0.010000 Mn	
1432	Mn2	1.0	-0.015184	0.077556	0.503059	Uiso 0.010000 Mn	
1433	Mn3	1.0	0.513070	0.821507	0.028718	Uiso 0.010000 Mn	
1434	Mn4	1.0	-0.015169	0.573386	0.501514	Uiso 0.010000 Mn	
1435	Mn5	1.0	0.227966	0.571696	1.014711	Uiso 0.010000 Mn	
1436	Mn6	1.0	0.016443	0.825107	0.236694	Uiso 0.010000 Mn	
1437	Mn7	1.0	0.770980	0.573025	0.011136	Uiso 0.010000 Mn	
1438	Mn8	1.0	0.968235	0.824988	0.768707	Uiso 0.010000 Mn	
1439	Mn9	1.0	0.224646	0.073347	1.011379	Uiso 0.010000 Mn	
1440	Mn10	1.0	0.016311	0.323868	0.237225	Uiso 0.010000 Mn	
1441	Mn11	1.0	0.769238	0.072862	0.009023	Uiso 0.010000 Mn	
1442	Mn12	1.0	0.968815	0.324214	0.768275	Uiso 0.010000 Mn	
1443	Zn1	1.0	0.482823	0.270441	0.515699	Uiso 0.010000 Zn	
1444							

-5 -6	#========= # todorokite	with corner	Zn + corner Zn	generated by Kid	eok Kwon		
.7 .8	#						
.0 .9	data VESTA	phase 1					
0	-						
1 2	_chemical_n	_	on	"			
3	_cell_length _cell_length			10.09522 5.76410			
4	_cell_length			9.66638			
5	_cell_angle_	alpha		90			
ò,	_cell_angle_			95.78459			
	_cell_angle_ _space_grou	0	M alt	90 'P 1'			
	_space_grou			1			
	_1 _0 ,						
	loop_						
	_space_grou	p_symop_o	peration_xyz				
	'x, y, z'						
	loop_						
	_atom_si	_					
		te_occupan	cy				
		te_fract_x te_fract_y					
		te_fract_y					
		te_adp_type	2				
	_atom_si	te_U_iso_o	r_equiv				
		te_type_syr		0.540(42	0.542544	LI: 0.010000 LI	
	H1 H2	1.0 1.0	0.472122 0.392869	0.549643 0.585332	0.743744 0.583717	Uiso 0.010000 H Uiso 0.010000 H	
	H3	1.0	0.643034	0.436447	0.383717	Uiso 0.010000 H	
	H4	1.0	0.525494	0.562986	0.382897	Uiso 0.010000 H	
	H5	1.0	0.373619	0.789745	0.399354	Uiso 0.010000 H	
	H6	1.0	0.288104	0.555216	0.385852	Uiso 0.010000 H	
	H7	1.0	0.654600	0.143853	0.618245	Uiso 0.010000 H	
	H8 H9	1.0 1.0	0.742558 0.282547	$0.369640 \\ 0.078643$	0.658576 0.356923	Uiso 0.010000 H Uiso 0.010000 H	
	H10	1.0	0.419732	0.071392	0.282933	Uiso 0.010000 H	
	H11	1.0	0.402547	1.063528	0.543140	Uiso 0.010000 H	
	H12	1.0	0.465257	0.975639	0.693390	Uiso 0.010000 H	
	H13	1.0	0.634421	0.759849	0.557645	Uiso 0.010000 H	
	H14 H15	1.0 1.0	0.698237 0.770179	0.805664 2.164562	0.712153 0.386634	Uiso 0.010000 H Uiso 0.010000 H	
	H16	1.0	0.670950	2.164832	0.252933	Uiso 0.010000 H	
	O1	1.0	0.451788	0.023936	0.126654	Uiso 0.010000 O	
	02	1.0	0.936898	0.015081	0.166213	Uiso 0.010000 O	
	O3	1.0	0.902779	0.017592	0.665981	Uiso 0.010000 O	
	O4 O5	1.0 1.0	0.560735 0.078192	0.014624 0.016516	0.905174 0.868787	Uiso 0.010000 O Uiso 0.010000 O	
	05 06	1.0	0.119993	0.017780	0.364526	Uiso 0.010000 O	
	07	1.0	0.162585	0.268296	0.126127	Uiso 0.010000 O	
	08	1.0	0.692316	0.267142	0.093387	Uiso 0.010000 O	
	09	1.0	0.923070	0.265755	0.414999	Uiso 0.010000 O	
	O10 O11	1.0 1.0	0.850827 0.319691	0.267032 0.265470	0.901157 0.929000	Uiso 0.010000 O Uiso 0.010000 O	
	011	1.0	0.319691 0.104190	0.265470	0.929000	Uiso 0.010000 O	
	012	1.0	0.447307	0.514552	0.122253	Uiso 0.010000 O	
	O14	1.0	0.934840	0.519170	0.168788	Uiso 0.010000 O	
	015	1.0	0.899778	0.515127	0.666665	Uiso 0.010000 O	
	O16 O17	1.0 1.0	0.561985	0.522528	0.903524	Uiso 0.010000 O Uiso 0.010000 O	
	017	1.0 1.0	0.079132 0.120432	0.515796 0.514383	0.869181 0.365345	Uiso 0.010000 O Uiso 0.010000 O	
	018	1.0	0.120432	0.767213	0.130606	Uiso 0.010000 O	
	O20	1.0	0.693604	0.764295	0.100527	Uiso 0.010000 O	
	O21	1.0	0.919087	0.764468	0.411156	Uiso 0.010000 O	
	O22 O23	1.0 1.0	0.846305	0.767426 0.769292	0.898249 0.938174	Uiso 0.010000 O Uiso 0.010000 O	
	023 024	1.0 1.0	0.318708 0.098022	0.769292 0.764104	0.938174 0.621677	Uiso 0.010000 O Uiso 0.010000 O	
	024	1.0	0.090022	0./04104	0.0210//	0180 0.010000 0	

O25	1.0	0.384730	0.552922	0.685044	Uiso 0.010000 O
O26	1.0	0.625339	0.563953	0.408267	Uiso 0.010000 O
O27	1.0	0.377504	0.618066	0.419154	Uiso 0.010000 O
O28	1.0	0.654695	0.317934	0.615727	Uiso 0.010000 O
O29	1.0	0.381223	0.069576	0.375576	Uiso 0.010000 O
O30	1.0	0.382300	1.031107	0.640687	Uiso 0.010000 O
O31	1.0	0.631043	0.861531	0.639677	Uiso 0.010000 O
O32	1.0	0.687208	2.084726	0.345051	Uiso 0.010000 O
Mn1	1.0	0.508154	0.268628	0.012029	Uiso 0.010000 Mn
Mn2	1.0	0.013182	0.014300	0.519079	Uiso 0.010000 Mn
Mn3	1.0	0.502436	0.767779	0.015466	Uiso 0.010000 Mn
Mn4	1.0	0.011436	0.516878	0.517795	Uiso 0.010000 Mn
Mn5	1.0	0.238055	0.520119	1.029275	Uiso 0.010000 Mn
Mn6	1.0	0.035161	0.766765	0.250284	Uiso 0.010000 Mn
Mn7	1.0	0.778579	0.515823	0.001329	Uiso 0.010000 Mn
Mn8	1.0	0.987846	0.766110	0.779482	Uiso 0.010000 Mn
Mn9	1.0	0.233464	0.015257	1.025761	Uiso 0.010000 Mn
Mn10	1.0	0.030444	0.266515	0.251281	Uiso 0.010000 Mn
Mn11	1.0	0.777335	0.018124	0.000517	Uiso 0.010000 Mn
Mn12	1.0	0.984529	0.265758	0.783488	Uiso 0.010000 Mn
Zn1	1.0	0.735840	0.756184	0.303774	Uiso 0.010000 Zn
Zn2	1.0	0.286532	0.277322	0.725163	Uiso 0.010000 Zn
	O26 O27 O28 O29 O30 O31 O32 Mn1 Mn2 Mn3 Mn4 Mn5 Mn6 Mn7 Mn8 Mn9 Mn10 Mn11 Mn12 Zn1	$\begin{array}{cccc} 026 & 1.0 \\ 027 & 1.0 \\ 028 & 1.0 \\ 029 & 1.0 \\ 030 & 1.0 \\ 031 & 1.0 \\ 032 & 1.0 \\ 031 & 1.0 \\ 032 & 1.0 \\ Mn1 & 1.0 \\ Mn2 & 1.0 \\ Mn3 & 1.0 \\ Mn3 & 1.0 \\ Mn4 & 1.0 \\ Mn5 & 1.0 \\ Mn6 & 1.0 \\ Mn6 & 1.0 \\ Mn7 & 1.0 \\ Mn7 & 1.0 \\ Mn8 & 1.0 \\ Mn9 & 1.0 \\ Mn9 & 1.0 \\ Mn9 & 1.0 \\ Mn10 & 1.0 \\ Mn11 & 1.0 \\ Mn12 & 1.0 \\ Zn1 & 1.0 \\ Zn1 & 1.0 \\ \end{array}$	O26 1.0 0.625339 O27 1.0 0.377504 O28 1.0 0.654695 O29 1.0 0.381223 O30 1.0 0.382300 O31 1.0 0.631043 O32 1.0 0.687208 Mn1 1.0 0.508154 Mn2 1.0 0.013182 Mn3 1.0 0.502436 Mn4 1.0 0.011436 Mn5 1.0 0.238055 Mn6 1.0 0.035161 Mn7 1.0 0.778579 Mn8 1.0 0.987846 Mn9 1.0 0.233464 Mn10 1.0 0.777335 Mn12 1.0 0.735840	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	O26 1.0 0.625339 0.563953 0.408267 O27 1.0 0.377504 0.618066 0.419154 O28 1.0 0.654695 0.317934 0.615727 O29 1.0 0.381223 0.069576 0.375576 O30 1.0 0.382300 1.031107 0.640687 O31 1.0 0.631043 0.861531 0.639677 O32 1.0 0.687208 2.084726 0.345051 Mn1 1.0 0.508154 0.268628 0.012029 Mn2 1.0 0.013182 0.014300 0.519079 Mn3 1.0 0.502436 0.767779 0.015466 Mn4 1.0 0.011436 0.516878 0.517795 Mn5 1.0 0.238055 0.520119 1.029275 Mn6 1.0 0.035161 0.766765 0.250284 Mn7 1.0 0.778579 0.515823 0.001329 Mn8 1.0 0.987846 0.766110<

1538	#						
1539	# todorokite with corner Zn generated by Kideok Kwon #						
1540 1541	#				-		
1542	data_VEST	A_phase_1					
1543							
1544 1545		name_commo	on	" 9.94486			
1546		_cell_length_a _cell_length_b					
1547		_cell_length_c					
1548		_cell_angle_alpha					
1549 1550	_cell_angle			93.43745 90			
1550	•	_cell_angle_gamma space group name H-M alt					
1552		up_IT_numb		'P 1' 1			
1553							
1554 1555	loop_						
1555	_space_gro 'x, y, z'	up_symop_op	peration_xyz				
1557	<i>A</i> , <i>J</i> , <i>L</i>						
1558	loop_						
1559		site_label					
1560 1561		site_occupand site fract x	cy				
1562		site_fract_x					
1563		site_fract_z					
1564		site_adp_type					
1565 1566		site_B_iso_oi site type syn					
1567	H	1.0	0.304897	0.487388	0.721618	Biso 1.000000 H	
1568	Н	1.0	0.352542	0.511284	0.566401	Biso 1.000000 H	
1569	Н	1.0	0.622892	0.439881	0.513773	Biso 1.000000 H	
1570 1571	H H	1.0 1.0	0.518967	0.525480	0.386922	Biso 1.000000 H Biso 1.000000 H	
1572	Н	1.0	0.287279 0.372125	0.576156 0.801219	0.364558 0.413990	Biso 1.000000 H	
1573	Н	1.0	0.646127	-0.249696	0.674843	Biso 1.000000 H	
1574	Н	1.0	0.504945	-0.101978	0.649735	Biso 1.000000 H	
1575	Н	1.0	0.273188	0.091565	0.371738	Biso 1.000000 H	
1576 1577	H H	1.0 1.0	0.406961 0.641848	0.074003 0.196092	0.293199 0.667202	Biso 1.000000 H Biso 1.000000 H	
1578	Н	1.0	0.524343	0.381423	0.675452	Biso 1.000000 H	
1579	Н	1.0	0.345430	1.118394	0.658197	Biso 1.000000 H	
1580	H	1.0	0.338452	0.904198	0.761934	Biso 1.000000 H	
1581 1582	H H	1.0 1.0	$0.628642 \\ 0.680459$	0.925717 0.809857	0.486242 0.347989	Biso 1.000000 H Biso 1.000000 H	
1583	0	1.0	0.450590	0.045569	0.126161	Biso 1.000000 O	
1584	О	1.0	0.922135	0.040819	0.147429	Biso 1.000000 O	
1585	0	1.0	0.884965	0.040261	0.660698	Biso 1.000000 O	
1586 1587	0 0	1.0 1.0	0.592619 0.071025	0.042563 0.041175	0.919584 0.848940	Biso 1.000000 O Biso 1.000000 O	
1588	0	1.0	0.100825	0.042338	0.342549	Biso 1.000000 O	
1589	0	1.0	0.151022	0.291394	0.108329	Biso 1.000000 O	
1590	Ο	1.0	0.693652	0.289998	0.113106	Biso 1.000000 O	
1591 1592	0	1.0	0.899591	0.288321 0.291311	0.396040	Biso 1.000000 O	
1592	0 0	1.0 1.0	0.843246 0.329970	0.291311	0.907938 0.930919	Biso 1.000000 O Biso 1.000000 O	
1594	Õ	1.0	0.078120	0.289599	0.599697	Biso 1.000000 O	
1595	0	1.0	0.450507	0.541467	0.125820	Biso 1.000000 O	
1596	0	1.0	0.923265	0.540752	0.146318	Biso 1.000000 O	
1597 1598	0 0	1.0 1.0	0.885353 0.592461	0.543023 0.544428	0.660409 0.920322	Biso 1.000000 O Biso 1.000000 O	
1599	0	1.0	0.072959	0.540905	0.920322	Biso 1.000000 O	
1600	0	1.0	0.102034	0.538106	0.341360	Biso 1.000000 O	
1601	0	1.0	0.150108	0.792022	0.107534	Biso 1.000000 O	
1602 1603	0	1.0	0.695218	0.791346	0.109611	Biso 1.000000 O	
1603	0 0	1.0 1.0	0.905829 0.841690	0.788243 0.790292	0.398841 0.905542	Biso 1.000000 O Biso 1.000000 O	
1605	0 0	1.0	0.331273	0.792464	0.930849	Biso 1.000000 O	
1606	О	1.0	0.080403	0.789067	0.600381	Biso 1.000000 O	

1607	0	1.0	0.360898	0.416543	0.654317	Biso 1.000000 O
1608	0	1.0	0.618098	0.491260	0.408708	Biso 1.000000 O
1609	0	1.0	0.370323	0.624121	0.417069	Biso 1.000000 O
1610	0	1.0	0.606294	-0.111290	0.630658	Biso 1.000000 O
1611	0	1.0	0.371753	0.074773	0.386872	Biso 1.000000 O
1612	0	$1.0 \\ 1.0$	0.621552	0.363409	0.654752	Biso 1.000000 O
1613	0		0.358678	0.947372	0.667112	Biso 1.000000 O
1614	0	1.0	0.643118	0.955201	0.381436	Biso 1.000000 O
1615	Mn	1.0	0.507150	0.293803	0.021721	Biso 1.000000 Mn
1616	Mn	1.0	-0.009262	0.037876	0.503795	Biso 1.000000 Mn
1617	Mn	1.0	0.511816	0.791464	0.025165	Biso 1.000000 Mn
1618	Mn	1.0	-0.009735	0.542192	0.503355	Biso 1.000000 Mn
1619	Mn	1.0	0.229429	0.542451	1.009925	Biso 1.000000 Mn
1620	Mn		0.017287	0.790201	0.236416	Biso 1.000000 Mn
1621	Mn	1.0 1.0	0.771657	0.543481	0.009258	Biso 1.000000 Mn
1622	Mn	1.0	0.976228	0.790621	0.767348	Biso 1.000000 Mn
1623	Mn	1.0	0.227847	0.041979	1.007894	Biso 1.000000 Mn
1624	Mn	1.0	0.018961	0.290760	$0.233930 \\ 0.007441$	Biso 1.000000 Mn
1625	Mn	1.0	0.770840	0.039067		Biso 1.000000 Mn
1626	Mn	1.0	0.974805	0.290775	0.767263	Biso 1.000000 Mn
1627 1628	Zn	1.0	0.706911	0.246373	0.315942	Biso 1.000000 Zn