1	Original Paper
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4	Copper isotope fractionation between aqueous compounds
5	relevant to low temperature geochemistry and biology
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25 Abstract:

Isotope fractionation between the common Cu species present in solution (Cu^{+}, Cu^{2+}) 26 hydroxide, chloride, sulfide, carbonate, oxalate, and ascorbate) has been investigated 27 28 using both *ab initio* methods and experimental solvent extraction techniques. In order to 29 establish unambiguously the existence of equilibrium isotope fractionation (as opposed 30 to kinetic isotope fractionation), we first performed laboratory-scale liquid-liquid 31 distribution experiments. Upon exchange between HCl medium and a macrocyclic complex, the 65 Cu/ 63 Cu ratio fractionated by -1.06 to -0.39‰. The acidity dependence 32 33 of the fractionation was appropriately explained by ligand exchange reactions between 34 hydrated H₂O and Cl⁻ via intramolecular vibrations. The magnitude of the Cu isotope 35 fractionation among important Cu ligands was also estimated by *ab initio* methods. The 36 magnitude of the nuclear field shift effect to the Cu isotope fractionation represents only 37 \sim 3% of the mass-dependent fractionation. The theoretical estimation was expanded to 38 chlorides, hydroxides, sulfides, sulfates, and carbonates under different conditions of 39 pH. Copper isotope fractionation of up to 2‰ is expected for different forms of Cu 40 present in seawater and for different sediments (carbonates, hydroxides, and sulfides). 41 We found that Cu in dissolved carbonates and sulfates is isotopically much heavier 42 (+0.6‰) than free Cu. Isotope fractionation of Cu in hydroxide is minimal. The 43 relevance of these new results to the understanding of metabolic processes was also 44 discussed. Copper is an essential element used by a large number of proteins for electron transfer. Further theoretical estimates of δ^{65} Cu in hydrated Cu(I) and Cu(II) 45 ions, Cu(II) ascorbates, and Cu(II) oxalate predict Cu isotope fractionation during the 46 47 breakdown of ascorbate into oxalate and account for the isotopically heavy Cu found in 48 animal kidneys.

50 Keywords: Copper, Cupric, isotope fractionation, ligand, quantum chemical calculation

1. INTRODUCTION

Copper has two stable isotopes, ⁶³Cu and ⁶⁵Cu, with respective average 53 54 abundances of 69.174% and 30.826% in the reference metal SRM-NIST 976 (Shields et 55 al., 1964). Variations of Cu isotope abundances in natural samples were identified by 56 Walker et al. (1958) using thermal ionization mass spectrometry, but the variability of 57 the mass bias at that time was too large with respect to the natural isotopic availability 58 to qualify Cu isotopes as a potential geochemical or biochemical tracer. The advent of 59 inductively-coupled plasma mass spectrometry (ICP-MS) instruments equipped with a 60 magnetic sector and multiple collection made precise isotopic analysis of Cu possible 61 (Albarède, 2004). Maréchal et al. (1999) published the first measurements of Cu isotope 62 compositions in a variety of minerals and biological materials. The broad range of 63 isotopic variations in Cu ores observed previously (Walker et al. 1958; Shields et al., 1965) was confirmed by subsequent work (Maréchal et al., 1999; Zhu et al., 2000; 64 65 Albarède, 2004; Klein et al., 2009). Hereafter, the δ notation with

$$\delta^{65} Cu = \left(\begin{array}{c} ([^{65} Cu]/[^{63} Cu])_{sample} \\ ([^{65} Cu]/[^{63} Cu])_{reference} \end{array} - 1 \right) \times 1000$$
(1)

66

67 is used throughout.

The narrow range of Cu isotope variation in basalts (Ben Othman et al., 2006; Archer and Vance, 2004; Herzog et al., 2009; Li et al., 2009; Moynier et al., 2010; Bishop et al., 2012) and granites (Li et al., 2009) suggests that high-temperature magmatic processes do not induce large Cu isotope fractionation. As a consequence, it makes Cu isotopes a tracer of choice for the study of sediments and more generally for

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low-temperature processes such as those involved in soil formation (Bigalke et al., 2010, 2011) and in biology (Balter and Zarro, 2011; Weinstein et al., 2011).

Substantial isotope fractionation during chemical exchange reactions occurs at room temperature and reflects isotopic differences between the equilibrium constants of Cu isotopologues. Enrichment in 65 Cu of up to 0.8‰ was observed during the chromatographic elution of compounds (Matin et al., 1998; Maréchal et al., 1999, 2002; Zhu et al., 2002). The same technique demonstrates isotope fractionation of ~0.4‰ upon chromatographic separation of Cu(I) from Cu(II) (Matin et al., 1998; Zhu et al., 2002).

At low temperature, Cu isotopic abundances are also affected by inorganic surface processes (physical adsorption/desorption). Isotope separation of up to a few permil upon adsorption/desorption of Cu onto/from inorganic and organic substrates was observed by several groups (Mathur et al., 2005; Balistrieri et al., 2008; Pokrovsky et al., 2008; Navarette et al., 2011).

87 Copper plays very important roles in biology (Lippard and Berg, 1994). The 88 metal centers of the 'blue' copper proteins are divided according to the coordination of 89 the Cu ion into three types that contain one or more copper ions as prosthetic groups 90 (Cowan, 1997). In plastocyanine, cysteine, histidine, and methionine 'residues' bind to 91 Cu and this plays a major role for electron transfer (redox reactions) (Freeman and Guss, 92 2001). In humans, ceruloplasmin is a ferroxidase enzyme involved in the iron cycle, 93 cytochrome c oxidase is involved in electron transfer across the membrane of 94 mitochondrion, and superoxide dismutase protects cells against superoxide. Blue (Cu) 95 proteins are ubiquitous and characterized by their multiple ways of bonding with amino 96 acids. (Cowan, 1997). Coupling of Fe and Cu isotope fractionation has been identified

97 in human blood components (Albarède et al., 2011a) and in mammals (Balter and Zazzo, 98 2011; Albarede et al., 2011b). Depletion of δ^{65} Cu at the permil level was found in plants 99 with respect to soils or nutrient solutions (Weinstein et al., 2011; Jouvin et al., 2012). 100 δ^{65} Cu in enzymes is a few permil negative compared with the Cu isotopic composition 101 of the host soil (Zhu et al., 2002). Positive δ^{65} Cu values up to 1.5‰ also were found in 102 the kidneys of sheep (Balter and Zazzo, 2011) and mice (Albarede et al., 2011b).

103 Contrary to Fe (Hill et al., 2010; Rustad et al., 2010, and references therein), Ni 104 (Fujii et al., 2011a), and Zn (Fujii et al, 2010,2011b; Black et al., 2011; Pons et al., 105 2011; Fujii and Albarède, 2012), the theoretical prediction of Cu isotope fractionation 106 among ligands (simple inorganic ligands like halide, sulfate, and phosphate ions, and 107 more complex organic ligands as humic acids and phytoplankton exometabolites) in 108 natural waters and biological fluids has only received minimal attention, which is a 109 major hindrance for an informed interpretation of observations. This is the main 110 motivation for the present work.

111 We first demonstrate that equilibrium isotope fractionation of Cu(II) does take 112 place in laboratory-scale experiments. In parallel, we calculated the molecular orbitals 113 of the corresponding Cu species to obtain the reduced partition function ratio (RPFR) of 114 isotopologues. The *ab initio* calculations were then extended to hydrated Cu(II) 115 complexes, hydroxide, sulfate, carbonate, ascorbate, and oxalate. The choice of 116 ascorbate and oxalate was motivated by evidence of isotopically heavy Cu in the 117 kidneys of sheep (Balter and Zazzo, 2011) and mice (Albarède et al., 2011b). Ascorbate 118 is allegedly catabolized into oxalic acid (Chai et al., 2004; Massey et al., 2005; González et al., 2005), which makes some patients develop a condition known as 119 120 hyperoxaluria and develop kidney stones, a calcium oxalate. The oxalate anions are

121 known to be strong complexing ligands of Cu(II) (Hayakawa et al., 1973), while 122 ascorbate oxidase, which oxidizes ascorbate into dehydroascorbate (Solomon et al., 123 1996) is a bis(histidine)copper(II) compound. Applications of our calculations to the 124 understanding of Cu isotope variability in marine sediments and to biological samples 125 are briefly outlined.

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127

2. METHODS

128 **2.1. Extraction experiments and isotopic analyses**

Dicyclohexano-18-crown-6 (DC18C6) (>97% purity) is a product of Fluka Chemie GmbH and Cu dichloride (hydrated, 99.9% purity) is a product of Wako Pure Chemical Industries, Ltd. Hydrochloric acid, in which the Cu impurity was certified to be less than 1.9 ppt, was purchased from Kanto Chemical, Co., Inc. Other chemicals were reagent grade.

134 $CuCl_2 \cdot 2H_2O$ was dissolved in HCl to prepare solutions of 0.1 mol dm⁻³ (M) 135 Cu(II) in 1 to 6 M HCl. DC18C6 was dissolved in 1,2-dichloroethane to prepare a 136 solution of 0.1 M DC18C6. Ten mL aqueous solution were combined with 15 mL 137 organic solution in a glass vial, which was sealed. The two phases were stirred for 30 138 minutes using a Teflon-coated magnetic bar. After equilibrium was achieved, the two 139 phases were separated by centrifugation at 2,000 rpm for 1 min, and an aliquot of the 140 aqueous solution taken for analysis. A 10 mL aliquot of the organic solution was saved 141 for back extraction of Cu in 15 mL of 0.02 M HCl. The extracted Cu was fully (>98%) 142 recovered in this procedure. All these steps were carried out at 298 K. Copper 143 concentrations in the equilibrated aqueous phase and the back extraction solution were determined by inductively-coupled plasma atomic-emission spectrometry (Thermo 144

145 Fisher Scientific, iCAP 6300 Duo). Aliquots of these solutions were saved for isotopic146 analysis.

147 Organic impurities carried over from the extraction step were separated from 148 Cu on AGMP1 anion exchange resin in 7 M HCl as described in Maréchal et al. (1999). 149 Copper isotopic compositions were measured on a Thermo-Finnigan Neptune 150 MC-ICP-MS at Washington University in St. Louis. The samples were introduced by 151 free aspiration in 0.1 M HNO₃ using a Teflon microcentric nebulizer (uptake rate of 100 152 μ L/min) and a glass cyclonic spray chamber. Masses 63 and 65 were positioned on the 153 L2 and axial collectors, respectively. Intensities of Zn isotopes (64, 66, 67 and 68) were measured on the collectors L1, H1, H2, H3, respectively. The intensity of ⁶²Ni was 154 155 monitored in order to correct for the potential interference of ⁶⁴Ni. Instrumental mass 156 fractionation was corrected by both Zn doping and standard-sample bracketing 157 (Maréchal et al., 1999; Albarède, 2004). The typical external reproducibility of replicate 158 analyses of the same samples carried out during different analytical sessions is 0.10% (2 σ) for δ^{65} Cu. 159

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161 **2.2. Computational methods**

Orbital geometries and vibrational frequencies of aqueous Cu(II) species were computed using the density functional theory (DFT) as implemented by the Gaussian09 code (Frisch et al., 2009; Dannington et al., 2009). The DFT method employed here is a hybrid density functional consisting of Becke's three-parameter non-local hybrid exchange potential (B3) (Becke, 1993) with Lee-Yang-and Parr (LYP) (Lee et al., 1988) non-local functionals. Using the 6-311+G(d,p) basis set or higher is recommended for calculating the Cu complexes by de Bruin et al. (1999). The 6-311+G(d,p) basis set, which is an all-electron basis set, was therefore chosen for H, C, O, S, Cl, and Cu. Molecules were modeled without any forced symmetry. An 'ultrafine' numerical integration grid was used and the SCF (self-consistent field) convergence criterion was set to 10^{-9} .

173 The isotope enrichment factor due to intramolecular vibrations can then be 174 evaluated from the reduced partition function ratio (s/s')f (Bigeleisen and Mayer, 1947), 175 also noted β ,

$$ln\frac{s}{s'}f = \sum \left[lnb(u_i') - lnb(u_i)\right]$$
⁽²⁾

176 where

$$\ln b(u_i) = -\ln u_i + \frac{u_i}{2} + \ln(1 - e^{-u_i})$$
⁽³⁾

177 and

$$u_i = \frac{hv_i}{kT} \tag{4}$$

178

179 in which v stands for vibrational frequency, s for the symmetry number of the Cu 180 compound, h the Planck constant, k the Boltzmann constant, and T the absolute 181 temperature. The subscript *i* denotes the *i*th mode of molecular vibration, and primed variables refer to the light isotopologue. The isotope enrichment factor due to molecular 182 183 vibrations can be evaluated from the frequencies (v) summed over all the different 184 modes. The isotopic difference in the stability constant of chemical reactions is identical 185 to the difference of $\ln \beta$ between related species. For example, a chemical exchange 186 reaction,

$$\operatorname{Cu}^{2^+} + \operatorname{Cl}^- \rightleftharpoons \operatorname{Cu}\operatorname{Cl}^+, \quad K_{CuCl^+}$$
(5)

188 with stability constant K_{CuCl^+} , the isotope fractionation between the hydrated Cu²⁺ and 189 CuCl⁺ is,

$$\ln \frac{K_{CuCl^{+}}(^{65}Cu)}{K_{CuCl^{+}}(^{63}Cu)} = \ln \frac{[^{65}CuCl^{+}]/[^{63}CuCl^{+}]}{[^{65}Cu^{2+}]/[^{63}Cu^{2+}]}$$
$$= \ln \beta_{CuCl^{+}} - \ln \beta_{Cu^{2+}}$$
(6)

191

190

192 Isotope fractionation also results from the mass-independent nuclear field shift effect (Bigeleisen, 1996; Nomura et al., 1996, Fujii et al. 2009) which is an 193 194 isotope-dependent shift reflecting that the electronic orbitals do not vanish at the 195 nucleus and depends on the nuclear size and shape (King, 1984). The latest version of 196 quantum chemical models have taken finite-size nuclei into account in ab initio 197 calculations (Schauble, 2007; Abe et al., 2008, 2010; Fujii et al., 2010, 2011a,c,d). For 198 light elements such as Ni (Fujii et al., 2011a) and Zn (Fujii et al., 2010), the nuclear 199 field shift effect was found to be small as 0.02-0.03‰/amu. The isotope enrichment 200 factor via the nuclear field shift $(\ln K_{fs})$ is given by

201

$$\ln K_{\rm fs} = \frac{hc}{kT} v_{fs} \tag{7}$$

202

203 where v_{fs} is the nuclear field shift and *c* the velocity of light.

204The contribution of the nuclear volume was estimated by the Dirac-Coulomb205Hartree-Fock (DCHF) method implemented in the UTChem program (see references in

Abe et al., 2008, 2010). We calculated the electronic structure of $Cu^0([Ar]3d^{10}4s^1)$. 206 $Cu^{+}([Ar]3d^{10}), Cu^{2+}([Ar]3d^{9})$ and a few other simple Cu molecules. The Cartesian 207 208 coordinates of Cu molecules optimized by the Gaussian09 code were supplied to the 209 DCHF calculation. Exponents of basis sets were taken from Faegri's four-component 210 basis for Cu (Faegri, 2001), third-order Douglas-Kroll basis for H and Cl (Tsuchiya et 211 al., 2001), and ANO-RCC for O (Roos et al., 2005). Contraction coefficients were 212 optimized by four-component atomic calculations (Koc and Ishikawa, 1994). After adding some diffuse and polarization functions, the final contraction form of the large 213 214 component basis sets was (19s14p9d)/[6s4p3d] for Cu, (16s11p1d)/[4s3p1d] for Cl, 215 (14*s*9*p*2*d*)/[3*s*2*p*2*d*] for O, and (8*s*2*p*)/[3*s*2*p*] for H.

The effect of nuclear spin can be neglected at equilibrium (Bigeleisen, 1996). Since the present crown ether experiments were run at equilibrium, the kinetic part of this effect also known as the magnetic isotope effect (Epov et al., 2011; Epov, 2011), will not be considered in this work.

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- 221

3. RESULTS AND DISCUSSION

3.1. Isotope fractionation of Cu by crown ether extraction

The Cu(II) species in HCl medium in the laboratory experiment are Cu²⁺, CuCl⁺, CuCl₂, and CuCl₃⁻ (Brugger et al., 2001), which are related through complexation reaction 5 and the following stepwise reactions, with stability constants K_{species} ,

$$\operatorname{CuCl}^{+} + \operatorname{Cl}^{-} \rightleftarrows \operatorname{CuCl}_{2}, \quad K_{CuCl}, \tag{8}$$

227 and

$$\operatorname{CuCl}_2 + \operatorname{Cl}^- \rightleftarrows \operatorname{CuCl}_3^-, \quad K_{CuCl_3^-}$$
(9)

The presence of $CuCl_4^{2-}$ was reported for highly concentrated HCl and $CuCl_2$ systems (Bell et al., 1973; Neilson, 1982; Tanimizu et al., 2007) or high-temperature systems (Collings et al., 2000) but this higher order of complexation is negligible at the low HCl concentrations and ambient temperature of the present experiments (Brugger et al., 2001).

234

$$CuCl_2 + L \rightleftharpoons CuLCl_2, \tag{10}$$

(10)

235

236 where L stands for DC18C6. The distribution ratio *D* is written as,

$$D = \frac{\Sigma[\operatorname{Cu}(\operatorname{II})]_{\operatorname{org}}}{\Sigma[\operatorname{Cu}(\operatorname{II})]_{\operatorname{aq}}} \approx \frac{[\operatorname{Cu}\operatorname{LCl}_2]}{[\operatorname{Cu}^{2^+}] + [\operatorname{Cu}\operatorname{Cl}^+] + [\operatorname{Cu}\operatorname{Cl}_2] + [\operatorname{Cu}\operatorname{Cl}_3^-]}$$
(11)

237

where the subscripts 'org' and 'aq' stand for the organic and aqueous phases,respectively.

240 The isotope separation factor α , between the aqueous and organic phases, is 241 defined as:

$$\alpha = \frac{\sum_{k} \phi_{\text{org}}^{k} ({}^{65}\text{Cu}/{}^{63}\text{Cu})_{\text{org}}^{k}}{\sum_{k} \phi_{\text{aq}}^{k} ({}^{65}\text{Cu}/{}^{63}\text{Cu})_{\text{aq}}^{k}}$$
(12)

where *k* refers to the *k*th compound in the organic and aqueous phases, respectively, and φ is the fraction of total Cu (proper ⁶³Cu) hosted in the *k*th compound. The isotope enrichment factor is defined as $\alpha - 1$. Because α is always close to unity, $\alpha - 1 \approx \ln \alpha$, and 1000 ln α therefore can be approximated by Δ^{65} Cu = δ^{65} Cu_{org} - δ^{65} Cu_{aq}. The Δ^{65} Cu values obtained are shown in Table 1 together with distribution ratios. Δ^{65} Cu ranged from -1.06 to -0.39‰.

249 The yield of Cu(II) extraction by crown ethers has been investigated by Yoshio 250 et al. (1980), Nakamura et al. (1982), and Contreras et al. (1993). A study on extraction 251 kinetics showed that equilibrium is achieved within 30 sec (Nakamura et al., 1982). 252 Hydration-dehydration is known to be the rate-limiting step of solvent extraction. The rate constant for H_2O substitution in the inner coordination sphere of Cu^{2+} is quite high 253 in the transition metal series, and it is comparable with that of Li^+ of the alkali metals as 254 observed by Eigen (1963). The isotope exchange rate kinetics of ⁷Li/⁶Li in the crown 255 256 ether systems has been studied in detail (Jepson and Cairns, 1979; Nishizawa et al., 1984), in which the isotopic equilibrium is achieved within 30-60 sec after contacting 257 258 aqueous and organic phases. The 30 min stirring period in the present study is therefore considered to be enough to achieve the isotopic equilibrium of Cu. The Δ^{65} Cu values 259 260 listed in Table 1 are hence attributable to the equilibrium isotope effect.

Figure 1a shows the mole fractions of Cu species estimated by using the stability constants of reactions 5, 8, and 9 (Brugger et al., 2001). Δ^{65} Cu was found to vary with HCl molarity (Fig. 1b), a trend that reflects the isotope fractionation of Cu in reactions 5 and 8-10. With increasing HCl molarity, more CuCl₂ forms, which in turn results in an increase of *D*. The magnitude of Δ^{65} Cu shown in Table 1 is similar to that found in other ligand exchange systems (Matin et al., 1998; Maréchal et al., 1999, 2002;
Zhu et al., 2002).

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269 **3.2.** Contribution of the mass-independent isotope effect

270 The total electronic energies calculated for Cu isotopes and isotopologues are shown in atomic units (a.u.) in Table 2, in which Cu^{2+} was chosen as the reference. The 271 maximum effect was found to be 0.015‰ for the Cu⁰-Cu(I) pair and 0.012‰ for the 272 Cu(I)-Cu(II) pair. The solvent extraction reaction is controlled by ligand exchange of 273 274 Cu(II). Based on the Cu(II) data in Table 2, the nuclear field shift effect in Cu(II) species is only ~3% of the smallest Δ^{65} Cu value listed in Table 1. The values of Δ^{65} Cu 275 276 reported in Table 1 are the values for the mass-dependent isotope fractionation and reflect intramolecular vibrations. 277

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279 **3.3. Mass-dependent isotope effects**

280 3.3.1. Hydrated Cu(II) ions

281 The hydrated Cu(II) ion is generally represented as the hexaaqua complex $Cu(H_2O)_6^{2+}$ (Fig. 2). It is known that $Cu(H_2O)_6^{2+}$ shows the Jahn-Teller distortion effect 282 283 (Sherman, 2001; Bersuker, 2006). Two Cu-O distances of the vertical axial bond 284 (Cu-Oax) are longer than four Cu-O distances in the equatorial plane (Cu-Oeq). The Jahn-Teller effect lowers the symmetry of $Cu(H_2O)_6^{2+}$ from octahedral T_h to D_{2h} . The 285 286 calculated bond distance of Cu-O and the literature values are shown in Table 3. The bond distances were calculated to be 2.02-2.03 Å (Cu-Oeq) and 2.30 Å (Cu-Oax), which 287 288 are consistent with those reported in the literature (see references in Table 3).

289 After evidence was found that the stable aqueous complex has a fivefold coordination (Pasquarello et al., 2001), the Jahn-Taller effect of $Cu(H_2O)_6^{2+}$ in aqueous 290 solutions has been questioned (Chabov et al., 2006). The $Cu(H_2O)_5^{2+}$ complex has five 291 identical Cu-O bonds with a length of 1.96 Å, which was identified by neutron 292 293 diffraction (ND) analysis combined with first-principles molecular dynamics (MD). The 294 common bond distance reflects the rapid switch between the square pyramid and 295 trigonal bipyramid configurations (Pasquarello et al., 2001; de Almeida et al., 2008). In 296 a Car-Parrinello MD simulation (Amira et al., 2005), a square pyramidal geometry with 297 four short (2.00 Å) and one long (2.45 Å) Cu-O bond could be optimized. A similar set 298 of bond distances, four short (1.97 Å) and one long (2.39 Å) Cu-O bond, has been 299 confirmed in an X-ray absorption study (Benfatto et al., 2002). Our calculation of the $Cu(H_2O)_5^{2+}$ geometry also resulted in the square pyramid rather than the trigonal 300 301 bipyramid of Fig. 2 and Table 4.

There is no clearly predominant structure among the four-, five-, and six- fold coordinated Cu(II) ions (Chaboy et al., 2006). In the present study, we tested both the coordination numbers five and six. We also tested a coordination number four to evaluate the possibility of steric hindrance and strong degree of covalency. The optimized structures in Cartesian coordinates are given in Table S1 of the electronic supplement.

308 The intramolecular vibrational frequencies were then calculated for aqueous 309 Cu(II) species with optimized geometries. The ln β [= ln (*s/s'*)*f*] values for the isotope 310 pair ⁶⁵Cu/⁶³Cu were determined by employing Eqs. 2-4 and are listed in Table 5. The 311 value of ln β of Cu(H₂O)₅²⁺ at 298 K is 0.26‰ larger than that of Cu(H₂O)₆²⁺.

313 3.3.2. Hydrated Cu(II) chlorides

The structure of the hydrated Cu(II) chloride $CuCl_m(H_2O)_n^{2-m}$ is shown in 314 315 Tables 6 and S1 together with literature values. For the Cu(II) *monochloride*, CuCl⁺, 316 both fivefold (D'Angelo et al., 1997) and sixfold (Rode and Islam, 1992; Texler et al., 1998) coordination have been reported. The bond distances obtained for the 317 fivefold-coordinated $CuCl(H_2O)_4^+$ reproduce the experimental results of D'Angelo et al. 318 (1997). Though the geometry of sixfold-coordinated $CuCl(H_2O)_5^+$ was optimized within 319 320 the SCF convergence criterion, this species still showed a few imaginary (negative) 321 frequencies. Tighter convergence criteria may be needed.

The Jahn-Teller effect should be detectable in the sixfold-coordinated Cu(II) *dichloride*, CuCl₂(H₂O)₄. For a geometry with an axial Cl-Cu-Cl bond, we found two long and two short Cu-O bonds in the equatorial plane (Fig. 2). Our calculation agreed with the bond distances determined experimentally (Neilson, 1982; Tajiri and Wakita, 1986). A fivefold coordinated structure was inferred from the experiments of Ansell et al. (1995), and, indeed, we found evidence for a stable fivefold-coordinated CuCl₂(H₂O)₃ structure with a distorted square pyramid (Fig. 2).

329 The X-ray diffraction study of Bell et al. (1973) reports the existence of 330 sixfold-coordinated Cu(II) *trichloride*, CuCl₃(H₂O)₃, which is an average structure of polymerized Cu(II) chlorides in a concentrated CuCl₂ solution. The complex has three 331 332 Cu-Cl and three Cu-O bonds, a very low symmetry, and is unstable. The isolated $CuCl_3^{-1}$ 333 anion in solution may have a different structure. There are no reports in the literature of 334 the fivefold-coordinated CuCl₃(H₂O)₂⁻. Convergence could not be achieved for the square pyramidal nor the trigonal bipyramidal geometry. A smaller coordination 335 number was found for hydrated CuCl₃⁻ (Collings et al., 2000). Under highly 336

concentrated HCl and CuCl₂ conditions, a higher order chlorinated CuCl₄^{2–} exists. It is disputed whether the structure is distorted octahedral CuCl₄(H₂O)₂^{2–} or tetrahedral CuCl₄^{2–} (see references in Table 6). Systematic analysis by the extended X-ray absorption fine structure (EXAFS) technique and Raman spectrometry suggested that a possible structure is the tetrahedral CuCl₄^{2–} (Tanimizu et al., 2007). This supports the existence of fourfold-coordinated CuCl₃H₂O[–]. The calculation of CuCl₃H₂O[–] successfully converged.

344 The ln β values of the hydrated chlorides are shown in Fig. 3 and Table 5. As for the hydrated Cu^{2+} case, ln β (298 K) of the fivefold-coordinated $CuCl_2(H_2O)_3$ was 345 346 found to be 0.26‰ larger than that of the sixfold-coordinated CuCl₂(H₂O)₄. A difference of this magnitude is assigned to a stereochemical effect. For the fivefold 347 coordination, the ln β value at 298 K for Cu(H₂O)₅²⁺ is 0.56‰ larger than that of 348 349 $CuCl_2(H_2O)_3$ (Table 5). Likewise, for the sixfold coordination, the ln β value at 298 K for $Cu(H_2O)_6^{2+}$ is also 0.56% larger than that of $CuCl_2(H_2O)_4$ (Table 5). This suggests 350 that, with the same coordination number, exchanging H₂O by Cl⁻ enhances 65 Cu/ 63 Cu 351 fractionation by approximately the same amount. 352

353

As shown in Fig. 3, ln β decreases when the fraction of H₂O exchanged with Cl⁻ increases. The variation of Δ^{65} Cu estimated from the ln β values (298 K) with the molarity of HCl is shown in Fig. 1b. A similar observation was previously made for Ni and Zn isotope fractionation (Fujii et al., 2010, 2011a).

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The isotopic mass balance is preserved in the extraction system as,

$$\frac{\Sigma[^{65}Cu]}{\Sigma[^{63}Cu]} = \frac{[^{65}Cu^{2+}] + [^{65}CuCl^{+}] + [^{65}CuCl_{2}] + [^{65}CuCl_{3}^{-}] + [^{65}CuLCl_{2}]}{[^{63}Cu^{2+}] + [^{63}CuCl^{+}] + [^{63}CuCl_{2}] + [^{63}CuCl_{3}^{-}] + [^{63}CuLCl_{2}]}$$
(13)

The mole fractions of related Cu(II) species are evaluated from the values of K_{CuCl^+} , 360 K_{CuCl_2} , K_{CuCl_3} (Brugger, 2001), and D (Eq. 11), all obtained experimentally. Each mole 361 fraction is the sum of fractions of ⁶³Cu and ⁶⁵Cu isotopologues. The isotopic fractions 362 within the species were estimated from $\ln \beta$. Since there is no structural information of 363 the Cu-DC18C6 complex in organic solution, β_{CuLCl_2} was set as a free parameter. The 364 results of the calculation are shown in Fig. 1b. The value of ln $\beta_{CuLCl_{2}}$ converged to 365 3.35‰. The bold and dotted curves shown in Fig. 1b are the values of Δ^{65} Cu calculated 366 for ln β_{CuLCl_2} = 3.35 ± 0.05‰. The results agree well with the experimental results. An 367 368 important aspect of the estimation is that the calculation correctly predicts the sigmoidal 369 dependence of $\ln \beta$ on the fraction of H₂O exchanged by Cl⁻ with increasing HCl 370 molarity.

371

372 **3.4. Application to hydrous geochemistry**

373 In the absence of organic ligands which are known to dominate Cu speciation in seawater (Moffet, 1995; Moffet and Brand, 1996; Hirose, 2006) the free Cu²⁺ ion is 374 dominant in freshwater, while the carbonate complexes $CuCO_3$ and $Cu(CO_3)_2^{2-}$ are 375 dominant in seawater (Albarède, 2004). Copper concentration in seawater increase 376 377 down the water column, which indicates both the uptake of Cu by plankton and 378 scavenging of Cu by settling particles (Boyle et al., 1977; Bruland, 1980). Dissolved Cu in seawater is isotopically heavy (δ^{65} Cu = +0.9 to +1.5‰) relative to rocks and 379 380 sediments (0.0 to +0.3%) and the riverine input (+0.69%) (Bermin et al., 2006; Vance 381 et al., 2008). It was suggested that this result is attributable to the partitioning of Cu isotopes between a heavy dissolved phase, where Cu is bound to organic ligands, and a light particulate phase, dominated by Fe-Mn oxides (Vance et al., 2008). δ^{65} Cu in Fe-Mn nodules (Albarède, 2004) and encrustations (Little et al., 2010) from the three major ocean basins varies between +0.04 and +0.6‰, which is significantly lighter than the δ^{65} Cu value in seawater (+0.9 to +1.5‰, Vance et al., 2008).

387 Surface seawater also contains various inorganic complexes and carbonates. 388 Speciation of Cu in seawater is very sensitive to the hydrolysis constants. Zirino and Yamamoto (1972) used the cumulative formation constant of Cu(OH)₂, log $\beta_2 = 14.3$, 389 which, as shown in Fig. S1a (see the electronic supplement), makes Cu speciation in 390 391 seawater dominated by Cu(OH)₂. Mole fractions have been calculated from the set of 392 stability constants recommended by Powell et al. (2007) with the second hydrolysis constant log $*\beta_2 = -16.65$, and the results are shown in Fig. 4a as a function of pH. At a 393 typical seawater pH of 8.22 (Macleod et al., 1994), CuCO₃ is the dominant species, 394 while the hydrated Cu^{2+} ion, chloride, sulfate, and hydroxide are subordinate. At pH \geq 395 8.22, $Cu(CO_3)_2^{2-}$ becomes an abundant species. These calculations are only relevant to 396 397 seawater with no organic ligands.

Figure 4b shows the isotope fractionation of Cu(II) among its various species as estimated from their mole fractions and ln β (Tables 5 and 7) at 298 K under conditions typical of seawater. The optimized geometries of the relevant species are shown in Fig. 2 and Table S1 and the temperature dependence of ln β (Table 7) in Fig. 5. The geometry of Cu hydroxides, CuOH⁺ and Cu(OH)₂, were calculated by dissociating H⁺ ions from the *fivefold*-coordinated Cu(H₂O)₅²⁺. The positions of OH⁻ ligands were determined by reference to CuCl⁺ and CuCl₂ (Fig. 2). The structure calculated for both 405 $CuOH(H_2O)_4^+$ and $Cu(OH)_2(H_2O)_3$ converged to a distorted square pyramid. With a 406 H_2O molecule at the vertex of the square pyramid, the bond distance is shorter for 407 $Cu-OH^-$ (1.8-1.9 Å) than for Cu-OH₂ (2.3-2.4 Å) (see Table S1). This suggests that the 408 stronger Cu-O bonds on the square plane loosen the hydration bond from the vertex. 409 This conclusion is consistent with the known structure of amorphous Cu(OH)₂ or 410 $Cu(OH)_2$ colloidal solution, which both show a layer structure of polymerized square 411 planes (Elizarova et al., 1999; Kriventsov et al., 1999).

412 Fourfold complexation occurs in both amorphous and crystalline Cu(OH)₂. A 413 stable fourfold complexation was also obtained for the Cu(II) carbonates, CuCO₃(H₂O)₂ and $Cu(CO_3)_2^{2-}$ (Fig. 2 and Table S1). CO_3^{2-} was treated as a bidentate ligand. The 414 415 structure converged to a distorted square plane. We tested positioning a water molecule 416 above the plane in the initial structure, but it moved out of the inner coordination shell 417 during the calculation. The structural optimization of CuCO₃OH⁻ was more problematic. Since CO₃²⁻ and OH⁻ occupy three coordination positions, one water molecule should 418 419 also bind to Cu(II) at the fourth coordination position, thereby giving the 420 CuCO₃(OH)H₂O⁻ compound. It was actually found that no water molecule is stable in 421 the inner coordination shell. From the stoichiometry, the stable model molecule may instead be $CuHCO_3(OH)_2^-$. This stable structure, in which HCO_3^- acts as a 422 423 quasi-bidentate ligand is shown in Fig. 2 and Table S1. The $\ln \beta$ values obtained are 424 shown in Fig. 5 and Table 7. The Cu hydroxides and carbonates have $\ln \beta$ values of 4.3-5.2‰ (298 K), in which compounds with higher complexation with OH^- and CO_3^{2-} 425 resulted in $\ln \beta \ge 5\%$ (298 K). 426

427 The structure of hydrated Cu(II) in sulfate solutions was previously 428 investigated by Ohtaki et al. (1976) and Musinu et al. (1983). In CuSO₄ solutions, Cu^{2+} 429 is surrounded by disordered ligands (Musinu et al., 1983), which suggests a direct interaction of Cu^{2+} with SO_4^{2-} . We used the structural model of Fe(III) sulfate proposed 430 by Magini (1979) as a reference for $CuSO_4(H_2O)_4$ (Fig. 2 and Table S1), in which SO_4^{2-} 431 432 is treated as a monodentate ligand, and obtained bond distances of 1.9-2.1 Å for Cu-OH₂ and 2.37 Å for Cu-OSO₃^{2–}. These values agree with the Cu-O bond distances 433 determined by X-ray diffraction (Musinu et al., 1983). The electron donor SO_4^{2-} 434 strongly attracts H^+ and the resulting structure is equivalent to $CuH_2SO_4(OH)_2$ (Fig. 2). 435 436 The magnitude of $\ln \beta$ for CuSO₄ is 5.14‰ at 298 K (Table 7) and matches that of $Cu(CO_3)_2^{2-}$ (Fig. 5). 437

438 As shown in Figs. 4a and 4b, at $pH \ge 8.22$, the dominant Cu(II) species are carbonates. We found that Cu in $Cu(CO_3)_2^{2-}$ (and $CuSO_4$) is isotopically 0.6% heavier 439 than Cu^{2+} ion. At pH = 8.22, the $\delta^{65}Cu$ value of CuCO₃ is slightly negative -0.17‰, 440 whereas those of $Cu(CO_3)_2^{2-}$ and $CuCO_3OH^-$ are substantially heavier, 0.75% and 441 442 respectively. At high pH, the relevant fractionation 0.58‰, factors are Δ^{65} Cu[Cu(CO₃)₂²⁻-CuCO₃] = 0.92‰ and Δ^{65} Cu[CuCO₃OH⁻-CuCO₃] = 0.75‰. Copper 443 in carbonates, whether in CuCO₃ or in solid solutions with carbonates of other elements 444 445 should therefore be isotopically light with respect to dissolved Cu.

At pH = 8.22, Cu(OH)₂ is +0.48‰ heavier than bulk seawater Cu (Fig. 4b). A caveat is the somewhat different fractionation obtained when the cumulative formation constant of Cu(OH)₂, log β_2 =14.3 of Zirino and Yamamoto (1972) is used. The major species are Cu(OH)₂ and CuCO₃, while mole fractions of other species are insignificant (Fig. S1a, Zirino and Yamamoto, 1972). The δ^{65} Cu shift of Cu(OH)₂ and CuCO₃ at pH = 8.22 with respect to dissolved Cu are +0.02‰ and -0.62‰, respectively (Fig. S1b).

453

Isotope fractionation may therefore not be seen in Cu(II) hydroxide, but Cu in CuCO₃ should nevertheless remain isotopically light with respect to the seawater value.

454 As sulfides have small $\ln \beta$ values (Fujii et al., 2011b; Pons et al., 2011), 455 sulfate-sulfide and carbonate-sulfide exchange are possibly determinant reactions for Cu isotope fractionation in seawater. $CuHS(H_2O)_4^+$ and $Cu(HS)_2(H_2O)_3$ were calculated 456 457 with the geometries shown in Table S1. The structures that successfully converged were 458 similar to those of the hydroxides shown in Fig. 2. The ln β values are shown in Fig. 5 459 and Table 7. It is clear that the $\ln \beta$ values of Cu(II) sulfides are smaller than those of all 460 the species shown in Fig. 3. In particular, the ln β values of Cu sulfides are 1.9-2.0% 461 smaller than those of Cu sulfate and carbonates. Diagenetic Cu(II)-bearing sulfides are 462 expected to be isotopically lighter than dissolved Cu, while Cu in the water table of 463 sulfide-rich terranes should be heavy (Mathur et al., 2005, 2012).

Isotope fractionation also occurs under reducing conditions through Cu(I)-Cu(II) redox reactions. The ln β values calculated for Cu(I) chlorides and hydrogensulfides are $\leq 2.6\%$ at 298 K (Seo et al., 2007), which suggests that Cu(I) complexes preferentially enrich ⁶³Cu and their precipitation results in the enrichment of ⁶⁵Cu in the fluid.

469 Cu in sulfides should be isotopically light with respect to dissolved aqueous Cu, 470 and the main form of Cu removal from aquatic systems is Cu sulfide. The heavy 471 isotopic composition of Cu in surface seawater (Vance et al., 2008) and groundwater 472 (Mathur et al., 2012) therefore likely reflects that Cu carbonate complexes are held back 473 in solution, while Cu is precipitated in sulfides. Complexation of Cu(II) with organic 474 matter in surface seawater (Moffett, 1995) should also cause Cu isotope fractionation. 475 Evidence for the large-scale deposition of sediments with isotopically light Cu required476 by mass balance is, however, still missing.

477

478 **3.5. Application to biological activity**

479 A variety of biological processes may induce Cu isotope fractionation. 480 Pokrosvky et al. (2008) demonstrated that Cu in bacteria is isotopically light relative to 481 the ambient solution. Pokrovsky et al. (2012) further identified S-coordinated Cu(I) 482 complexes at the surface and inside bacterial cells that may preferentially concentrate ⁶³Cu over ⁶⁵Cu. It is therefore appealing to assign isotope fractionation in biological 483 484 material to competing Cu(II) and Cu(I) complexation. Mammal physiology is also a 485 cause of Cu isotope fractionation. Albarede et al. (2011a) found a ~1‰ fractionation between human erythrocytes and serum. High δ^{65} Cu values of 1.5‰ were found in the 486 487 kidney of both sheep (Balter and Zazzo, 2011) and mice (Albarède et al., 2011b) with 488 respect to the rest of the body.

489 Oxalic acid is an ubiquitous toxic organic acid in body fluids. Oxalate is 490 adsorbed in the intestinal track, but the question of the origin of high oxalate contents in 491 urine and plasma found in patients prone to kidney damage is still contentious. This is a 492 genuine medical concern as ascorbate (vitamin C) supplementation has been argued to 493 increase the urinary oxalate levels, and therefore ascorbate may be a risk factor for 494 individuals predisposed to kidney stones (Chai et al., 2005; González et al., 2005; 495 Massey et al., 2005). Ascorbate is efficiently converted to oxalate when the coexisting 496 copper concentration is high (Hayakawa et al., 1973).

497 Hayakawa et al. (1973) suggest that Cu(II) is readily reduced by ascorbic acid
498 (H₂A) to Cu(I),

$$H_2A + 2Cu(II) \rightarrow A + 2Cu(I) + 2H^+$$
(14)

The ascorbate anion A^{2-} here is oxidized to A, dehydroascorbic acid $C_6H_6O_6$. In the 500 501 catalytic reaction by a multicopper oxidase, ascorbate oxidase, the oxidation of 502 ascorbate to dehydroascorbate occurs via the disproportionation of the 503 semidehydroascorbate radical (Solomon et al., 1996). The enzymatic reactions may be 504 identical to the catalytic reaction 14.

The monohydroascorbate anion HA⁻ forms a complex with Cu²⁺, CuHA⁺.

$$Cu^{2+} + HA^{-} \rightleftharpoons CuHA^{+}, \quad K_{CuHA^{+}}$$
 (15)

506

- 507 Degradation of ascorbate via dehydroascorbate forms oxalate. A pathway of ascorbate508 degradation has been proposed by Green and Fry (2005).
- 509 The oxalic acid also forms a complex with Cu^{2+} , CuC_2O_4 .

$$Cu^{2+} + C_2O_4^{2-} \rightleftharpoons CuC_2O_4, \quad K_{CuC_2O_4}$$
 (16)

511 The value of the standard redox potential Eh with respect to the standard hydrogen 512 electrode (SHE), E_0 , for the Cu redox reaction,

$$Cu^+ \neq Cu^{2+} + e^- \tag{17}$$

513

514 is $E_0 = 0.153 + 0.591 \log ([Cu^{2+}]/[Cu^{+}])$ (Pourbaix, 1974). We used the following acid 515 dissociation constants and equilibrium constants $pK_a = 4.03$ (ascorbic acid), $pK_{a1} =$ 516 1.252, $pK_{a2} = 4.266$ (oxalic acid) (Smith and Martell, 1989), $\log K_{CuHA^+} = 2.32$

(Jameson et al., 1976), and log $K_{CuC_2O_4} = 4.5$ (Foreci et al., 1995). Figure 6a shows the 517 mole fractions of Cu^+ , Cu^{2+} , $CuHA^+$, and CuC_2O_4 as functions of Eh. The total 518 concentrations of Cu and ascorbic acid are $\sim 10^{-6}$ M (Lech and Sadlik, 2007) and $\sim 10^{-4}$ 519 520 M (Margolls and Davis, 1988), respectively. The pH and Eh in the body are known to 521 be ~7.4 and 0.27 V (van Rossum and Schamhart, 1991). These values were used to 522 estimate the mole fractions of Cu species using the assumption that 10% of ascorbic 523 acid is converted to oxalic acid. We further assumed that no other degradation products 524 were present. Figure 6b shows the speciation as a function of the reaction progress. As 525 expected, oxidation of Cu(I) to Cu(II) proceeds with increasing Eh, which in turn 526 enhances Cu-ascorbate and Cu-oxalate formation. The degradation of ascorbate 527 increases oxalic acid concentration and therefore promotes the production of 528 Cu-oxalate.

529 Figure 6c and 6d show the range of isotope fractionation among Cu species as 530 estimated from their mole fractions and ln ß at 298 K (Tables 5 and 8). The optimized geometries of related species are shown in Table S1. Hydrated Cu⁺ was treated as free 531 532 Cu(I). Cu(I) complexes possess simple linear structures (Fulton et al., 2000a,b). Our ln β value of 2.87‰ at 289K for Cu(H₂O)₂⁺ (Table 8) is consistent with the value of 533 2.89‰ estimated by *ab initio* methods for CuClH₂O (Seo et al., 2007). Cu(H₂O)₅²⁺ was 534 535 treated as free Cu(II), and its $\ln \beta$ is given in Table 5. For the geometry of Cu(II) ascorbate CuHA⁺, we used the copper monomer (Ünaleroğlu et al., 2001). The structure 536 537 calculated for Cu(II) in fivefold coordination with L-ascorbate and H₂O is shown in Fig. 538 2 and Table S1. The value of ln β calculated at 298 K is 3.32‰. The ln β value of 539 D-ascorbate at 298 K (also 3.38‰) is indistinguishable from that of L-ascorbate. The

geometry of Cu(II) oxalate was quite similar to the structure of the hydrated Cu carbonate. It successfully converged to a distorted plane configuration (Fig. 2 and Table S1) and led to large ln β values (Fig.7 and Table 8).

Figures 6c and 6d show the δ^{65} Cu values of the different species relative to the 543 544 bulk solution as a function of Eh and extent of oxalate formation. Figure 6c shows that, 545 with respect to free Cu(II), free Cu(I) is enriched in the light isotope. When Cu(I) and Cu(II) coexist, the δ^{65} Cu of the total free Cu ions is only slightly lighter than the sum of 546 all the Cu species. The δ^{65} Cu of the Cu ascorbate varies from -1.0 to +0.5% when Eh 547 548 increases from -1 V to +1 V, but its mole fraction remains very small. The prominent 549 feature of Fig. 6c is the heavy isotope enrichment (+0.6 to +2.5%) of the Cu oxalate 550 relative to total Cu. Figure 6d was drawn for Eh = 0.27 V, which is a typical Eh value of 551 human blood. It is expected that degradation of ascorbate and excretion of oxalate 552 should leave isotopically heavy Cu in the kidney. With respect to food, which has a 553 δ^{65} Cu value of about 0‰ (Balter and Zazzo, 2011), when even trace amounts of oxalate form, it should leave behind copper with a δ^{65} Cu of ~1.4‰. This value is very close to 554 the δ^{65} Cu (of 1.5%) found in sheep (Balter and Zazzo, 2011) and mice (Albarède et al., 555 2011b) kidneys. The positive δ^{65} Cu found in kidneys may result from isotopically heavy 556 557 Cu left in the kidney by the degradation of ascorbic acid. Copper isotopes therefore 558 represent a potential marker of the troubles associated with hyperoxalury and kidney 559 stones.

- 560
- 561

4. CONCLUSIONS

562 We first demonstrated that Cu isotope fractionation at equilibrium is observed in laboratory-scale experiments. The effect on the ⁶⁵Cu/⁶³Cu ratio of Cu(II) partitioning 563 564 between HCl and DC18C6 depends on acid molarity. The effect is mainly governed by 565 the mass-dependent fractionations as a result of intramolecular vibrations. The nuclear 566 field shift effect accounts for less than 3% of the mass-dependent fractionation. We then 567 computed the coefficients of isotope fractionation of Cu for several species (hydrated 568 Cu ion, hydroxide, chloride, sulfide, sulfate, carbonate, oxalate, and ascorbate). It was 569 found that Cu in dissolved carbonates and sulfates is isotopically heavier than free Cu and Cu in sulfides. The theoretical estimation of δ^{65} Cu in ligand exchange between 570 571 inorganic ligands including carbonate anions may be useful to understand the heavy Cu 572 isotopic compositions found in both seawater and groundwater in the absence of strong organic ligands. Further theoretical estimation of δ^{65} Cu in hydrated Cu(I) and Cu(II) 573 574 ions, Cu(II) ascorbates, and Cu(II) oxalate suggests that Cu isotope fractionation during 575 the breakdown of ascorbate into oxalate results in the isotopically heavy Cu found in 576 mammal kidneys.

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- 953
- 954

967 Table 1 Isotopic fractionation of Cu(II) during
968 exchange experiments between HCl medium and a
969 macrocyclic complex.

macrocycnic con	npiex.	
[HCl] (M)	D	Δ^{65} Cu ^a (‰)
1	1.3×10^{-4}	-1.06
2	1.8×10^{-3}	-0.84
3	2.4×10^{-2}	-0.77
4	1.5×10^{-1}	-0.65
5	4.6×10^{-1}	-0.61
6	9.8×10^{-1}	-0.39
<i>a</i> =		

970 ^{*a*} Errors are $\pm 0.10\%$ for 2σ .

Table 2 Nuclear field shift effect of Cu isotopes and isotopologues at 29

1	a .	Total ene	ergy (a.u.)	ΔE (a.u.)	Nuclear Field
Valence	Species	⁶³ Cu	⁶⁵ Cu	(10^{-5})	shift effect (‰)
Cu(0)	Cu ⁰	-1653.451898480	-1653.451827445	7.1035	-0.014
Cu(I)	Cu^+	-1653.210583752	-1653.210512731	7.1021	0.001
	CuCl	-2114.481787065	-2114.481716036	7.1029	-0.007
	$\mathrm{CuH_2O}^+$	-1729.354508434	-1729.354437410	7.1024	-0.002
	CuOH	-1728.963440770	-1728.963369739	7.1031	-0.010
Cu(II)	Cu^{2+}	-1652.578696630	-1652.578625608	7.1022	0
	$CuCl^+$	-2114.155110496	-2114.155039471	7.1025	-0.003
	CuH_2O^{2+}	-1728.798913392	-1728.798842366	7.1026	-0.004
	CuOH^+	-1728.656622258	-1728.656551226	7.1032	-0.011
	CuO	-1728.336259331	-1728.336188299	7.1032	-0.011

981 ^{*a*} The nuclear field shift effect was calculated by $(\delta E_{\text{reference}} - \delta E_{\text{species}})/kT$ at T = 298K. 982 The root-mean-square charge radii $\langle r^2 \rangle^{1/2}$ reported for ⁶³Cu (3.8823×10⁻¹⁵ m) and ⁶⁵Cu 983 (3.9022×10⁻¹⁵ m) (Angeli, 2004) were used. Cu²⁺ was set as the reference. Energy of 984 photons of 1 a.u. is equal to 4.3597×10⁻¹⁸ J.

993 Table 3 Bond distances determined for $Cu(H_2O)_6^{2+}$.

	(= /*	
	Method ^{<i>a</i>}	Reference
2.02-2.03	MO	This study
2.06	MO	Beagley et al., 1989
2.05	MO	Åkesson et al., 1992
2.06-2.07	MO	Breza et al., 1997
2.033	MO	Bryantsev et al., 2008
1.97-2.09	MD	Liu et al., 2010
1.94	QM	van Duin et al., 2010
2.07	QM/MM MC	Marini et al., 1999
1.99-2.07	QM/MM MD	Schwenk and Rode, 2003a,b
1.955	EXAFS	Sham et al., 1981
2.00	EXAFS	Tajiri and Wakita, 1986
1.96	EXAFS	Nomura and Yamaguchi, 1988
1.99	EXAFS	Beagley et al., 1989
1.969-1.971	EXAFS	Fulton et al., 2000a,b
1.955	EXAFS	Persson et al., 2002
1.99	EXAFS+XANES	Benfatto et al., 2002
1.96	EXAFS+XANES	Chaboy et al., 2006
1.96	XANAS	Garcia et al., 1989
1.96	ND	Salmon et al, 1988
1.95	ND	Okan and Salmon, 1995
1.94	XRD	Ohtaki and Maeda, 1974
1.976	XRD	Magini, 1982
	$\begin{array}{c} 2.05\\ 2.06\text{-}2.07\\ 2.033\\ 1.97\text{-}2.09\\ 1.94\\ 2.07\\ 1.99\text{-}2.07\\ \hline 1.99\text{-}2.07\\ \hline 1.955\\ 2.00\\ 1.96\\ 1.99\\ 1.969\text{-}1.971\\ 1.955\\ 1.99\\ 1.96\\ 1.96\\ 1.96\\ 1.96\\ 1.95\\ 1.94\\ \end{array}$	(Å)Method2.02-2.03MO2.06MO2.05MO2.06-2.07MO2.033MO1.97-2.09MD1.94QM2.07QM/MM MC1.99-2.07QM/MM MD1.955EXAFS2.00EXAFS1.96EXAFS1.955EXAFS1.96EXAFS1.96EXAFS1.95ND1.96ND1.96ND1.96ND1.95ND1.95ND1.96XANAS1.96ND1.95ND1.94XRD

^a MO (molecular orbital), EXAFS(extended x-ray absorption fine structure), MD
 (molecular dynamics), ND (neutron diffraction), QM(quantum mechanics), MM
 (molecular mechanics), MC (Monte Carlo), XANES (x-ray absorption near-edge

⁹⁹⁷ structure), and XRD(x-ray diffraction).

Table 4 Bond distances determined for $Cu(H_2O)_5^{2+}$.

Cu-O _{long} (Å)	Cu-O _{short} (Å)	Method ^a	References
2.20	1.99-2.02	МО	This study
2.181	2.011	MO	Bryantsev et al., 2008
2.45	2.00	CPMD	Amira et al., 2005
2.22	2.04-2.07	CPMD	de Almeida et al., 2009
2.31	2.03-2.09	MD	Liu et al., 2010
2.34	1.963	EXAFS	Persson et al., 2002
2.39	1.97	EXAFS+XANES	Benfatto et al., 2002
2.36	1.96	EXAFS+XANES	Chaboy et al., 2006
_	1.96	ND	Pasquarello et al., 2001

^{*a*} CPMD (Car-Parrinello MD)

1050 Table 5 Logarithm of the reduced partition function, $\ln \beta$, for the pair ⁶⁵Cu-⁶³Cu.

1051	Hydrated Cu	(II) ions	and ch	lorides.

Crasica	_		Tempera	ature (K)		
Species	273	298	323	373	473	573
$Cu(H_2O)_5^{2+}$	5.355	4.546	3.905	2.968	1.876	1.290
$Cu(H_2O)_6^{2+}$	5.053	4.288	3.682	2.798	1.767	1.215
$CuCl(H_2O)_4^+$	4.906	4.161	3.572	2.712	1.711	1.176
$CuCl_2(H_2O)_3$	4.709	3.988	3.420	2.592	1.633	1.120
$CuCl_2(H_2O)_4$	4.397	3.724	3.193	2.421	1.525	1.046
$CuCl_3H_2O^-$	3.530	2.985	2.556	1.933	1.214	0.832

Table 6 Bond distances determined for $CuCl_m(H_2O)_n^{2-m}$.

Possible	т	10	Cu-Cl ^a	Cu-O ^a	Method	References
species	т	п	(Å)	(Å)	Method	Kelefences
$CuCl(H_2O)_4^+$	1	4	2.20	2.04-2.06(3) 2.35(1)	МО	This study
$CuCl(H_2O)_4^+$	1^b	4	2.29 ^b	1.968(3) 2.27(1)	EXAFS	D'Angelo et al., 1997
$CuCl(H_2O)_5^+$	1.1	5.2	2.55	2.00	MC	Rode and Islam, 1992
CuCl ₂ (H ₂ O) ₃	2	3	2.25	2.07-2.08(2) 2.35(1)	МО	This study
$CuCl_2(H_2O)_3$	2.0	3.3	3.1	1.96	ND	Ansell et al, 1995
CuCl ₂ (H ₂ O) ₄	2	4	2.27	2.11(2) 2.43(2)	МО	This study
CuCl ₂ (H ₂ O) ₄	2	4	2.22	2.01(2) 2.28(2)	EXAFS	Tajiri and Wakita, 1986
CuCl ₂ (H ₂ O) ₄	2.8	4.3	2.56	2.05(2) 2.5(2)	ND	Neilson, 1982
CuCl ₃ H ₂ O ⁻	3	1	2.22(1) 2.28(2)	2.28	МО	This study
$CuCl_3H_2O^-$	2.1	1	2.28	1.95	EXAFS	Collings et al., 2000
CuCl ₃ (H ₂ O) ₃ ⁻	3.3	2.7	2.43	1.90-1.95	XRD	Bell et al., 1973
CuCl ₄ ^{2–}	3.8	0	2.24	_	EXAFS	Tanimizu et al., 2007
$CuCl_4(H_2O)_2^{2-}$	3.8	1.9	2.23	2.18	EXAFS	Tanimizu et al., 2007
$CuCl_4(H_2O)_2^{2-}$	4.2	2.3	2.56	2.05	ND	Neilson, 1982
$CuCl_4(H_2O)_2^{2-}$	3.6	2.4	2.43	1.90-1.95	XRD	Bell et al., 1973

 ^a Number of bonds is shown in parentheses.
 ^b An additional Cu-Cl bond distance of 2.85 Å reported may be too long to treat as chemical 1078 bonding.

1096	Table 7 Logarithm of the reduced partition function, $\ln \beta$, for the pair ⁶⁵ Cu- ⁶³ Cu.
1097	Cu(II) hydroxides, carbonates, sulfate, and sulfides.

Chasica			Tempera	ature (K)		
Species	273	298	323	373	473	573
$CuOH(H_2O)_4^+$	5.307	4.517	3.889	2.967	1.883	1.298
$Cu(OH)_2(H_2O)_3$	5.814	4.966	4.288	3.286	2.098	1.451
$CuCO_3(H_2O)_2$	5.091	4.323	3.715	2.825	1.787	1.230
$Cu(CO_3)_2^{2-}$	6.176	5.239	4.498	3.416	2.158	1.483
CuHCO ₃ (OH) ₂ ⁻	5.951	5.075	4.376	3.346	2.130	1.471
$CuSO_4(H_2O)_4$	6.041	5.144	4.430	3.381	2.148	1.481
$CuHS(H_2O)_4^+$	4.002	3.386	2.900	2.194	1.377	0.942
$Cu(HS)_2(H_2O)_3$	3.855	3.264	2.797	2.119	1.333	0.914

- 1099

1115 Table 8 Logarithm of the reduced partition function, $\ln \beta$, for the pair ⁶⁵Cu-⁶³Cu. 1116 Hydrated Cu(I) ion, Cu(II) ascorbates, and Cu(II) oxalate.

Success	Temperature (K)						
Species -	273	298	323	373	473	573	
$Cu(H_2O)_2^+, Cu(I)$	3.368	2.867	2.468	1.882	1.193	0.822	
$CuH(L$ -ascorbate) $(H_2O)_4^+$	3.924	3.324	2.850	2.161	1.362	0.935	
$CuH(D$ -ascorbate) $(H_2O)_4^+$	3.989	3.380	2.899	2.199	1.386	0.951	
$CuC_2O_4(H_2O)_2$	6.236	5.302	4.561	3.474	2.202	1.516	

1122 Figure captions

Figure 1 Mole fractions of Cu species and isotope fractionation of Cu in HCl 1123 1124 solutions. a) Distribution of Cu chlorides. The mole fractions were calculated from the stability constants of Brugger (2001). b) Isotope fractionation is shown as $\Delta^{65}Cu =$ 1125 δ^{65} Cu_{org} – δ^{65} Cu_{aq} upon extraction of Cu by dicyclohexano-18-crown-6 (DC18C6) from 1126 1127 the aqueous solution. The solid curves represent our estimates from the ln β values (298) K) of fivefold-coordinated $Cu(H_2O)_5^{2+}$, $CuCl(H_2O)_4^{2+}$, and $CuCl_2(H_2O)_3$, and 1128 CuCl₃H₂O⁻ (Table 5), ln β_{CuLCl_2} (3.35‰), and *D*. The dotted curves represent the effect 1129 of $\pm 0.05\%$ errors on ln β_{CuLCl_2} . 1130

- 1131 Figure 2. Molecular structures of hydrated Cu⁺, Cu²⁺, and aqueous Cu(II) species.
- 1132 The structures are drawn using GaussView5 (Gaussian Inc.) (Dannington et al., 2009).
- 1133 Symbol keys: Cu (vermilion), Cl (green), S (yellow), O (red), C (gray), and H (white).

1134 **Figure 3. Temperature dependence of ln \beta.** The ln β values of hydrated Cu²⁺ and 1135 Cu(II) chlorides (see Table 5) are shown as linear functions of T^2 .

Figure 4. Mole fractions of Cu(II) species and Cu isotopic variations as functions of 1136 **pH at 298 K.** a) Mole fractions of Cu species, b) Species δ^{65} Cu relative to the bulk 1137 1138 solution. Literature values of formation constants at ionic strength I = 0.70 M (Powell et al., 2007) were used for the calculations. Σ [Cu(II)] was set to 10⁻⁹ M and it was 1139 assumed that the system was in equilibrium with air having a CO_2 fugacity of $10^{-3.5}$ bar 1140 (1 bar = 10^5 Pa) (Powell et al., 2007). We further assumed log $K[CO_2(g) = CO_2(aq)] =$ 1141 -1.5 (Morel and Hering, 1993) and concentrations of Cl⁻ = 0.55 mol kg⁻¹ and SO₄²⁻ = 1142 $0.029 \text{ mol kg}^{-1}$ (Macleod et al., 1994). 1143

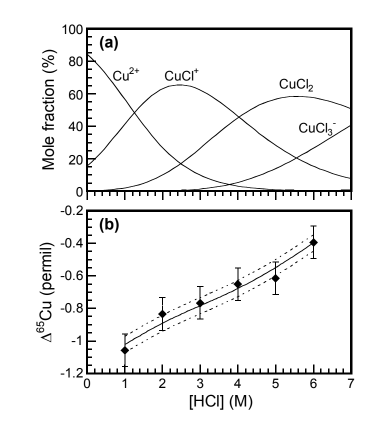
1144 **Figure 5. Temperature dependence of ln \beta.** The ln β values of hydrated Cu(II) 1145 hydroxides, carbonates, sulfate, and sulfides (see Table 7) are shown as linear functions 1146 of T^2 .

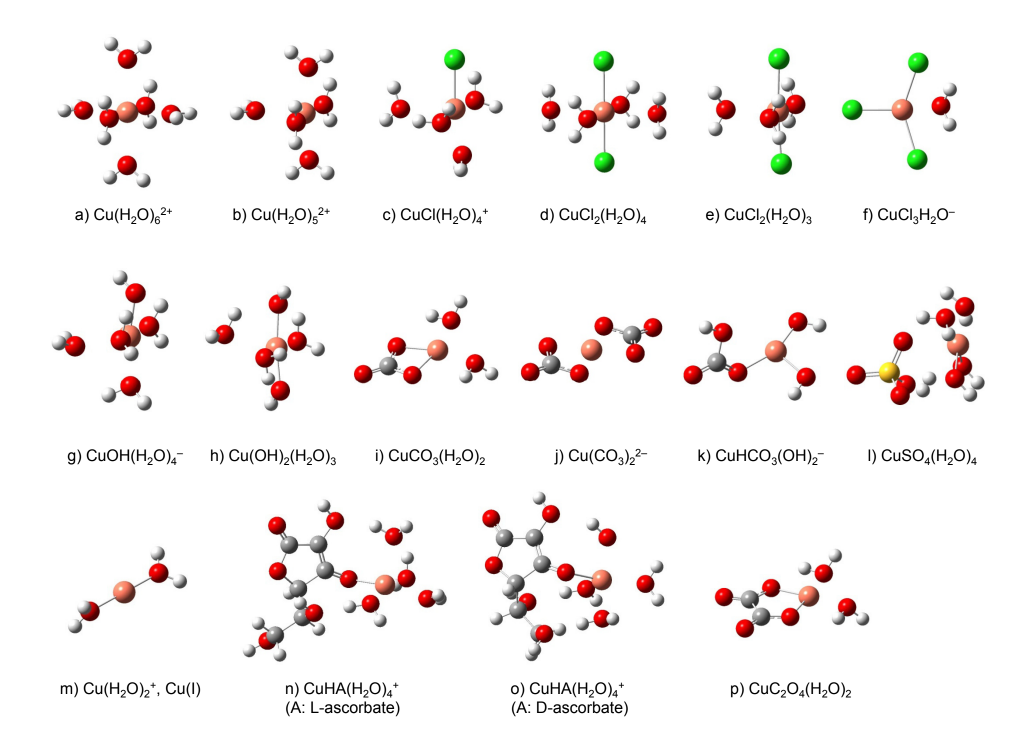
1147 Figure 6. Mole fractions of Cu(I) and Cu(II) species and Cu isotopic variations as 1148 functions of Eh and extent of oxalate formation at 298 K. pH and total Cu 1149 concentration were set to be 7.4 and 1 μ M, respectively a) Mole fractions of Cu species 1150 as functions of Eh. Concentrations of ascorbate and oxalate were set to 90 μ M and 10 1151 µM, respectively. b) Mole fractions of Cu species as functions of proportion of oxalate formed. Eh was set to be 0.27 V. c) δ^{65} Cu relative to the bulk solution as functions of 1152 Eh. Conditions are identical to those of a, and d) δ^{65} Cu relative to the bulk solution as 1153 1154 functions of extent of oxalate formation. Conditions are identical to those of b. **Figure 7. Temperature dependence of ln \beta.** The ln β values of hydrated Cu⁺ and 1155

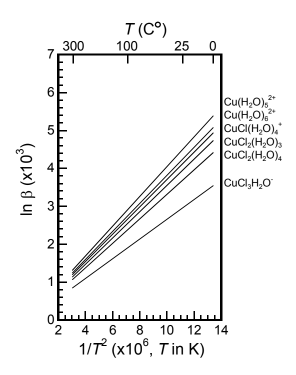
1155 Figure 77 reinperturice dependence of in p. The in p values of hydrated ed and 1156 Cu(II) oxalate and ascorbate (see Table 8) are shown as linear functions of T^2 . The ln β 1157 values of pentaaqua Cu²⁺ (see Table 5 and Fig. 3) are shown together.

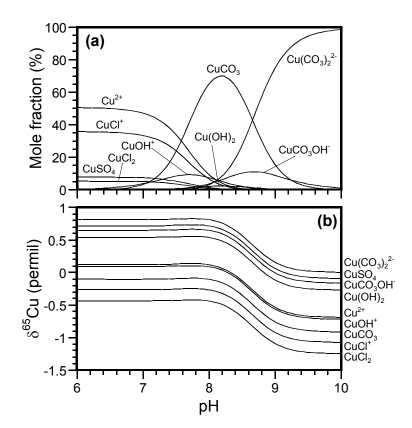
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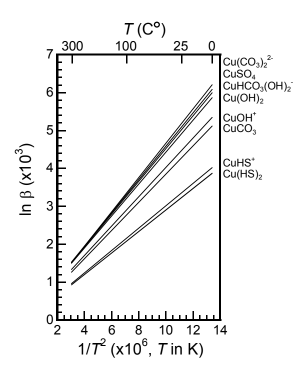
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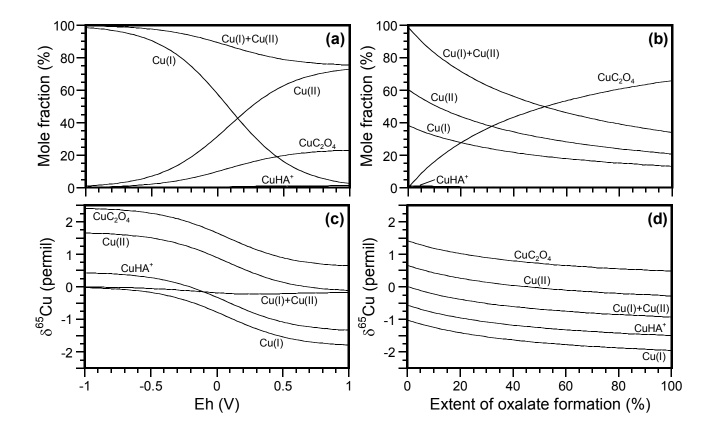


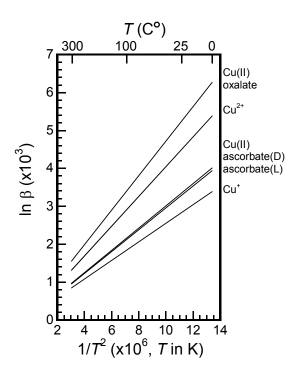












Supporting Information

Copper isotope fractionation between aqueous compounds

relevant to low temperature geochemistry and biology

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Table S1. Optimized structure Cartesian coordinates of hydrated Cu(I) and Cu(II) ions and Cu(II) complexes. (see Figure 2).

0	TT	\mathbf{O}	2+
Cu	H_{2}	$_{2}O)$	5

$Cu(\Pi_2O)_5$			
Element	Х	Y	Ζ
Cu	0.000000	0.000000	0.180286
0	-0.000218	1.972347	0.416359
0	-2.015695	0.000003	0.367094
0	0.000218	-1.972347	0.416359
Ο	2.015695	-0.000003	0.367094
0	0.000000	0.000000	-2.017628
Н	-0.788497	2.520100	0.544468
Н	0.787668	2.520277	0.546113
Н	-0.787668	-2.520277	0.546113
Н	0.788497	-2.520100	0.544468
Н	-2.541191	0.000171	1.181710
Н	-2.620652	-0.000935	-0.390193
Н	2.541191	-0.000171	1.181710
Н	2.620652	0.000935	-0.390193
Н	-0.000087	0.774145	-2.598417
Н	0.000087	-0.774145	-2.598417

$Cu(H_2O)_6^{2+}$					
Element	Х	Y	Ζ		
Cu	0.000038	-0.000034	0.000063		
0	-0.088873	2.019575	0.031113		
0	-0.024969	-0.086591	2.300836		
0	2.025399	0.080251	0.024068		
0	0.088810	-2.019630	-0.031022		
0	0.024925	0.086561	-2.300618		
0	-2.025376	-0.080204	-0.023865		
Н	0.668784	2.613652	0.123299		
Н	-0.902057	2.538568	0.101414		
Н	0.902001	-2.538615	-0.101323		
Н	-0.668838	-2.613716	-0.123240		
Н	-0.067738	0.644710	2.931884		
Н	0.005459	-0.892336	2.834501		
Н	-0.005323	0.892306	-2.834292		
Н	0.067755	-0.644755	-2.931647		
Н	2.576945	0.077097	0.819263		
Н	2.591990	0.154602	-0.756953		
Н	-2.591929	-0.154550	0.757185		
Н	-2.576976	-0.076949	-0.819023		

 $CuCl(H_2O)_4^{2+}$

	/)4		
Element	Х	Y	Ζ
Cu	-0.016654	0.212129	-0.176174
Cl	-0.368293	-1.902143	-0.655033
0	-0.263347	-0.047793	1.850095
0	2.209959	-0.237365	0.440996
0	-0.036021	2.218349	0.250375
0	-0.057535	0.711653	-2.157294
Н	0.598254	-0.331813	2.193790
Н	-0.889110	-0.757015	2.061119
Н	0.438028	2.925105	-0.204498
Н	-0.248494	2.513422	1.145250
Н	3.038932	0.256513	0.412129
Н	2.414530	-1.133283	0.138808
Н	-0.785481	1.250808	-2.496952
Н	-0.059913	-0.113878	-2.668013

$CuCl_2(H_2O)_3$

Element	Х	Y	Ζ
Cu	-0.053962	0.000888	-0.169852
Cl	0.052761	2.249760	-0.117204
Cl	0.050030	-2.247826	-0.115168
0	-0.166668	0.001703	1.905225
0	0.472730	-0.000037	-2.167140

Ο	-2.337733	0.004542	-0.702549
Н	0.114046	0.787034	-2.597917
Η	0.111330	-0.785773	-2.598069
Н	0.267547	-0.788948	2.253624
Η	0.269249	0.791416	2.253590
Η	-2.746356	0.783330	-0.306552
Η	-2.752723	-0.768865	-0.302717

CuCl₂(H₂O)₄

Element	X	Y	Ζ
Cu	0.000113	-0.000205	0.000036
Cl	0.001187	-2.268694	0.000102
Cl	-0.000962	2.268241	-0.000026
0	0.417498	0.000178	2.071588
0	2.327682	0.000039	-0.692256
0	-0.417251	-0.000257	-2.071517
0	-2.327453	-0.001831	0.692327
Н	-2.708483	-0.776704	0.262520
Н	-2.710171	0.772049	0.262246
Н	0.031352	0.780962	-2.422591
Н	0.032951	-0.780315	-2.423107
Н	2.709142	-0.774664	-0.262522
Н	2.709964	0.774089	-0.262095
Н	-0.031968	0.780871	2.422726
Н	-0.031827	-0.780407	2.423132

$CuCl_3H_2O^-$

Element	Х	Y	Ζ
Cu	0.009270	0.089718	-0.002037
Cl	2.197046	0.600440	0.407279
Cl	-0.053553	-0.775059	-2.050390
Cl	-2.075181	0.930696	0.398702
0	-0.031900	-0.399468	2.220232
Н	-0.770277	0.213170	2.362199
Н	0.792853	0.090825	2.362505

$CuOH(H_2O)_4^+$

	71		
Element	Х	Y	Ζ
Cu	-0.022412	-0.011572	0.135042
0	2.055857	0.070828	-0.052489
0	-2.050422	0.169765	-0.181485
0	0.159947	1.789601	-0.261094
0	-0.205225	-2.061829	0.224458
0	0.053073	-0.361275	2.421138
Н	2.051118	1.010485	-0.319345
Н	2.660380	-0.414596	-0.627116
Н	-2.204931	0.986925	-0.676734

Н	-2.650954	-0.512055	-0.507452
Н	-0.097467	2.412135	0.429233
Η	-0.047966	-2.399730	1.116909
Η	0.091646	-2.725054	-0.410229
Η	0.872675	-0.124189	2.873930
Н	-0.665321	-0.109440	3.015233

$Cu(OH)_2(H_2O)_3$

Element	X	Y	Ζ
Cu	-0.258245	-0.086613	-0.062472
0	-0.074679	-0.232907	1.820662
0	2.081764	0.067472	0.259624
0	-0.399476	2.106958	-0.460200
0	-0.677539	0.056454	-1.875154
0	-0.551081	-2.201797	0.273191
Н	-0.175399	-0.497968	-2.477903
Н	-1.413113	-2.553797	0.027322
Н	-0.566410	-1.987168	1.228999
Н	2.369866	0.976539	0.124900
Н	1.738922	0.025664	1.174903
Н	-1.195357	2.594308	-0.223017
Н	-0.543869	1.724067	-1.363498
Н	-0.468696	0.448284	2.370949

$CuCO_3(H_2O)_2$

<u>eace3(11</u>	20)2		
Element	Х	Y	Ζ
Cu	0.248982	0.000870	-0.001777
0	1.470106	1.668045	-0.237640
0	1.470728	-1.665996	0.239991
0	-1.340955	1.081270	-0.128922
0	-1.338478	-1.082522	0.125223
0	-3.325503	-0.003074	-0.002008
С	-2.117328	-0.001419	-0.001917
Н	2.092931	1.971316	0.432853
Н	0.791443	2.353819	-0.329161
Н	0.791117	-2.351065	0.329900
Н	2.096954	-1.971245	-0.426543

$Cu(CO_3)_2^{2-}$

Cu (CC3)2			
Element	Х	Y	Ζ
Cu	0.000000	0.000000	0.000000
0	1.670093	1.094791	0.000000
0	1.670093	-1.094791	0.000000
0	-1.670093	1.094791	0.000000
0	-1.670093	-1.094791	0.000000
0	3.658669	0.000000	0.000000
0	-3.658669	0.000000	0.000000

С	2.413378	0.000000	0.000000	
С	-2.413378	0.000000	0.000000	

CuHCO₃(OH)₂⁻

Element	X	Y	Ζ
Cu	0.328828	-0.274402	0.333280
0	-0.217938	0.652837	1.889544
0	2.152371	-0.434121	0.505155
0	-1.358315	-0.720389	-0.512057
0	-2.850925	-0.477652	-2.183668
0	-0.850477	0.562886	-2.262179
С	-1.778817	-0.270627	-1.625885
Н	2.446380	-0.021985	1.321861
Н	-1.181615	0.642755	1.857756
Н	-1.280353	0.840660	-3.079424

CuSO₄(H₂O)₄

Element	X	Y	Ζ
Cu	0.747523	-0.018631	0.075914
S	-2.464712	0.089386	0.319198
0	0.888308	1.472448	-1.374296
0	1.260468	-1.547098	-1.216183
0	-1.460921	0.074115	-0.784172
0	-3.860376	0.146652	-0.015297
0	-2.105782	1.314366	1.248990
0	-2.198978	-1.199463	1.194935
0	0.426005	1.463763	1.245656
0	0.334120	-1.521773	1.204522
Н	0.822684	1.412610	2.120663
Н	0.727408	-1.537786	2.082364
Н	-1.070215	1.435319	1.324962
Н	-1.181852	-1.395474	1.269655
Н	-0.031570	1.461652	-1.687718
Н	0.699514	-1.631741	-1.997308
Н	0.934346	2.226004	-0.763219
Н	0.994031	-2.244346	-0.588664

$CuHS(H_2O)_4^+$

Element	X	Y	Ζ
Cu	0.005656	0.054181	0.049629
S	0.046759	2.225860	-0.652513
0	2.066189	0.034639	0.017085
0	-2.046642	0.067176	-0.000969
0	-0.098475	-2.084607	0.119384
0	-0.014729	-0.342819	2.333240
Н	2.342733	0.886375	-0.359080
Н	2.619756	-0.660348	-0.362321

Н	-2.350685	0.888640	-0.417416
Н	-2.561894	-0.668691	-0.356925
Η	0.013829	2.775894	0.581471
Η	-0.038801	-2.444466	1.013933
Η	0.055219	-2.798994	-0.509691
Η	0.752827	-0.111541	2.871191
Н	-0.791745	-0.101300	2.852979

$Cu(HS)_2(H_2O)_3$

	-20)3		
Element	Х	Y	Ζ
Cu	-0.314974	-0.153083	0.141975
S	-0.177791	-0.026204	2.425827
S	-0.977088	-0.352010	-2.086603
Ο	2.022133	0.047647	0.018214
Ο	-0.359897	2.013095	-0.277947
Ο	-0.584294	-2.264904	0.309083
Η	0.260936	-0.547023	-2.586321
Н	-1.025393	-2.457587	-0.533692
Н	-1.180639	-2.509597	1.028124
Н	2.206699	0.965967	-0.211249
Η	2.078845	0.012600	0.986178
Η	-1.007256	2.546627	0.198468
Н	-0.707717	1.864863	-1.179080
Н	-0.366876	1.299105	2.585329

$Cu(H_2O)_2^+$, Cu(I)

000(1120)2	, •••(-)		
Element	Х	Y	Ζ
Cu	0.000000	0.000000	0.000000
0	1.917316	0.000000	0.000000
0	-1.917316	0.000000	0.000000
Н	2.479474	0.000000	0.785072
Н	2.479474	0.000000	-0.785072
Н	-2.479474	0.785072	0.000000
Н	-2.479474	-0.785072	0.000000

CuH(L-ascorbate)(H₂O)₄⁺

Curr(E ub						
Element	Х	Y	Ζ			
Cu	-2.554119	0.204618	-0.182677			
Ο	2.495659	-1.159318	2.512367			
0	2.452051	-0.123655	0.482443			
Ο	0.832979	2.264059	0.187868			
0	2.921441	1.402529	-2.662114			
0	-0.251520	-1.875139	2.006688			
0	-0.902877	-0.653920	-0.701190			
0	-1.802772	1.980765	0.480774			
0	-2.984382	-1.344893	1.462135			
0	-3.638265	-1.258716	-1.373516			

Ο	-4.488981	1.096700	-0.188664
С	0.210170	-0.634730	-0.055714
С	0.533793	-1.180051	1.159370
С	1.917511	-0.858682	1.501999
С	1.476723	0.028661	-0.581649
С	1.324332	1.507831	-0.932285
С	2.611406	2.124521	-1.479680
Η	0.282491	-2.124993	2.781107
Η	1.856949	-0.495139	-1.462939
Η	0.550722	1.582431	-1.699857
Η	2.426476	3.184812	-1.685448
Η	3.412119	2.041566	-0.733982
Η	1.495728	2.278551	0.891165
Η	3.769228	1.694906	-3.011156
Η	-0.822376	2.131725	0.453305
Η	-2.229187	2.799140	0.202579
Η	-5.124762	0.619254	-0.737069
Η	-4.976734	1.533791	0.518984
Н	-2.135960	-1.697050	1.787704
Н	-3.557018	-1.253973	2.232088
Н	-3.692109	-2.023413	-0.782479
Н	-3.169074	-1.562546	-2.161632

CuH(D-ascorbate)(H₂O)₄⁺

Element	X	Y	Z
Cu	1.338456	0.278439	2.323761
О	0.071296	1.001644	1.087992
О	3.072810	0.897735	1.231687
Ο	1.773378	-1.729001	1.305373
Ο	-0.285916	0.071930	3.660925
Ο	2.706954	0.025220	3.885105
Ο	1.836297	1.689085	-1.258622
Ο	0.437176	0.641637	-3.585234
Ο	-1.131435	-0.102936	-2.109935
Ο	-0.309048	-2.292050	-0.476613
0	-2.377214	-0.591814	1.927738
С	-0.098546	0.703108	-0.149174
С	0.661040	1.030157	-1.240427
С	0.029557	0.534435	-2.460046
С	-1.310506	-0.058681	-0.671084
С	-1.489555	-1.515966	-0.211950
С	-1.770799	-1.723141	1.289766
Н	-2.225667	0.499580	-0.459825
Н	-2.314597	-1.919188	-0.807171
Н	-0.261507	-2.495742	-1.418672
Н	-2.380007	-2.618680	1.425075
Н	-0.818875	-1.901148	1.791013

Н	-3.309482	-0.757823	2.100138
Η	2.102750	1.825492	-2.183748
Η	2.843084	1.455503	0.467547
Η	3.464206	0.098983	0.853094
Η	3.521725	0.542562	3.868942
Η	2.519606	-0.218994	4.798761
Η	-1.112200	0.003512	3.130698
Η	-0.438922	0.769088	4.309435
Η	1.151785	-2.028910	0.608694
Η	2.088039	-2.518759	1.759410

 $CuC_2O_4(H_2O)_2$

	-)-		
Element	Х	Y	Ζ
Cu	0.220299	0.001436	-0.001762
Ο	-1.136604	0.668638	1.127124
0	-1.137806	-0.672486	-1.125207
Ο	-3.370391	0.723223	1.234860
Ο	-3.371751	-0.739772	-1.222759
Ο	1.501827	0.874109	1.359112
Ο	1.503177	-0.864008	-1.365652
С	-2.354928	0.396973	0.678715
С	-2.355545	-0.407526	-0.671427
Н	0.833390	1.221592	1.973878
Н	2.155906	0.391795	1.877813
Н	2.155176	-0.378487	-1.884016
Н	0.837250	-1.215488	-1.980679

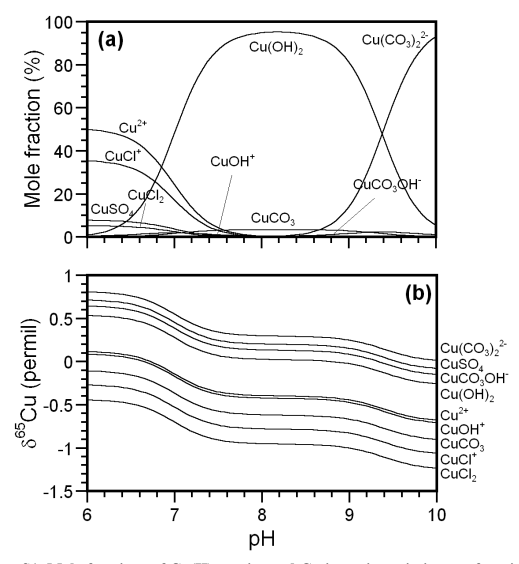


Figure S1. Mole fractions of Cu(II) species and Cu isotopic variations as functions of pH at 298 K. a) Mole fractions of Cu species, b) Species δ^{65} Cu relative to the bulk solution. Conditions are the same with those of Fig. 4 except for second hydrolysis constant. log* $\beta_2 = -16.65$ used in Fig. 4 was substituted by log $\beta_2 = 14.3$ and ion product $pK_W = 14$ (ionic strength I = 0).