1	Atomic-scale characterization of the oxidation state of Ti in meteoritic
2	hibonite: Implications for early solar system thermodynamics
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14	ABSTRACT
15	Calcium-aluminum-rich inclusions (CAIs) in chondritic meteorites are composed of refractory
16	minerals thought to be the first solids to have formed in the solar nebula. Among them, hibonite,
17	nominally $CaAl_{12}O_{19}$, holds particular interest because it can incorporate significant amounts of Ti into
18	its crystal structure in both Ti ³⁺ and Ti ⁴⁺ oxidation states. The relative amounts of these cations that are
19	incorporated reflect the redox conditions under which the grain formed or last equilibrated and their
20	measurement can provide insight into the thermodynamic landscape of the early solar nebula. Here
21	we develop a new method for the quantification of Ti oxidation states using electron energy-loss
22	spectroscopy (EELS) in an aberration-corrected scanning transmission electron microscope (STEM) in
23	order to apply it to hibonite.

24 Using a series of Ti-bearing oxides, we find that the onset intensity of the Ti L_{2.3} edge decreases as 25 a function of increasing Ti-oxidation state, which is corroborated by simulated Ti-oxide spectra using 26 first-principles density-functional theory. We test the relationship on a set of synthetic hibonite grains 27 with known Ti⁴⁺/ Σ Ti values and apply the developed method on a hibonite grain from a compact type 28 A inclusion in the Northwest Africa (NWA) 5028 CR2 carbonaceous chondrite. The STEM-EELS data show that the chondritic hibonite grain is zoned with a Ti⁴⁺/ Σ Ti ratio ranging from 0.78±0.04 to 29 30 0.93±0.04 over a scale of 100 nm between the core and edge of the grain, respectively. The Ti substitution sites are characterized by experimental and calculated high-angle annular-dark-field 31 32 (HAADF) images and atomic-level EEL spectrum imaging. Simulated HAADF images reveal that Ti is 33 distributed between the M2 and M4 sites while Mg sits on the M3 site. Quantitative energy-dispersive 34 X-ray spectroscopy shows that this grain is also zoned in Al and Ti. The Mg distribution is not well-35 correlated with that of Ti and Ti⁴⁺/ Σ Ti at the nanoscale.

36 The spatial decoupling of the element composition and Ti-oxidation states suggests a multistage evolution for this hibonite grain. We hypothesize that Ti and Mg were incorporated into the structure 37 during condensation at high temperature through multiple reactions. Transient heating, presumably in 38 the solar nebula, adds complexity to the crystal chemistry and potentially redistributed Ti and Mg. 39 40 Concurrently, the formation of oxygen vacancies as a result of a reducing gas, led to the reduction of Ti⁴⁺ to Ti³⁺. The multiple defect reactions occurring in this single hibonite crystal preclude a simple 41 42 relationship between the $Ti^{4+}/\Sigma Ti$ and the fO_2 of formation. However, moving forward, these 43 measurements are fundamental input for modeling of the thermodynamic conditions under which 44 hibonite formed in the early solar nebula.

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Keywords: hibonite, Ti oxidation state, STEM-EELS, chondrites, CAIs, thermodynamic, early solar
system, atomic scale, DFT calculations.

48	INTRODUCTION
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50	The sun and planets formed out of a rotating disk of gas and dust over four billion years ago. It was
51	hypothesized that the temperature of the inner solar nebula was high enough such that solid materials
52	experienced melting, evaporation, and condensation (Cameron 1962). Thermodynamic models
53	subsequently predicted that as the solar gas cooled, solid materials condensed out of it according to
54	their volatility, with the most refractory phases forming at the highest temperatures (Lord 1965;
55	Larimer 1967; Grossman 1972; Lewis 1974; Yoneda and Grossman 1995; Ebel and Grossman 2000;
56	Lodders 2003; Ebel 2006; Grossman et al. 2008; Scott and Krot 2014; Zega et al. 2021). Evidence for
57	such condensation is provided by the mineral assemblages found in Ca-Al-rich inclusions (CAIs) within
58	primitive chondrites. With radiometric age dates of 4.5673 ± 0.00016 billion years (Amelin et al.
59	2009; Connelly et al. 2012), CAIs mark time zero for our solar system, and our chronological
60	understanding of its evolution is anchored to that time stamp.
61	Common CAI phases include melilite, spinel, perovskite, hibonite, Ca-pyroxene and forsterite
62	usually with lesser amounts of grossite, metal, and corundum (MacPherson 2014). The composition
63	and structure of these CAI phases are key to determining the chemical pathways through which they
64	formed and to quantifying the conditions under which they condensed or were subsequently altered,
65	whether in the solar nebula or on chondrite parent bodies (Beckett et al. 1988; Simon et al. 1991,
66	1997; Keller and Buseck 1994; Connolly and Burnett 2003; Grossman et al. 2008; Chi et al. 2009). Thus,
67	measurement of CAI phases from the bulk scale down to the atomic level allows us to gain insight into
68	the thermodynamic landscape surrounding their origins.
69	Hibonite (nominally $CaAl_{12}O_{19}$) is among the first few phases predicted to form, at temperatures
70	between 1400 and 1900 K, in thermodynamic models that consider a cooling gas of solar composition

- 71 (e.g., Yoneda and Grossman 1995; Ebel and Grossman 2000; Lodders 2003; Zega et al. 2021). Hibonite
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can incorporate a variety of solutes into its crystal structure, including: Mg²⁺, Zn²⁺, Fe²⁺ and Fe³⁺, V³⁺ 72 and V⁴⁺, Ti³⁺ and Ti⁴⁺, Si⁴⁺ and Th⁴⁺ as minor and trace elements (Allen et al. 1978; Beckett et al. 1988; 73 74 Bermanec et al. 1996; Rakotondrazafy et al. 1996; Simon et al. 1997). Titanium is particularly 75 important because it is often found in significant concentrations in meteoritic hibonite (>0.1 cations 76 per formula unit (Brearley and Jones 1998) and it can occur in multiple oxidation states, thereby providing a redox sensor of nebular conditions (Allen et al. 1978; Armstrong et al. 1982; Beckett et al. 77 78 1988; Doyle et al. 2014; Ardit et al. 2016). Quantitative measurement of the Ti oxidation state in 79 hibonite can, therefore, provide fundamental information on the thermodynamic conditions under 80 which it formed or last equilibrated, e.g., composition of the gas, temperature of condensation or 81 thermal processing, and oxygen fugacity (fO_2) (Beckett et al. 1988). 82 Hibonite is part of the magnetoplumbite group of minerals. It has a space group symmetry of 83 P6₃/mmc and is characterized by unit cell parameters a \approx 5.5Å, c \approx 22.0Å (Curien et al. 1956; Bermanec 84 et al. 1996; Nagashima et al. 2010; Holtstam and Hålenius 2020). This structure contains six cation 85 sites and five oxygen sites. Ca occupies a 12-coordinated polyhedron, whereas the aluminum is distributed among five crystallographic sites (named hereafter M1 to M5). M1, M4, and M5 are 86 87 octahedra, M3 is a tetrahedron, M2 is a trigonal bipyramid, and the atoms within these sites form 88 polyhedral layers perpendicular to the c axis of the hexagonal cell. Atomic substitutions occur 89 abundantly in the hibonite structure, and so the detailed crystal chemistry becomes fundamental to 90 inferring precisely the thermodynamic conditions of the nebular gas at its origin. However, the 91 reactions leading to the Ti incorporation, oxidation, and the location of the substitution in the 92 structure are still debated. Ti can be incorporated into the hibonite structure mainly through two pathways. Ti³⁺ can substitute directly for Al³⁺, or Ti⁴⁺ and Mg²⁺ can couple substitute for two Al³⁺ (Allen 93 94 et al. 1978; Bermanec et al. 1996; Nagashima et al. 2010; Doyle et al. 2014; Berry et al. 2017). The 95 oxidation state of Ti can be further affected by the presence of O vacancies and cation defects, both of

96 which are expected to occur in the structure (Beckett et al. 1988; Asaduzzaman et al. 2021). Regarding 97 the location of the Ti substitution within the M sites (Al), Beckett et al. (1988) proposed that Ti³⁺ occupies the M2 site while Ti⁴⁺ occupies the M4 site. More recently, Doyle et al. (2014, 2016) 98 proposed that Ti³⁺ does not occupy the M2 site, but instead occupies the M4 octahedral site, and Ti⁴⁺ 99 100 partitions between the M2 trigonal bipyramidal and the M4 octahedral sites. The Mg, when balancing 101 the charge of the Ti, is expected to occupy the M3 site. Oxygen vacancies can also occur and modify 102 the Mg and Ti partitioning (Beckett et al. 1988). The distribution of Ti within the M2 and M4 sites was 103 recently observed by coupling STEM-HAADF and energy dispersive spectroscopy (EDS) measurements (Han et al. 2021). However, the $Ti^{4+}/\Sigma Ti$ distribution over these sites has not yet been investigated by 104 105 direct measurement. Thus, establishing a clear connection between the cation site occupancy and the 106 oxidation state of Ti in hibonite is crucial to understanding the nature of the substitution reactions, 107 their thermodynamic history, and what they tell us about the thermodynamic landscape of the solar 108 nebula. Here, we examine the oxidation state of Ti and the chemistry of hibonite via EDS and EELS in an 109

110 aberration-corrected scanning transmission electron microscope. We: (1) report an alternative 111 method for the quantification of Ti-oxidation state in Ti-bearing oxides using EELS; (2) examine the 112 $Ti^{4+}/\Sigma Ti$ variations in meteoritic hibonite down to the atomic scale; (3) provide new information on the Ti site substitution; and (4) link the $Ti^{4+}/\Sigma Ti$ variation as well as chemical zonation to the 113 114 thermodynamic history of a hibonite grain from the Northwest Africa (NWA) 5028 CR2 chondrite. We 115 propose that the approach used here is an important steppingstone for the analysis of other Ti-116 bearing oxides in primitive planetary materials and rigorous treatment of the kinds of defect reactions 117 recorded by them.

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SAMPLES AND METHODS

120 EELS has been used for decades to obtain information on oxidation states of 3d metals in a wide 121 range of earth, planetary, and synthetic materials from micro to atomic scales (Leapman et al., 1982; 122 Brydson et al., 1993; Garvie et al., 1994; Van Aken et al., 1998; Garvie and Buseck, 1998; Van Aken and 123 Liebscher, 2002; Zega et al., 2003; Muller et al., 2008; Chi et al., 2009; Shao et al., 2010; Tan et al., 124 2011; 2012). Fitting of linear mixing models on energy-loss near-edge structure (ELNES) is one such 125 approach but requires standards whose valence states, coordination polyhedra, and crystal fields 126 ideally match those of the unknown (e.g., Garvie and Buseck, 1998a; Zega et al., 2003). In comparison, 127 the white-line method, so called because the absorption edges of EELS spectra were first observed on 128 photographic film as white lines (Azaroff and Pease 1974), monitors the chemical shift of EELS spectra 129 to higher energy loss with increasing oxidation state using integration windows (Leapman et al. 1982; 130 Garvie et al. 1994; Garvie and Buseck 1998; Van Aken et al. 1998; Van Aken and Liebscher 2002; 131 Stoyanov et al. 2007). The advantage of the white-line method is that it can accommodate materials 132 with varied crystal chemistry (electrical properties, orientation, site symmetry, etc.) without the need to synthesize endmembers (with systematically varied $Ti^{4+}/\Sigma Ti$ ratios) as is the case with a method 133 134 based on curve fitting. Previous workers developed a calibration curve for Ti based on the white-line approach (Stoyanov et al. 2007). Thus, our purpose here is to modify it as needed to reflect our 135 136 laboratory and electron-optical conditions.

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138 Samples

To reproduce the white-line ratio method, we acquired seven high-purity (>99.8%) standards with known amounts of Ti^{4+} and Ti^{3+} . These materials and their Ti-oxidation states include: TiO (Ti^{2+}), Ti_2O_3 (Ti^{3+}), Ti_3O_5 (Ti^{4+} : Ti^{3+} = 1:2), Ti_4O_7 (Ti^{4+} : Ti^{3+} = 1:1), TiO_2 (Ti^{4+}), $FeTiO_3$ (Ti^{4+}), $CaTiO_3$ (Ti^{4+}). All were acquired from commercial suppliers (CERAC, Alfa Asear, Phelly Materials, and SigmaAldrich) in powder form except Ti_4O_7 , which was kindly provided by Dr. Lindsay Keller (Keller 1995). Standards were crushed in

an agate stone mortar and pestle in dry conditions and deposited onto lacey-carbon films supported
by Cu-mesh TEM grids (Fig. S1a). The dry conditions avoided the use of solvents and hence possible
alteration of the samples but some grain clustering (up to 200 nm) occurred during the sprinkling of
the particles. We measured only those particles that are monodisperse and electron transparent (< 50
nm in thickness, see Fig. S1).
We also examined synthetic hibonites equilibrated under conditions of controlled temperature

and oxygen fugacity (fO_2). We measured the 'ALL' samples reported in Ihinger and Stolper (1986) and Beckett et al. (1988) because their composition and Ti oxidation states are comparable to meteoritic samples. The compositions of these synthetic hibonite samples correspond to the average composition of hibonite in the CG-11 inclusion in the Allende CV3 chondrite. They were chosen because they contain significantly lower concentration of V (which can occur as V⁴⁺ and V³⁺) and a higher Ti concentration than the Blue Angel (BA) inclusion from the Murchison meteorite (0.39 vs 1.1 V₂O₃ wt.% and 7.3 vs 4.1 TiO2 wt%, respectively; see Armstrong et al. (1982).

The Ti⁴⁺/ Σ Ti of these samples was originally measured by electron spin resonance (ESR) spectroscopy (Beckett et al. 1988). We selected the extreme and an intermediate samples of the series that correspond to Ti⁴⁺/ Σ Ti of 0.85±0.01, 0.91±0.01, and 0.95±0.01 for ALL 2-55, ALL2-57, and ALL2-6, respectively. Chips from these experiments were embedded in epoxy bullets and sectioned by with a diamond knife (in a microtome) to obtain a flat surface. Our goal with these measurements was to confirm the accuracy of the calibration using orthogonal approaches of EELS and ESR, and to relate the Ti⁴⁺/ Σ Ti ratio to the fO_2 under which meteoritic hibonite formed or last equilibrated.

To test the efficacy of our calibration and apply it to meteoritic hibonite, we examined a CAI within the CR2 chondrite NWA 5028 (thin section from the Center for Meteorite Studies, Arizona State University collection #1845-5). The CAI was identified as a compact type-A (CTA) (Ramprasad et al. 2018). The hibonite grain is attached to the rim of the CAI (see the RS2 area in Ramprasad et al. 2018)

and identified and quantified using a Cameca SX-100 electron microprobe, located in the Kuiper Materials Imaging and Characterization Facility (KMICF) at the Lunar and Planetary Laboratory, University of Arizona. A 1-µm probe with an acceleration voltage of 15 kV and beam current of 20 nA was used for the analyses, with counting times of 8 to 20 s on the peaks and 4 to 10 s on the background for each element. Mg, Na, Si and Al intensities were acquired on a TAP crystal; Ca, Ti, K, Cr, P and Cl on a PET crystal; and Fe, Mn and V on an LIF crystal. Petrographic description of the CAI is

174 reported in Ramprasad et al. (2018).

We prepared electron-transparent cross sections of the four hibonite samples (3 synthetic and 1 175 176 meteoritic hibonite) using the ThermoScientific (formerly FEI) Helios NanoLab 660 G³ focused-ion-177 beam scanning-electron microscope (FIB-SEM) also located in KMICF. The general procedures of FIB 178 sample preparation, e.g., coarse cutting, lift out, and in situ thinning were previously described (Zega 179 et al., 2007; 2020) but we note some exceptions briefly here. Rather than attach the FIB sections to Cu 180 support grids in flagpole-style geometry, we used the top of one of the posts that has an M-shape 181 configuration (Fig. S1b) to aid in securing the section on both sides and minimize bending. We created 182 electron-transparent windows of 6 μ m x 8 μ m. Other areas of the synthetic hibonite FIB sections were 183 kept thick ~500 nm, i.e., electron opaque. This mounting approach provided additional stability to the 184 FIB section that was necessary in order to obtain very thin (<40 nm) samples for atomic-resolution 185 analyses. All samples were ion polished down to 8 keV acceleration voltage to remove the amorphous 186 damage layer on each surface created by higher-voltage milling (Schaffer et al. 2012).

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188 **STEM**

The Ti standards and the natural and synthetic hibonites were measured using a 200 keV aberration-corrected Hitachi HF5000 scanning TEM (S/TEM). The HF5000 is equipped with a cold fieldemission gun, a third-order spherical-aberration corrector for STEM mode, bright-field (BF), and

annular dark field (ADF) STEM detectors, an Oxford Instruments X-Max N 100 TLE EDS system with dual 100 mm² windowless silicon-drift detectors with solid angle (Ω) of 2.0 sr, and a Gatan Quantum ER (model 965) electron energy-loss spectrometer (EELS). The Quantum ER is a Gatan Imaging Filter (GIF) equipped with a 2048 × 2048 CCD detector and dual EELS capabilities for simultaneous acquisition of low- and core-loss spectra. STEM-HAADF images were acquired using a 200 keV acceleration voltage, a 10 μ A emission current, and 3.45 eV extraction voltage, a STEM condenser aperture of 25 μ m, and a 2 μ s dwell time.

199 Three types of EELS spectrum-imaging experiments were performed. These experiments include: 200 (1) large $(30 \times 30 \text{ nm up to } 150 \times 150 \text{ nm})$ field-of-view measurements to compare the Ti-oxide 201 standards and the hibonite samples (both natural and synthetic) under the same electron-optical 202 conditions; (2) large field-of-view (135 x 135 nm) measurements using a higher probe current and a 203 larger collection angle than in (1); and (3) atomic-resolution maps (field of view: 6.2 nm x 2.5 nm) of 204 the natural hibonite. Each experiment was performed in STEM mode using a similar condition as for 205 the HAADF images. All EELS spectra were corrected for channel-to-channel gain variation and dark 206 current. Each standard and synthetic sample was analyzed two times to verify the reproducibility of 207 results. For the standards, the two measurements represent different grains with different 208 orientations located in different areas on the lacey film of the TEM grid. For the synthetic hibonites, 209 spectra were acquired from different locations in the FIB section.

The experimental conditions for the EELS spectrum imaging are as follows. (1) For large field-ofview analysis, the microscope was operated using a 100 pm probe, a STEM condenser aperture of 25 μ m, a 105 pA probe current, and convergence angle (α) of 28 mrad. Spectra were acquired using an EELS entrance aperture of 2.5 mm and a collection angle (β) of 24 mrad. The energy dispersion was set to 0.025 eV/ch and a drift tube offset of 440 eV, enabling acquisition of the Ti L_{2,3} edge with an energy range of ~ 50 eV and analysis of its ELNES. At this scale, samples are not beam sensitive and the drift is

216 negligible. Therefore, maps were acquired with a single frame but a large pixel time of 2 s for core loss 217 and 0.01 s for the low loss. Quantitative analysis was performed on summed spectra to enhance the signal-to-noise (S/N) ratio. (2) For EELS large field-of-view analysis, we used a different set of 218 219 conditions because we wanted a quantifiable spectrum at each pixel. We used a condenser aperture 220 size of 35 μ m and a probe current of 330 pA. We increased β to 63 mrad by using high-angle EELS 221 mode, which provides a shorter camera length than standard EELS mode. We also used a 5 mm EELS 222 entrance aperture and a dispersion of 0.25 eV/channel to increase the EELS S/N ratio. Spectral images 223 were acquired by averaging multiple passes (5 frames). A pixel time of 0.2 s for the core loss was used 224 and 0.001 s for the low loss. (3) Atomic-resolution EELS requires lower probe current and shorter 225 acquisition time to minimize the beam damage and the carbon deposition, respectively. Thus, we used 226 a condenser aperture of 30 μ m and reduced the probe current to 150 pA. Further, we used the liquid 227 nitrogen cold trap on the HF5000 to limit the carbon deposition during beam rastering and to 228 decrease the sample drift. We used a large EELS spectrometer aperture (5.0 mm), the high-angle EELS 229 mode (β = 63 mrad), and a 0.5 eV/channel dispersion to ensure a high S/N ratio and short acquisition 230 time. To obtain EELS spectra from individual atomic columns on the hibonite grain from the NWA 5028 meteorite, we oriented it along $[11\overline{2}0]$, with a goniometer tilt of α = 16.4° and β = -2.0°, enabling 231 232 analysis of the stacking along [0001]. The drift tube offset was set at 400 eV, and EELS maps were 233 acquired with a single frame, a pixel dwell time of 0.2 s for the core loss and 0.001 s for the low loss. 234 These atomic-resolution experiments limited our acquisition time and probe current, and so EDS 235 spectra could not be obtained at this scale of measurement.

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237 Data Processing

238 Spectra were processed using personal Python scripts (available at: 239 https://github.com/ZanettaPM/Ti-Oxidation-EELS-data-processing) constructed using Hyperspy (de la

240 Peña et al. 2021). The energy position of the low-loss and high-loss spectra from different pixels were 241 calibrated via zero-loss peak (ZLP) centering. The average energy resolution calculated from the full-242 width-half-maximum intensity of the ZLP over all standard spectra (experimental condition 1) is 243 approximately 0.26 eV. The π^* peak of the C K-edge at 285 eV acquired with the same energy 244 dispersion served as a calibration point. A shift of 1.7 eV is commonly observed at the π^* peak energy 245 (after ZLP centering) which corresponds to a \sim 2.5 eV shift at the Ti L_{2.3} edge energy. An inverse power 246 law (AE^{-r}) model, where E is the energy loss and A and r are constants, was used to subtract the 247 background (random plural and single-scattering contributions). An energy window measuring 20 eV, 248 placed between 430 eV and 450 eV (for a 0.025 eV/channel dispersion), was used to fit the 249 background function. This range is located after potential ghost peak channels (due to the previous 250 ZLP calibration) and at lower energy than possible Ti L_{2.3} pre-edge peaks. The plural scattering 251 contributions due to sample thickness were removed by Fourier-ratio deconvolution. We also 252 removed the continuum beneath the white lines to isolate their intensities. There are varied methods 253 for such removal, including a linear fit between edge onset and offset (Garvie and Buseck 1998; Zega 254 et al. 2003), but for the purposes of reproducibility and consistency with previous work (Stoyanov et 255 al. 2007) we applied a double-step function. The onset and the offset refer to the position at which the 256 energy rapidly decreases to a ground-state intensity before the first peak and after the last peak in the 257 Ti L_{2.3} edge respectively. The double-step function was initially implemented by Van Aken and 258 Liebscher (2002) for the Fe $L_{2,3}$ edge and later modified by Stoyanov et al. (2007) for Ti in the form of:

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$$f(E)=h*(arctan((\frac{\pi}{w})*(\Delta E - E1)) + \frac{\pi}{2})) + h*(arctan((\frac{\pi}{w})*(\Delta E - E2)) + \frac{\pi}{2})$$
 (1)

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where ΔE is the electron energy varying over a range of tens of eV; w represents the width of the functions fixed to 2 eV; E1 and E2 are the inflection points fixed at 457.5 and 463.0 eV, respectively;

and h is the height of the individual arctan functions. In this work, h is a free parameter fit to the spectrum after the L_2 edge between 473 and 477 eV depending on the shape of the offset (Giannini 266 2014).

267 EDS data were also processed using Hyperspy. We fit the spectra, i.e., background and peaks, 268 using a linear combination of the different components and a least square minimization method. The 269 model of the background and its implementation are described in Zanetta et al. (2019). Net peak 270 counts are given by the Gaussian areas which allows us to obtain element maps. Quantification was 271 achieved using the Cliff-Lorimer method (Cliff and Lorimer 1975) and an absorption-correction routine 272 developed by Le Guillou et al. (2018) and based on Horita et al. (1987). We used theoretical k-factors 273 provided by Oxford Instruments[®]. Hyperspectral maps, measuring about 0.4 μm² in area, with a pixel 274 size of ~0.5 nm, were acquired with a dwell time of 30 μ s during 1h. These maps were then binned to 275 achieve a suitable S/N ratio, quantified, and recalculated as cation per formula unit (pfu). The first set 276 of maps described below is 160×160 pixels and is used to display the elemental distributions of the 277 cations. The second set of maps measures 10×10 pixels and is used to achieve a high number of 278 counts and plot element variations. Major elements like Al reach 50,000 counts; Ca, Ti and Mg reach 279 around 8,000 counts while minor elements like Fe and Si reach 1000 counts per pixel, per integrated 280 peak. The high number of counts, the accurate fitting of the spectrum, and the absorption-correction 281 procedure, which is particularly important to account for the thickness variation and the effect of 282 heavy elements on lighter elements, allow us to give confident errors on the quantified results (less 283 than 3% on major elements including Mg and Ti).

Experimental HAADF images were compared to simulated images computed using the QSTEM software, a multislice image-simulation code (Koch 2002). Two cases were investigated. We initially simulated a nominal hibonite (CaAl₁₂O₁₉). Then, using the Supercell software (Okhotnikov et al. 2016), we generated a hibonite solid solution of CaAl₁₀Ti₁Mg₁O₁₉ with Ti and Mg substituting on M4 sites and

288 M3 sites, respectively with an occupancy of 0.5. We also investigated CaAl₁₀Ti₁Mg₁O₁₉ with Ti 289 substituting between M2 and M4 sites. However, replicating the low concentration of Ti on the M2 290 site (<1 atom/unit cell for the M2 site, i.e., an occupancy of 0.5) is complex and requires a large 291 supercell. So, for simplicity, we assumed in our QSTEM simulation that the M2 site contains only Al, 292 and as discussed later, this assumption is valid (see Fig. 4b). From this hibonite structure, we simulated 293 HAADF images to compare to our experimental HAADF images. We also added spherical aberration to 294 our simulations of end-members: $CaAl_{12}O_{19}$ and $CaAl_{10}Ti_1Mg_1O_{19}$, accounting for the A1, B1, and C3 295 components of the aberration function, e.g., Urban et al. (1999) and Chang et al. (2006). The model 296 consists of a 60×60×30 Å supercell. The scan window is 80×80 pixels with a resolution of 0.25 Å/pixel. 297 We fixed the number of slices to 21 with a slice thickness of 1.43 Å.

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299 Density Functional Theory

300 To understand the chemical shift and ELNES of our EELS spectra, we used density functional theory 301 (DFT) to simulate spectra. We computed the electronic band structure of the Ti-oxides by means of 302 the linearized augmented plane wave (LAPW) method, as implemented in the Wien2k package (Blaha 303 et al. 2020). Electronic structure calculations of various Ti-bearing compounds were then conducted 304 starting from experimentally reported crystal structures (Table 1). These structures are further relaxed 305 by employing both local spin density approximation (LSDA) and the generalized gradient approach 306 (GGA) (Perdew et al. 1996) exchange-correlation functionals. Both approaches are found to yield 307 similar EELS spectra. Initialization parameters and self-consistent field (SCF) calculation convergence 308 rules were set according to Hébert (2007). The energy to separate core and valence states were fixed 309 at -6 Rydberg (Ry; 1 Ry ≈ 13.605 eV), except for FeTiO₃ and hibonite where values of -7 Ry and -8 Ry, 310 respectively, were adopted because of their atomic configurations. After structural relaxation, we 311 used an atomic sphere radius, i.e. muffin-tin radius (RMT), as large as possible but avoiding any

312 overlap. Within the RMT, the Wien2K code uses spherical harmonics, whereas in the interstitial region, 313 plane waves are instead used. Final RMT values range from 1.63 to 1.99 atomic units (a.u.) for the Ti (Table 1). The matrix size RKmax ($R_{mt} * K_{max}$) was always fixed at 7.0, where K_{max} is the plane wave 314 315 cutoff and R_{mt} is the smallest atomic sphere. Spin-polarized calculations were performed for Ti₂O₃, 316 Ti₃O₅, and Ti₄O₇. Convergence criteria for the SCF cycle were fixed at 0.00001 Ry and 0.5 mRy/au for 317 the energy and the force respectively. After optimization of the structure, we proceeded to the next 318 iteration with higher k-mesh points (sampling points in the irreducible wedge of the first Brillouin zone 319 of the material) until the theoretical electron energy-loss near-edge spectra (TELNES) between two 320 iterations became nearly identical (peak positions and width differing by <0.1 eV). Final k-mesh sizes 321 are presented in Table 1.

322 The TELNES3 program implemented in Wien2k was used to model EELS spectra (Jorissen 2007). 323 Spectra were calculated using the same parameters as our experiments. We used a fixed splitting 324 energy of 5.5 eV between the L_3 and L_2 transitions and a branching ratio (intensity ratio between the L_2 325 and L_3 lines) of 0.8 (Leapman et al. 1982; Stoyanov et al. 2007). We assumed no orientation 326 dependence and so we averaged our calculations over all possible directions of the scattering vector 327 with respect to the crystal. To improve the match between the model and the experimental results, 328 we used 2×2×2 supercells. A full core-hole calculation for every species was achieved by removing one 329 core electron that was later added to the valence electrons. The spectral broadening, which is mostly 330 dependent on instrumental broadening, was always simulated with the same conditions, i.e., no 331 linearly energy dependent valence broadening and spectrometer broadening set at 0.6 eV 332 (experimentally determined). We kept the core-hole lifetime of the two edges as tabulated in the 333 TELNES3 program.

334 We calculated EELS hibonite spectra for two Ti oxidation states. We investigated two cases with 335 $Ti^{4+}/\Sigma Ti$ ratios of 1 and 0.5, respectively, and a composition of CaAl₁₀Ti₁Mg₁O₁₉, and CaAl₉Ti₂Mg₁O₁₉

respectively. We produced the structures using the Supercell program (Okhotnikov et al. 2016). For the $Ti^{4+}/\Sigma Ti$ ratio of 1, we assumed that all the Ti occurs in the M4 site. For the ratio of 0.5, we randomly distributed the Ti between the M2 and M4 site with a 1:1 ratio. Mg is set to occupy the M3 site.

340

RESULTS

341 Chemical shift calibration and white-line ratio

We acquired EELS spectra at the Ti $L_{2,3}$ edge for standards to develop a calibration curve relating 342 the white-line ratio to the $Ti^{4+}/\Sigma Ti$ ratio under the electron-optical conditions for our laboratory. Figure 343 344 1 shows the processed spectra normalized to the individual maxima acquired from Ti-bearing 345 reference materials (the absolute intensity variations are normalized). The spectra show a chemical shift, marked by the edge onset, of 454.0 to 457.5 eV at the Ti $L_{2,3}$ edge that correlates with the 346 347 valence state of the material. We observed doublets (labeled a, b and c, d) at the L₃ and L₂ edges 348 similar to those reported by Stoyanov et al. (2007). The ELNES for these samples increases in complexity from reduced Ti²⁺ in TiO to oxidized Ti⁴⁺ in TiO₂. The spectrum of CaTiO₃, containing Ti⁴⁺, 349 350 shows four well-resolved broad peaks labeled a, b, c and d. In comparison, spectra from samples with 351 mixed oxidation state display secondary peaks convoluted with the major ones. As the Ti-oxidation 352 state decreases, peak intensity a and c decline in comparison to b and d peaks. The spectrum of TiO 353 contains only two major peaks and low-intensity pre-peaks a and c (Table S1). The energy positions of the peaks (Table S2) are in agreement with previous measurements of Ti L_{2,3} spectra of Ti-oxides 354 355 (Leapman et al. 1982; Brydson et al. 1987, 1989; Henderson et al. 2002; Stoyanov et al. 2007).

Hibonite samples (synthetic and natural) that contain a high $Ti^{4+}/\Sigma Ti$ ratio (>0.8) also show four major peaks (a, b, c, and d) with secondary peaks on the b and the c peak. As shown in Fig. 2, the four spectra display ELNES features similar to the Ti-oxide standards. A shift in energy is visible from 458.05 to 458.4 eV at the a peak position (Table S3).

360 Figure 3a shows the $I(L_2)/I(L_3)$ white-line intensity ratio as a function of Ti oxidation state for the 361 standard reference materials and the synthetic hibonites after removal of background and continuum intensity and integration of 1 eV wide windows centered at energies of 455.8 and 465.25 eV (cf., Fig. 1 362 and 2). We obtained an exponential relationship between the $I(L_2)/I(L_3)$ and the $Ti^{4+}/\Sigma Ti$ ratios. Ti-oxide 363 364 standards and synthetic hibonites with high $Ti^{4+}/\Sigma Ti$ values (>0.8) display strong (±50%) I(L₂)/I(L₃) 365 variation (Fig. 3a). To investigate the effect of the second window position on the IL_2/IL_3 ratio, we 366 normalized all spectra to their individual maxima, and we plotted the value of the L_2 integration window as a function of the Ti⁴⁺/ΣTi ratio (Fig. 3b). The data show significant scatter in the integrated 367 intensity of the samples with a high $Ti^{4+}/\Sigma Ti$ ratio. We observe a general positive slope as a function of 368 369 increasing Ti-oxidation state, but the large scatter in the data precludes a strong correlation between the white-line and $Ti^{4+}/\Sigma Ti$ ratios (Fig. 3a). 370

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372 Comparison of experimental and simulated spectra

373 Given the complexities of the Ti L_{2,3} ELNES, we performed first-principles calculations for the 374 reference oxides and hibonites based on previous experimentally determined structures. The goal of 375 these calculations is to verify the electronic-transition energies of the measured materials and to gain 376 insight into the underlying physics. The input structure and parameters for the TELNES 3 calculations 377 are presented in Table 1. To first order, the calculated spectra presented in Figure 4 match well to the 378 experimental data. We observe the same peaks a through d in the simulated spectra (solid lines), 379 giving reasonable 1:1 correspondence with experimental spectra (dashed lines). We observe that the 380 chemical shift, peak position, and peak width are reproduced. What differs between experimental and 381 simulated spectra is the peak heights. Even if the branching (L_3 to L_2) ratio of 0.8 is well reproduced (Leapman et al. 1982), the ratio of the doublets (i.e., a, b and c, d) are not consistent in certain spectra. 382 In the experimental data, the peak a and c (corresponding to the 2t_{2g} level, see §4.1) are always less 383

intense than the peak b and d (3_{eg} level). In comparison, most of the calculated spectra (Ti_2O_3 , Ti_4O_7 , FeTiO₃, TiO₂ and CaTiO₃) exhibit the opposite trend. The most significant difference between the experimental and calculated spectra occurs for CaTiO₃. The calculated CaTiO₃ spectrum shows wider peaks and a significant peak-height variation compared to the experimental spectrum.

388 The hibonite model spectra shows similar results to the standard calculated spectra. The chemical shift is clearly visible in the calculated spectra for the two selected cases where $Ti^{4+}/\Sigma Ti=0.5$ and 389 Ti⁴⁺/ΣTi=1 (Fig. 4b, red and purple curves, respectively). Compared to these calculated spectra, the 390 experimental spectra with Ti⁴⁺/ΣTi~0.9 (ALL 2-6, ALL 2-57 and ALL 2-55 samples) display an edge onset 391 392 energy located between these two endmembers (456.5 – 457.8 eV). Similar to the oxide standards, we 393 find that the experimental ratios between the peaks a and b, and c and d are not reproduced by the 394 calculation (cf., solid and dashed curves in Fig. 4). If calculated and experimental spectra are compared 395 in more detail, the ELNES of the Ti⁴⁺/ Σ Ti=1 case where Ti is only located on the M4 site (purple curve) shows a close match to the experimental spectra. In comparison, the model in which $Ti^{4+}/\Sigma Ti=0.5$ and 396 Ti is equally distributed between Al2 and Al4 sites (red curve) shows the presence of b" and d" peaks 397 398 that are not observed in the experimental spectra.

Regardless of the specific site occupancies and those effects on the ELNES, our TELNES calculations show that the onset position of the Ti $L_{2,3}$ is predominantly dependent on the oxidation state. However, the position of the offset appears to depend on both the chemical shift and other crystalfield effects with peak position varying for similar Ti⁴⁺/ΣTi values (e.g, peak d in FeTiO₃ vs TiO₂), which is corroborated in the experimental data (cf., Figs. 3a,b). These observations required us to develop a modified approach to quantify the Ti⁴⁺/ΣTi ratio.

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406 Edge onset correlation and coupled EELS/EDS NWA 5028 hibonite analysis

407 As the onset position is only dependent on oxidation state for the Ti-bearing materials investigated in this study, we decided to explore its possible use for quantifying the $Ti^{4+}/\Sigma Ti$ ratio. Thus, 408 409 we quantified the onset intensity (all spectra are normalized to their individual maxima) of the L_3 edge 410 using a window of 2 eV ranging from 455 to 457 eV (Fig. 1 and 2. single window highlighted in blue) for 411 the Ti-oxides. The window size and width were selected to give the best calibration (Fig. S3). We find 412 that the L_3 integrated intensity decreases as a function of increasing oxidation state (Fig. 5). We fit the 413 data to a polynomial function, which yields a 0.99 correlation coefficient (Fig 5). The obtained calibration was compared to the quantified $Ti^{4+}/\Sigma Ti$ ratios of previously synthesized hibonites (ALL 414 415 series) and this led to a close match (Fig.5). Based on scattering of the data points from the oxide 416 standards around the fitted curve, we estimate an absolute error of $\sim 1\%$ for the determination of the $Ti^{4+}/\Sigma Ti$ ratios (dashed black curves, Fig. 5). 417

418 A view of the complete CTA CAI and the local area from which the section of NWA 5028 was 419 extracted is shown in Figure 6. We extracted a hibonite grain that occurs at the edge of the inner part 420 of the rim layer (towards the core side of the CAI) with spinel and perovskite and is surrounded by a 421 mixture of melilite and diopside. The hibonite grain contains up to 2 at% of Ti (8.3 wt% TiO₂) and Mg 422 (2.4 wt% MgO), and low V 0.16 at % (0.56 wt% V_2O_3) (Table 2) and appears homogeneous in BSE 423 contrast. Inclusions of spinel and perovskite are also observed in the grain. A large field-of-view STEM-424 HAADF image of the FIB section shows the hibonite grain and surrounding context (Fig. 7a). The 425 absolute thickness of the lower-contrast zone is estimated to be <30 nm according to the method in 426 Lakoubovskii et al. (2008). We oriented the Hib. 1 grain in Fig. 7a along $[11\overline{2}0]$, enabling imaging along 427 [0001] (Fig. 7b). Analysis of an SAED pattern shows the 2.2 nm spacing along the [0001] direction (Fig. 428 7c), which appears in the lattice-fringe contrast of the HRTEM image (Fig. 7b). The SAED pattern 429 reveals sharp and discrete reflections indicating local long-range order, free of stacking defects. 430 However, stacking disorder (not shown) was observed in other locations. We focused our study on the

Hib. 1 hibonite grain in Fig. 7a, but measurements were also acquired from the Hib. 2 hibonite grain for comparison. We applied the new calibration curve to the Hib. 1 grain (Fig. 5). We obtained a mean $Ti^{4+}/\Sigma Ti$ value of 0.86±0.01 using the same acquisition procedure we applied to the standard oxides and synthetic hibonites. This value is in the same range as the orange hibonite from inclusion CG-11 of Allende or hibonite in the Blue Angel inclusion from Murchison (Beckett et al. 1988). This $Ti^{4+}/\Sigma Ti$ value is also comparable to those determined by Simon et al. (2009), Zega et al. (2012), Giannini (2014), and

437 Han et al. (2021) even though their measurements were not based on a standard calibration.

438 In order to evaluate the homogeneity of the composition of the hibonite grain extracted from the 439 CTA CAI of the NWA 5028 CR2 carbonaceous chondrite, we acquired simultaneous EDS and EELS 440 spectra from the same area. We quantified each spectrum contained in every pixel of the EDS maps 441 (Fig. 7a, white-dashed box, labeled Map. Fig. 8). Figure 8 shows the elemental distribution per formula 442 unit (pfu) extracted from the X-ray lines present in the spectra of the map. The Ca map shows some spatial variations, likely due to the thickness of the sample, but the magnitude of variation is low and 443 444 within error. In contrast, Ti and Mg exhibit significant spatial variations (0.65±0.03 to 0.8±0.04 cation 445 pfu) and show inverse spatial correlations with Al. We note that Ti is quantified based on the K α line 446 which is not convoluted with any other line.

447 To further assess the mechanism at the origin of the Ti incorporation and gain insight into cation 448 substitutions, we plotted four pertinent chemical fields based on quantified maps with the highest number of counts (10x10 pixels and thousand counts per peak; Fig. 9). Previous studies indicated that 449 Ti can be incorporated into the structure either by direct substitution of Ti³⁺ for Al³⁺ or Ti⁴⁺ coupled 450 substitution with Mg^{2+} for two Al^{3+} , and therefore we focused on these three elements (Allen et al. 451 1978; Beckett et al. 1988; Doyle et al. 2014; Berry et al. 2017). If cations are normalized to their sum, 452 453 i.e. AI+Ca+Ti+Mg+V+Fe+Cr=13, we obtain values of $AI=10.5\pm0.1$, $Ca=1\pm0.02$ and $O=19\pm0.5$. 454 Ti+Mg+V+Fe+Cr make up the 1.5 pfu difference. Si counts were near the detection limit (see Table 2;

455 Si=0.04 cation pfu and Fig. S4). Ca is homogeneous with a value close to 1 cation pfu, even as Al varies 456 significantly. The Al concentration linearly decreases as Ti+Mg increases; a strong correlation with a 457 slope \sim -1.02 and R² = 0.9 is observed (Fig. 9a). The Ti and Mg concentrations spatially correlate (Fig. 8) 458 but do not match perfectly (and anticorrelate with Al) with a range of 0.65 to 0.80 cations pfu (Fig. 9c). 459 The concentration of Mg is approximately equivalent to Ti on average, but a significant dispersion 460 around an ideal 1:1 correlation between Ti and Mg is observed (Fig. 9c). The Ca site shows a variation 461 of ±0.02. In comparison, Ti and Mg (or Ti+Si and Mg+Fe) show a variation of ±0.15 cation pfu. Fe, Si, Cr 462 and V are usually associated with the same substitution sites and can modify the relationship between 463 Al and the other cations. Since Cr and V are negligible (Fig. S4) we plotted the effect of Fe and Si on the 464 correlation (Fig. 9 d). Their low concentrations did not significantly affect the relationship (M2/M4 465 sites vs M3 site). We note that on average the concentration of Ti is approximately equivalent to Mg 466 and the Ti+Si concentration is approximately equivalent to Fe+Mg as shown by the microprobe bulk 467 measurement (Table 2). However, we emphasize that the nanoscale measurements reveal a 468 significant heterogeneity that is not evidenced by the lower scale microprobe measurements.

EELS spectral images were acquired with EDS maps, and so we evaluated the Ti $L_{2,3}$ edge acquired 469 over the same field of view. We quantified the chemical shift to obtain a $Ti^{4+}/\Sigma Ti$ ratio (Fig. 10) using 470 471 our onset calibration (Fig. 5). The ratio varies from 0.78±0.04 to 0.93±0.04; note that the errors on 472 these values are large due to the use of a smaller energy dispersion for this map (0.25 eV/channel vs 0.025 eV/channel). The map exhibits higher Ti³⁺ relative concentrations towards the grain boundary 473 474 (cf., Fig. 7a). In comparison, the EELS spectrum image (not shown) of the hibonite grain in the top left 475 corner of the FIB section (Fig. 7a) is homogeneous with a mean value of 0.95±0.04. We used the 476 Ti⁴⁺/ΣTi EELS ratio for assessing the electroneutrality of the sample. The ideal charge of 19 O pfu is subtracted from the charge of the cation sum, i.e., Al³⁺+Ti³⁺+Ti⁴⁺+Mg²⁺+V³⁺+Fe²⁺+Cr³⁺+Si⁴⁺. The 477 observed distribution of Ti³⁺, Ti⁴⁺ and Mg²⁺ and the low concentrations of other solutes (Fe, Cr, V, Si) 478

does not yield to electroneutrality, even if errors on quantification and possible variations in oxidation
states for the other solutes are considered (Fig. S7).

To evaluate the crystal chemistry and the atomic-scale variation of the Ti⁴⁺/ΣTi ratio, we performed aberration-corrected STEM-HAADF imaging. Such imaging reveals atomic columns arranged in horizontal planes along the c-axis (Fig. 11). The planes are characterized by a high-contrast central row of atomic columns juxtaposed by slightly less intense atomic columns above and below (Fig. 11b,c). The atomic columns form an echelon-like motif that is repeated vertically along [0001] (Fig. 11b,c).

487 Hibonite is a member of the magnetoplumbite group (Holtstam and Hålenius 2020). Its structure can be described as a sequence of spinel blocks (or S-blocks) interspersed with conductor layers (or R-488 489 blocks), which are visible in Fig. 11d (Nagashima et al. 2010). The Ca, O and Al sites (M1 to M5) are 490 represented with different colors (see the ball-and-stick model). The spinel block consists of the M1, 491 M3, and M5 sites (yellow, orange and green balls) giving a stoichiometry of Y₆O₈ where Y represents a 492 small cation like AI (see Doyle et al., 2014 for graphical representation). The R-Block contains large 493 cations such as Ca along with the M2 and M4 octahedral sites (red, light blue and dark blue), yielding 494 an overall composition of AY_6O_{11} , where A represents a large cation (Ca for instance). In hibonite, the 495 S- and R-blocks alternate, hence the final formula is AY₁₂O₁₉. In the model, the Ca atoms are shown in 496 red and match the central atom of the R-block in the HAADF image with the highest brightness (Fig. 497 11d). In this orientation (and in contrast to Han et al., 2021), Ca is adjacent to the M2 and M4 sites, 498 i.e., the light and dark blue spheres respectively in the model that complete the R-block (Fig. 11c, d, e, 499 f, g). The M1 and M3 sites occur in the darkest areas of the HAADF image, revealing clear separation of 500 each echelon-like motif (Fig. 11 f, g). Finally, the M5 columns (green spheres in the model) surround 501 the R-block and are visible as bright columns. In this orientation [11-20], every other M5 column is 502 brighter due to a higher atomic occupancy. The relatively darker M5 columns adjacent to the bright

503 M5 columns contain lower occupancy (50% lower in this direction). Light elements such as O are 504 generally not detectable with HAADF imaging (see Carter and Williams, 2016) and are therefore not 505 visible on the image simulation (Fig. 11d, f). The simulation accounting for the Ti and Mg substitutions 506 and aberrations in Fig. 11g closely matches the experimental HAADF image (Fig.11c), in particular, 507 reproducing the 'dumbbell' configuration where Ti atoms substitute onto the two adjacent M4 508 columns (white ellipses). The simulation of Mg substituting on the M3 site gives darker areas, 509 reproducing the contrast observed in the HAADF image (Fig. 11c and 11g). The Ti concentration within 510 the M2 site in this local area of the experimental image is below the detection limit.

511 We performed aberration-corrected STEM-EELS mapping of the local region of hibonite at its 512 boundary with perovskite (Fig. 12) to determine its atomic-scale composition (cf., Fig. 7a). Our focus 513 here is on Ti but because the Ca $L_{2,3}$ edge occurs at an energy close to that of Ti (346 eV and 456 eV, 514 respectively), our energy dispersion allows the acquisition of both the Ca $L_{2,3}$ and Ti $L_{2,3}$ edges. We 515 therefore mapped both cations. We note that the intensity of the Al L and K edges tend to be weak 516 and located at lower and higher energy losses, (78 and 1560 eV, respectively) than Ca $L_{2.3}$ and Ti $L_{2.3}$ 517 and are therefore not represented here. The O atoms, which are distributed throughout the structure 518 are generally not detected with HAADF (although they can be observed using annular-bright-field 519 STEM) and can be challenging to resolve by EELS, and so are also not represented here. The STEM-520 EELS mapping shows that the Ca columns are well defined (Fig.12b) and match well to the model with 521 atoms in the center of the R-block (cf., Fig. 11c-g). All columns present relatively similar intensities 522 suggesting that their occupancy is similar over the sampled area. The Ti $L_{2,3}$ map reveals that it occurs 523 in the region of the M2 and the M4 sites adjacent to the Ca (Fig. 12c). The relative concentrations of Ti 524 in M2 and M4 are difficult to determine by EELS, but the high-intensity pixels are distributed around 525 the line defined by the Ca atoms, suggesting higher Ti concentration on the M4 site (Fig. 12c), as 526 shown by the HAADF image simulation (Fig. 11g). We applied our edge onset calibration to the Ti $L_{2,3}$

edge to determine the $Ti^{4+}/\Sigma Ti$ ratio of this local area (Fig. 12d). The data show that the $Ti^{4+}/\Sigma Ti$ ratio is heterogeneous at the atomic scale with potentially more hotspots of higher Ti^{4+} in the center of the R block even if Ti^{3+} hotspots are also observed in the center of R-block in the bottom part of the map. We observe pure Ti^{4+} atomic columns but no pure Ti^{3+} columns.

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DISCUSSION

533 Validity of the single-window method

In EELS, edges result from the transition of core electrons to unoccupied states above the Fermi 534 535 level. The observed peaks in the Ti-oxide standards can mostly be explained by four possible 536 ionizations. The interaction between the Ti 3d, 4s, and 4p atomic orbitals and oxygen 2s and 2p atomic 537 orbitals gives rise to the following molecular orbital levels: $1a_{1g}(\sigma)$, $1t_{1u}(\sigma)$, $1e_g(\sigma)$, $2a_{1g}(\sigma)$, $2t_{1u}(\sigma)$, $2e_{g}(\sigma)$, $1t_{2g}(\pi)$, $3t_{1u}(\pi)$, $t_{2u}(\pi^{0})$, $t_{1g}(\pi^{0})$, $2t_{2g}(\pi^{*})$, $3e_{g}(\sigma^{*})$, $3a_{1g}(\sigma^{*})$, $4t_{1u}(\sigma,\pi^{*})$ (Fischer 1970; Stoyanov et al. 538 2007). In all of the Ti-oxides measured here, the molecular orbitals are filled up to the $t_{1g}(\pi^0)$. The 539 540 dipole selection rules exclude the $4t_{1u}(\sigma, \pi^*)$ as a possible final state, and the $3a_{1g}(\sigma^*)$ level, at higher energy, is a lower probability transition. However, evidence of secondary peaks at 472 and 477 eV was 541 542 observed in the TiO₂ spectrum that could correspond to $2p_{1/2}, 2p_{3/2} \rightarrow 3a_{1g}$ transitions. The number and intensity of the Ti L_{2,3} ELNES peaks are, therefore, mostly controlled by the $2p_{1/2}, 2p_{3/2} \rightarrow$ 543 $2t_{2g}(\pi^*)$, $3e_g(\sigma^*)$ transitions and the degree of occupancy of the $2t_{2g}$ level (Fig.1 and Table S1; Stoyanov 544 545 et al. 2007).

In contrast to the oxide standards measured here, where Ti is restricted to one or two crystallographically distinct octahedral sites, hibonite has three distinct octahedra, M1 (point symmetry D3d), M4 (C3v) and M5 (Cs), a M3 tetrahedron (C3v), and the unusual fivefold-coordinated trigonal bipyramid M2 (D3h). Previous work suggested that Ti mostly occupies the M4 site (Beckett et al. 1988; Doyle et al. 2014, 2016; Ardit et al. 2016; Berry et al. 2017). Ti is therefore located in a site

551 with an octahedral coordination (except for low Ti concentration e.g., Berry et al., 2017) similar to the 552 Ti-oxide standards (Table 1). This similar Ti environment explains why ELNES of the Ti standards with 553 $Ti^{4+}/\Sigma Ti$ ratio=1 (TiO₂, CaTiO₃, and FeTiO₃) and the hibonite spectra show similar peak doublets. 554 The quantized electronic transitions mainly control the observed peak positions, e.g., Fig. 4. 555 However, previous work showed that ELNES are also affected by other interactions. For instance, 556 distortion of the Ti polyhedra, depending on the material, can add complexity to the Ti L_{2.3} ELNES 557 (Stoyanov et al. 2007). The peak width, which depends on the time it takes for the ionized electron to 558 decay to the ground state, can also vary according to the electronic environment (Egerton 1996). 559 Decoupling the portion of the chemical shift due to the oxidation state is therefore challenging. The 560 white-line approach has the advantage of tracking the shift in energy as a mean to quantifying 561 oxidation-state ratios rather than precise fitting of the ELNES which is particularly challenging for early 562 3d metals that contain complex crystal-field effects superimposed on a chemical shift (Fischer 1970; 563 Leapman et al. 1982; De Groot et al. 1990; Brydson et al. 1993; Garvie and Buseck 1998; Stoyanov et 564 al. 2007). The white-line approach has been applied to 3d metals such as Fe (Van Aken et al. 1998; Van 565 Aken and Liebscher 2002) and Ti (Stoyanov et al. 2007). In the latter work, Stoyanov et al. (2007) 566 showed a relationship between the white-line intensity ratio (L_2/L_3) and the oxidation state of Ti. Thus, 567 we attempted a similar approach to obtain a calibration of the shift of the Ti $L_{2,3}$ edge as a function of 568 Ti oxidation state under the electron-optical conditions in our laboratory (Figure 1, 2, 3a). However, 569 because the Ti-oxide standards and synthetic hibonites with high $Ti^{4+}/\Sigma Ti$ values (>0.8) display large 570 (±50%) IL₂/IL₃ variations (Fig. 3a), the fit that we obtained is markedly different from the one reported 571 by Stoyanov et al. (2007). We attribute this difference to the position of the edge offset window (IL₂: 465.25 to 466.25 eV; cf., Fig. 1 and Fig. 3b). For standards with similar $Ti^{4+}/\Sigma Ti$ values (>0.8), we note 572 573 that the L_2 integration window (465.25 to 466.25 eV) can occur on the left side of the d peak, as can be 574 seen for FeTiO₃ and TiO₂ standards or directly on the crest as in CaTiO₃ (Fig.1). Thus, this difference in

575 position will significantly impact the final L_2 intensities and hence the IL_2/IL_3 white-line ratio. 576 Moreover, our experimental results and DOS calculations demonstrate that the dependency of the edge-offset position on other effects, e.g., electronic structure, does not relate to our experimental 577 578 microscope conditions (Fig. 4). These observations suggest that dependency of the white-line ratio on 579 both oxidation and crystal-field effects complicates the development of a universal curve for Ti. 580 Difficulty in reproducing the white-line ratio method for Ti was also noted by Giannini (2014). As 581 an alternative method, Giannini (2014) proposed using relative heights of the four peaks that occur in 582 hibonite spectra as a possible proxy for the oxidation state. To tackle the crystal-field dependency, we 583 first tried various methods that could relate the features present in the spectra to the oxidation state. Spectral parameters that are related to $Ti^{4+}/\Sigma Ti$, but not to the site occupancy, distortion, or Ti content 584 585 would be ideal for a Ti-valence calibration curve. We therefore tested a variety of potential tracers for 586 the valence state including: the position of the d peak (Fig. 2), the doublet dissociation in energy, the 587 edge center energy (defined by the difference between the onset and the offset), the edge variance, 588 and the relative heights (as proposed by Giannini (2014). We concluded however, that all of these 589 parameters depend on effects other than just the chemical shift. The only alternative method leading 590 to a satisfactory calibration curve was obtained when taking the ratio of the peak d energy range 591 (464.5 to 468.5 eV) divided by the L_2 energy range (462.5 to 468.5) (Le Guillou et al. 2015). This 592 method leads to a larger dispersion around the calibration curve but might be considered as an 593 alternative if the S/N ratio of the acquisition is low (since it integrates energy ranges with higher 594 intensities than the single window approach). The IL₃ onset window (455.8 to 456.8 eV) shows a more 595 accurate correlation with the Ti oxidation state. We therefore used the IL₃ onset window for our 596 $Ti^{4+}/\Sigma Ti$ calibration. However, the use of a wider window (455 to 457 eV) increases the final R² (Fig. 5) 597 and reduces artifacts that are background related (Fig. S3). We note that the almost linear relationship between the IL_3 intensity and the Ti⁴⁺/ Σ Ti ratio is linked to the evolution of the width of the band gap 598

599 for the various oxidation states. For TiO, which shows metallic conductivity, the conduction and the 600 valence bands overlap, but for an insulator such as TiO_2 (or hibonite), the band gap becomes larger 601 and the electronic screening poorer, leading to a shift to higher energy relative to metallic Ti. As a test of the efficacy of the alternative method developed here, we find that the $Ti^{4+}/\Sigma Ti$ ratios 602 603 of the ALL sample series quantified with our approach (Fig. 5) are ALL2-6=0.91±0.01, ALL2-604 57=0.88±0.01, and ALL2-55=0.84±0.01. These values compare favorably with the corresponding ratios 605 of 0.95±0.01, 0.91±0.01, and 0.85±0.01, measured by Beckett et al. (1988) using electron spin 606 resonance (ESR) spectroscopy. The agreement between the data sets from these orthogonal 607 approaches and the lower data dispersion around the fitted curve in comparison to the previously 608 discussed white-line ratio method suggests that the alternative method provides a higher accuracy 609 under our optical conditions.

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611 Error estimation on the bulk Ti⁴⁺/ΣTi ratio determination and mapping limitations

612 The single-window method described above based on the edge-onset position requires that each 613 spectrum must first be normalized to its own maximum to remove the intensity variations due to the 614 sample thickness or Ti concentration. This normalization is possible because the d peak always 615 corresponds to the maximum value of the spectra in the studied materials. The use of only one window in front of the edge potentially makes this method sensitive to the background noise 616 especially for samples with high Ti⁴⁺/ Σ Ti ratio (>0.8). Such noise sensitivity does not apply in the two-617 618 window method (e.g. Stoyanov et al., 2007) because at least one window is placed at energies where 619 the signal/background (S/B) ratio is high (S/B>10). In comparison, the edge-onset method described 620 here requires 1) a high S/B ratio (S/B>5) to avoid any noisy spike that would impact the integrated 621 value and 2) correct removal of the background. The background should ideally fall to zero intensity

before the edge. To increase the reproducibility of $Ti^{4+}/\Sigma Ti$ quantification, we fixed the energy window

623 used for the background removal to a constant energy range and position (see the methods section).

For either the two-window method or our approach here, the calibration of the position of the 624 edge is essential. In our case, we fixed the π^* peak of the C, K edge at 285 eV for the standard 625 626 calibration. Keeping the same dispersion and acquiring the C edge of amorphous carbon immediately after the Ti edge maps provided reproducible measurements of the Ti edge with shift variations < 0.04 627 628 eV between all the standard carbon π^* peak positions. This protocol resulted in a small shift of about 2 629 eV of the Ti edge to lower energy among all the standards (Table S2). In comparison, Stoyanov et al. (2007) used a dispersion of 0.3 eV/channel for the carbon calibration step and a dispersion of 0.05 630 631 eV/channel for the Ti edge acquisition. This difference in choice of dispersion is partly responsible for 632 why we obtain a different exponential calibration curve shown in Fig. 3a in comparison to Stoyanov et 633 al. (2007). The data show that standards define an error of ± 1.1 % around the calibration curve for a dispersion of 0.025 eV/channel (Fig. 5). The Ti⁴⁺/ Σ Ti ratio is therefore well constrained for bulk 634 measurements. This error is approximately similar to the 10% error on the Ti³⁺ wt% value determined 635 by Beckett et al. (1988) for values contained between 0.8 to 1 $Ti^{4+}/\Sigma Ti$. For lower $Ti^{4+}/\Sigma Ti$ values, our 636 approach presents a higher accuracy than ESR measurements (if the Ti³⁺ wt% increases relatively to 637 Ti⁴⁺). 638

We find that errors on the Ti⁴⁺/ Σ Ti ratio are larger for nanoscale mapping than for the bulk values (experimental condition 1 in §2.2), which is mostly related to the larger energy dispersion. The higher probe current and collection angle increase the S/N ratio and enable correct removal of the background intensity, which are, as we note above, the main sources of uncertainties in this method. The lower dispersion (0.25 ev/ch), however, gives lower spectral resolution and generates, therefore, an inherently less accurate integration of the onset-position intensity. To determine the accuracy of the Ti⁴⁺/ Σ Ti ratio, we measured the Ti₃O₅ standard (Ti⁴⁺/ Σ Ti ratio=0.5) with two different dispersions,

646 0.025 and 0.25 eV/channel (noted as experimental conditions 1 and 2 in §2.2). We estimate a 647 difference in the $Ti^{4+}/\Sigma Ti$ ratio between these two dispersions of ±0.04, which implies a relative error 648 of ±8% for the Ti_3O_5 standard and a ±5% relative error for $Ti^{4+}/\Sigma Ti=0.86$ in the NWA 5028 hibonite 649 grain.

650 Atomic-scale measurements are more complex than the nanoscale measurements described 651 above because the sample is beam-sensitive and carbon deposition can occur for exposure times 652 >0.2s. These experimental limitations force short acquisition times, which gives a lower S/N ratio than in the nanoscale maps. The background is therefore noisier, leading to larger errors in the $Ti^{4+}/\Sigma Ti$ ratio 653 654 determination. This error is added to the error due to lower energy dispersion (0.5 eV/ch). Even though the pre-edge background can be strongly affected by the noise, the S/N ratio of the Ti $L_{2.3}$ edge 655 is still above the detection limit and the shift is easily resolved (e.g., Fig. S5). The range of $Ti^{4+}/\Sigma Ti$ 656 657 obtained at the atomic level is consistent with bulk values obtained on the same samples. Based on the noise level, we estimate an absolute error in atomic scale maps of ± 0.08 for the Ti⁴⁺/ Σ Ti 658 659 determination, which represents a relative error of ±12%. The bulk measurement allows an accurate guantification of the mean $Ti^{4+}/\Sigma Ti$ value of the grain, whereas maps constrain variations in $Ti^{4+}/\Sigma Ti$ 660 661 down to the atomic scale.

662

663 Heterogeneities at the nanoscale

The microprobe measurement (Table 2) of the hibonite that we extracted from the CAI in NWA 5028 gives a composition (Ca=1.0, Al=10.49 and Ti = 0.70 pfu and Mg ~ Ti) that is within the range of previous measurements for meteoritic hibonites (e.g., Keil and Fuchs 1971; Allen et al. 1978; Armstrong et al. 1982; MacPherson and Grossman 1984; Ihinger and Stolper 1986; Beckett et al. 1988; Kööp et al. 2016), although it is Ti-rich (0.12 to 0.62 Ti pfu: Fig. S6 and Grossman 1975; Paque et al. 1987; Bischoff et al. 1993; Weber and Bischoff 1997; Srinivasan et al. 2000; Lin et al. 2003; Sapah

670 2015). Other solutes such as Fe, Si, Cr or V occur in relatively low concentration (<0.05 pfu), which 671 simplifies the crystal chemistry of the hibonite in this CAI. However, hibonite grains in chondrites can exhibit heterogeneous compositions at the nanometer scale (Han et al. 2015, 2021; Doyle et al. 2016). 672 673 Such variations reflect the spatial and temporal evolution of the thermodynamic conditions, i.e., 674 chemical potential, temperature, and pressure of the system in which grains formed or last 675 equilibrated. Our TEM data do show that the hibonite chemistry is heterogeneous at the nanoscale, 676 which is particularly clear in the elemental maps for Ti, Mg and Al (Fig. 8 and 9). The plotted elemental 677 compositions (Fig. 9) shows that Ca has negligible variation, whereas Al and Ti+Mg are anticorrelated, 678 suggesting that Ti and Mg are only present on Al sites. We emphasize that the bulk sample exhibits 679 approximately equal Mg and Ti concentrations (Table 2), but that compositional heterogeneities occur 680 at the nanoscale for these two elements (Fig. 8 and 9c). The highlighting of such nanoscale 681 heterogeneities could suggest that similar nanoscale variations occur in other hibonite grains.

682 **Crystal chemistry and Ti substitution sites**

The crystal chemistry of hibonite solid solutions, including description of the site occupancies, oxidation states of Mg, Ti, and other solutes is fundamental to the understanding of precise defect reactions that have occurred during the formation and equilibration of the phase. Recent studies showed that Ti mostly occupies the M4 site but that Ti⁴⁺ can also occupy the M2 site (Nagashima et al. 2010; Doyle et al. 2014; Berry et al. 2017; Asaduzzaman et al. 2021; Han et al. 2021). The Ti⁴⁺

distribution between the two sites is dependent on the Ti concentration (Berry et al. 2017).

In our hibonite DFT calculations, we set the Ti to the M4 site or homogeneously distribute it between the M2 site and the M4 site according to the literature (Doyle et al. 2014). Such relatively simple site distributions facilitate investigation of the effect of the Ti occupancy on the ELNES. In our calculations, the equal distribution of the Ti atoms on the M2 and M4 sites in hibonite gives a spectrum containing groups of peaks equal in intensity, e.g., b and b' (or d and d') and b'' (or d'', see

Fig. 4b, red spectrum). These ELNES features are not representative of the experimental spectra likely because of the high concentration of Ti on the M2 site. Rather, a closest match between experimental and simulated spectra, in terms of ELNES features, occurs where Ti occupies only the M4 site (%Ti⁴⁺=1 case; Fig. 4b, purple spectrum). Therefore, the ELNES modelled by our DFT calculations confirm that Ti mostly occupies the M4 site and that Ti is more concentrated on the M4 site rather than the M2 site, consistent with previous observations and computational work (Doyle et al. 2014; Berry et al. 2017;

700 Asaduzzaman et al. 2021; Han et al. 2021).

701 In this study, the use of aberration corrected STEM coupled with image simulation and a 702 hyperspectral data-processing script developed in-house, allowed for the first time a direct 703 measurement of the Ti distribution between the M2 and M4 site in the hibonite structure (Fig. 11 and 704 12). STEM-HAADF images show brighter spots on the M4 site and the weaker intensity of the M3 sites. 705 The image simulations confirm that these features are due to the Ti and Mg substituting in M4 and M3 706 sites, respectively (Fig. 11). We did not observe strong contrast on the M2 position of the HAADF 707 image, suggesting this site contains a low concentration of Ti. Al and Ca sites exhibit homogeneous 708 intensities similar to the simulations (Fig 11 c). Vacancies may occur but are under the HAADF 709 simulation detection limit (2 to 3 at%).

710 EELS spectrum imaging of the Ca L_{2.3} edge corroborates the STEM-HAADF data, i.e., homogeneous 711 intensity of the atomic columns, and suggests that there is no significant cation defect on the Ca site 712 (Fig 12 a, b). In comparison, the EELS spectrum image of the Ti $L_{2,3}$ edge shows high intensity pixels 713 that are dispersed around the Ca site. This dispersion around the Ca rows confirms the HAADF 714 measurements and indicates that Ti mostly occupies the two adjacent face-sharing octahedral M4 715 sites (which appear as dumbbells in projection). It was proposed that this configuration is energetically 716 favored due to stabilization of the Ti-Ti interaction (Doyle et al. 2014; Asaduzzaman et al. 2021). In addition to the Ti substitution site determination, the data show that pure Ti^{4+} columns ($Ti^{4+}/\Sigma Ti=1$) are 717

more abundant in the center of the row where the M2 sites are located (Fig. 12 d). Although the uncertainties on the $Ti^{4+}/\Sigma Ti$ ratio are larger at the atomic level than the nanoscale or bulk, the spatial distribution suggested by our STEM-EELS data indicate that if Ti^{4+} is distributed between the M2 and M4 sites, then Ti^{3+} mainly occupies the M4 site, which is consistent with previous work (Doyle et al. 2014). The observations made at the atomic scale allow us to discuss the mechanism of incorporation which we discuss below.

724

725 Mechanism of incorporation and Ti oxidation state

The observed crystal chemistry and the heterogeneities in conjunction with the knowledge of substitution mechanism(s) of solutes into the measured hibonites can shed light on the thermodynamic conditions that prevailed during condensation and subsequent alteration in the early solar nebula. Ti and Mg are the most concentrated solutes in chondritic hibonites (Brearley and Jones 1998), and the two key substitution reactions hypothesized for their incorporation are:

731
$$Ti(g) + Al_{Al}^X \rightleftharpoons Ti_{Al}^X + Al(g)(2)$$

732 and

733
$$Ti(g) + Mg(g) + 2Al_{Al}^X \rightleftharpoons Ti_{Al}^{\prime} + Mg_{Al}^{\prime} + 2Al(g) (3)$$

Where Al_{Al}^{X} represents an aluminum ion sitting on an aluminum lattice site, with a Al^{3+} charge that 734 define the reference state (Kröger and Vink 1956). Ti_{Al} is a titanium ion sitting on an aluminum lattice 735 site, with a single positive charge (in comparison to the reference charge, i.e., 4+) and Mg'_{41} a 736 737 magnesium ion sitting on an aluminum lattice site, with a single negative charge (2+). In other words, in this equation (2), the cations are in the following valence state: AI^{3+} , Mg^{2+} and Ti^{4+} . Thus, coupling 738 739 the analysis of the spatial distribution of Mg and Ti and its oxidation state is fundamental to testing the hypothesized substitution reactions in hibonite. Previous work showed that most meteoritic 740 741 hibonite grains exhibit a nearly 1: 1 correlation of Mg:Ti (Allen et al. 1978; Armstrong et al. 1982;

742 Doyle et al. 2014; Berry et al. 2017). This well-known 1:1 Mg:Ti correlation led previous workers to conclude that reaction (3) was the main pathway for Ti (and Mg) incorporation (Allen et al. 1978; 743 744 Armstrong et al. 1982; Beckett et al. 1988; Han et al. 2021). The dispersion around the 1:1 line (Fig. 9c) was also observed and attributed to a minor amount of Ti³⁺ being directly incorporated through 745 746 reaction (2) without Mg. Similarly, by plotting Si+Ti vs Fe+Mg, previous authors questioned the 747 incorporation mechanisms of Si and Fe (frequently present in the hibonite crystal chemistry) and their 748 impact on the 1:1 correlation (Simon et al. 2006, 2019; Han et al. 2015, 2021). Part of the dispersion 749 around the 1:1 curve (e.g., see Fig. 11b of Han et al. 2021), not explained by reaction (2) and (3), was therefore interpreted as being due to secondary substitution of Fe²⁺ for Mg²⁺ and Si⁴⁺ for Ti⁴⁺ in parent 750 751 body settings.

752 In our study of this hibonite grain, several lines of evidence cannot be explained by a combination 753 of reactions (2) and (3), or by the incorporation of Fe and Si, and lead us to propose new mechanisms 754 of Ti and Mg incorporation into the hibonite structure. To be clear, we are not suggesting that 755 previously proposed mechanisms are not possible, but rather that they cannot explain the crystal 756 chemistry of the hibonite grain that we measure here. First, some points in Fig. 9c occur above the 1:1 757 correlation line and show that Mg is locally more concentrated than Ti which is unexpected if only 758 reaction (2) and (3) occur. Second, Fe and Si are in low concentrations, and so the magnitude of the 759 dispersion around the curve (Fig. 9d) cannot be explained by the substitution of these cations during a 760 secondary event as previously proposed Simon et al. (2006, 2019) and Han et al. (2015, 2021). Lastly, 761 the Ti oxidation state distribution of the NWA 5028 hibonite grain does not correlate with the Mg and 762 Ti distribution. If, as previously proposed in the literature, reaction (3) is predominant and the direct 763 substitution reaction (2) is responsible for the dispersion around the Ti:Mg 1:1 line, then the Ti 764 oxidation state should correlate with the Ti and Mg distribution. To highlight this discrepancy, we calculated the excess of Mg (i.e., the ratio of Mg²⁺ relative to Ti⁴⁺ cations). Our data show that Mg is in 765

recess in most of the pixels, and that a significant amount of Mg is therefore not associated with Ti⁴⁺

767 (Fig. 13). These data indicate that part of Mg has been introduced independently from Ti into the 768 structure and that a portion of Ti^{4+} was reduced to Ti^{3+} .

Based on the EDS nanoscale quantification (Figs. 8 and 9) and the comparative EDS and EELS data (Fig. 13), we infer that additional reactions must be considered to explain both the Mg concentration and the Ti oxidation state distribution. If Mg substitutes independently in the structure, an oxygen vacancy is locally formed to ensure local charge neutrality.

773
$$Mg(g) + Al_{Al}^{X} \rightleftharpoons Mg'_{Al} + v_{o}^{"} + e' + Al(g)$$
(4)

774
$$O_o = v_o^{"} + 2e' + \frac{1}{2} O_2$$
(5)

(e.g, Kröger and Vink, 1956) where O_o represents an oxygen anion, $v_o^{"}$ a doubly ionized oxygen vacancy, and e' a free electron. If Mg substitutes near a Ti³⁺ already present in the structure, then the Ti is oxidized to Ti⁴⁺:

778
$$Mg(g) + Ti_{Al}^{X} + Al_{Al}^{X} \rightleftharpoons Mg_{Al}' + Ti_{Al}' + Al(g)$$
(6)

Lastly, we showed that Ti^{4+} could also be reduced to Ti^{3+} (Fig. 13) by trapping the free electrons originating from the formation of oxygen vacancies (5), i.e.,

781
$$Ti_{Al}^{\cdot} + e' \rightleftharpoons Ti_{Al}^{X}$$
 (7)

782 In order to test the hypothesis that oxygen vacancies could occur and play a key role in dictating the concentration of Ti³⁺ (and the distribution of Ti and Mg) in the structure, similar to the mechanism 783 originally proposed by Beckett et al. (1988), we compared our $Ti^{4+}/\Sigma Ti$ maps to the total charge of the 784 785 cation sum subtracted by 19 O pfu (see §3.3). As noted, electroneutrality was not obtained, and we find that the charge difference matches our $Ti^{4+}/\Sigma Ti$ maps (Fig. S7). Based on quantification of the 786 electroneutrality from our EDS and EELS data, we estimate that oxygen vacancies represent 0.25 to 0.5 787 pfu of the grain (Fig. S7). An oxygen vacancy of 0.5 pfu represents 1/12 of the atoms on the O4 788 789 Wyckoff site (which is the closest site to the M3 where Mg substitute) and 1 oxygen atom per unit cell

(which contains 38 oxygen atoms). Several additional pieces of evidence support that the charge difference is certainly due to O vacancies including: (1) neither Ca nor Al cation vacancies were observed in the experimental HAADF images, which was confirmed by comparison to the calculated images; (2) no abundant Ca vacancies were observed on the Ca L_{2,3} EELS spectrum image; (3) other solutes (V, Cr, Fe) are in too low concentration to explain the charge difference; and (4) substitutions (2), (3), or (6) are unlikely to explain the Ti⁴⁺/ Σ Ti distribution because the Mg/Ti and Ti⁴⁺/ Σ Ti spatial distribution are not correlated (Fig. 8, 10 and 13).

797 We conclude that multiple reactions are needed to correctly describe the crystal chemistry of this 798 specific hibonite grain including reactions producing oxygen vacancies. Therefore, the Ti oxidation 799 state distribution in this grain is not only controlled by the Mg/Ti ratio, proposed previously for 800 hibonite (Berry et al. 2017). As shown by reaction (6), the sequence of occurrence of these reactions is 801 also important from the geological context and is discussed in detail in the next section. These multiple reactions preclude direct connection of $Ti^{4+}/\Sigma Ti$ to the fO_2 of the system. Instead, we suggest 802 that the Ti (μ_{Ti}) and Mg chemical potential (μ_{Mg}) as well as the fO₂ of the gas play an important role in 803 the final Ti⁴⁺/ Σ Ti ratio recorded by the grain. Under reducing conditions, the fO_2 of the gas might 804 805 prevail over the μ_{Mg} , whereas in oxidizing environment, the μ_{Mg} could prevail over the fO_2 .

806

807 Cosmochemical Implications

NWA 5028 is a CR2 chondrite, and therefore, is not expected to have experienced hightemperature processing (thermal metamorphism with T> 400°C) on its parent body (Harju et al. 2014). In contrast, evidence for parent body aqueous alteration in the form of Fe-silicates within veins in spinel was observed in the chondrite (Ramprasad et al. 2018). However, we hypothesize that such low-temperature processes would not significantly modify the Ti⁴⁺/ Σ Ti gradient or the total Ti/Mg ratio in hibonite over the timescales inferred for parent-body processing. We base this hypothesis on the

low diffusion rates of Ti and Mg in spinel (Brady and Cherniak 2010), which, as described in detail above, is a fundamental unit of the hibonite structure. Fe^{2+} could however exchange with Mg²⁺ and increase the diffusion rate of the two cations but the Fe concentration in this hibonite is low (<0.04 pfu) suggesting that Fe^{2+} did not replace a significant amount of Mg²⁺ (Simon et al. 2006, 2019). Thus, it is unlikely that parent-body processing impacted the parameters of interest in this study, and so we explore nebular processes below.

820 CAIs are classified into different types based on their mineralogy and morphology. Fluffy type-A 821 (FTA) CAIs are melilite-rich objects that contain highly irregular and nodular structures and are 822 believed to have been formed by gas-solid condensation in the early solar nebula. In comparison, CTAs 823 exhibit more spheroidal shapes and are thought to have undergone partial to complete melting and 824 recrystallization sometime after condensation but prior to incorporation into their parent bodies 825 (MacPherson and Grossman 1984; Simon et al. 1999). The CAI selected here is typed as a CTA 826 (Ramprasad et al. 2018). Therefore, at least two stages need to be considered as potential source of 827 the heterogeneity in the grain (Fig. 9c and d). The grain could have incorporated homogeneous 828 amounts of Ti and Mg that were later redistributed in the grain. Alternatively, the heterogeneity could 829 have been directly inherited from the gas phase during initial condensation of the grain. Lastly, both 830 scenarios could have occurred consecutively. We note that this hibonite grain is at the boundary of 831 the Wark-Lovering (WL) rim (Fig. 6) and exhibits a low concentration of Si (Table 2), which suggests 832 that this hibonite did not equilibrate with a silica-rich melt that might have formed from the secondary 833 event that modified the morphology of the CTA. The heterogeneous AI, Ti, and Mg concentrations observed at the nanoscale (Fig. 8,9) as well as a $Ti^{4+}/\Sigma Ti$ gradient directed toward the grain boundary 834 835 (away from the capping layer Fig. 7, 10) indicate that the thermal event that led to the chemical, 836 structural, and morphological modification of the CAI was, locally, either too brief or at too low a 837 temperature to fully re-equilibrate this hibonite grain with surrounding material.

838 The complex history recorded by this hibonite grain is shown schematically in Fig. 14. We consider two scenarios. In scenario 1, the heterogeneity is inherited during nebular equilibration, i.e., before 839 840 the transient thermal event. 1.a) we hypothesize that the hibonite grain is formed through 841 backreaction with previously condensed material, in accordance with thermodynamic predictions, 842 producing a nominal CaAl₁₂O₁₉ composition; 1.b) Ti is incorporated at high temperature through a 843 direct substitution (reaction 2). 1.c) Mg solubility in hibonite increases as the temperature decreases 844 (e.g., Lodders, 2003) and is added through multiple reactions (4 and 6). Equilibrium between the grain 845 and the gas was not achieved and an elemental gradient was recorded, i.e., the grain became zoned in 846 Ti and Mg and Ti⁴⁺/ Σ Ti. 1.d) a transient thermal event occurred, presumably in the solar nebula but 847 prior to accretion of the CAI in its parent body. This secondary event may have remodified the Ti, Mg, and $Ti^{4+}/\Sigma Ti$ distribution. For instance, if the fO_2 was sufficiently low during the thermal processing, 848 formation of new oxygen vacancies (reaction 5) could have led to the reduction of Ti⁴⁺ to Ti³⁺ via 849 850 reaction (7).

In the second scenario, both 2.a) solid-gas interaction or 2.b) a direct condensation of molecular 851 species to form the hibonite grain could have led to a homogeneous composition. Ti and Mg are 852 incorporated simultaneously through a coupled substitution (reaction 3 above) at high temperature 853 into the hibonite structure. Indeed, Doyle et al. (2014) showed that a local Mg²⁺ and Ti⁴⁺ association in 854 the structure (replacing two nearest Al^{3+}) are energetically favored in hibonite rather than the 855 856 formation of isolated defects (i.e., dissociated Ti and Mg atoms). If Ti facilitates the incorporation of 857 Mg at temperature higher than its normal condensation temperature (according to thermodynamic predictions, e.g., Lodders, 2003), then it is likely that the Mg²⁺ and Ti⁴⁺ association (i.e., the coupled 858 859 substitution reaction 3) will dominate. A small amount of direct substitution reaction (2) could have occurred. 2.c) At this stage, the Mg/Ti distribution matches the $Ti^{4+}/\Sigma Ti$ distribution and the 1:1 860 861 correlation still occurs in the grain. 2.d) The heterogeneity is inherited from the stage 2. The transient
thermal event impacting the CTA modifies the distribution of Ti and Mg and oxygen vacancies are

863 formed. The Ti⁴⁺/ΣTi is modified by reaction (7) reducing Ti⁴⁺ in the structure to Ti³⁺. The 1:1 Mg:Ti

864 correlation is lost and the Ti and Mg distributions do not match the Ti⁴⁺/ΣTi distribution.

It is difficult to conclude whether scenario 1 or 2 is the correct interpretation for the history of this grain because thermodynamic models of Mg and Ti in hibonite are not available in the literature. Thermodynamic equilibrium calculations predict hibonite forms as a relatively pure Ca-Al-oxide. Therefore, measurement of hibonite grains extracted from FTA CAIs and the development of a more comprehensive thermodynamic model would be important steppingstones in understanding which scenario occurred for this grain and what such defects can tell us about the conditions in which hibonite formed or last equilibrated.

872

873 Hibonite as nebular oxygen barometer?

874 Hibonite is among the first several phases thermodynamically predicted to form in a gas of solar 875 composition (Yoneda and Grossman 1995; Ebel and Grossman 2000; Lodders 2003). According to such 876 models, corundum (Al₂O₃) which is calculated to condense first, can back react with Ca (as atomic Ca 877 or molecular CaO) in the nebular gas to form hibonite. If the hibonite grain equilibrates with the gas, then the Ti⁴⁺/ Σ Ti in hibonite should reflect the T, P, and fO_2 of the system. It was proposed that under 878 sufficiently oxidizing conditions, the Ti⁴⁺/ Σ Ti in hibonite is linearly correlated to the oxygen fugacity 879 880 (fO_2) of the environment in which it formed (Beckett et al. 1988). This relationship was established assuming that: 1) most of the Ti in hibonite is Ti⁴⁺ and incorporated through the coupled substitution 881 reaction (3) and 2) Ti⁴⁺ was reduced to Ti³⁺ by trapping free electrons originating from the formation of 882 883 oxygen vacancies (reactions 5 and 7). In their study, Beckett et al., (1988) investigated this relationship for specific hibonite compositions, i.e., with homogeneous Mg concentrations and hence a fixed 884 885 chemical Mg potential (μ_{Mg}). In comparison, Doyle et al. (2014) and Berry et al. (2017), argued that the

Ti³⁺ concentration in the hibonite structure is directly dependent on the Mg activity in the gas (reaction 7) and that Ti⁴⁺/ Σ Ti might be independent of the fO_2 . This scenario implies that in the absence of Mg, all Ti in hibonite must be Ti³⁺. We note that in their studies they investigated fO_2 conditions similar to those synthesized by Beckett et al. (1988) but on samples with different Mg concentrations, i.e., variable μ_{Mg} . Thus, moving forward, we underline three uncertainties that must be resolved for hibonite to be used as an oxybarometer.

First, the analysis of hibonite in the CTA from NWA 5028 reveals that Mg played a key role in the Ti oxidation state recorded by the hibonite grain. The final Mg concentration must therefore be considered because the relationship between $Ti^{4+}/\Sigma Ti$ and fO_2 will depend on the availability of Mg according to reaction (4). The synthesis of hibonite matching the compositions of specific samples (BA and ALL samples) allowed Beckett et al. (1988) to determine a relationship for a specific set of conditions and a fixed Mg chemical potential. Thus, understanding the relationship of $Ti^{4+}/\Sigma Ti$ and fO_2 in other hibonite samples, with different compositions, requires knowledge of the μ_{Mg} .

Another complexity in relating $Ti^{4+}/\Sigma Ti$ and the fO_2 is that numerous 3d metals can substitute into 899 900 the hibonite crystal, i.e., Fe, Cr, V (Allen et al. 1978; Beckett et al. 1988; Bermanec et al. 1996; 901 Rakotondrazafy et al. 1996; Simon et al. 1997). Of these, V can occur in relatively high concentration in 902 certain types of CAIs, e.g., the blue angel inclusion from Murchison that contains up to 1.8 wt% V_2O_3 (Armstrong et al. 1982). If, e.g., V^{3+} or V^{4+} compete with Ti for similar AI sites, the measured Ti⁴⁺/ Σ Ti 903 904 ratio alone will not be representative of the fO_2 of the material when it last equilibrated. Other 905 elements can also compete with Ti in meteoritic hibonites. Even if they occur in lower abundance than 906 V, the estimation of their concentration is also important to establishing the relationship between $Ti^{4+}/\Sigma Ti$ and the fO_2 . 907

908 In addition, the fO_2 variation as a function of temperature should be considered. For Ti⁴⁺/ Σ Ti to be 909 useful in constraining redox conditions for natural hibonites, the temperature pathway experienced by

910 the grain would ideally be known. For example, the temperature of formation and hence fO_2 of a 911 hibonite grain extracted from an FTA CAI preserving primary signatures of condensation could be 912 estimated based on equilibrium thermodynamic predictions. However, the case of a CTA CAI, as 913 measured here, is more complex. On the one hand, it is likely that kinetics played an important role on 914 the recorded Ti-oxidation state during the brief and intense thermal processing that the CTA may have experienced, precluding correlation of the $Ti^{4+}/\Sigma Ti$ ratio to the fO_2 without an appropriate kinetic 915 916 model. On the other hand, the elemental and the oxidation gradients could be used to determine the 917 time-temperature path associated with the diffusion process from profile modeling (Cuvillier et al. 918 2015). However, the temperature and the duration of thermal processing of this CTA CAI was not 919 estimated because it requires disentangling if the zonation was inherited from growth during 920 condensation or because of diffusion in a subsequent event. Deciphering these signatures is beyond 921 the scope of this paper.

Developing methods to quantify the nebular fugacity is challenging because the entire crystal chemistry, including all possible crystal defects, must be known. A complete model considering the parameters discussed above, e.g., μ_{Mg} , temperature history, the solutes competing with Ti, and the reaction paths for Ti incorporation would give a correct estimation of the fO_2 of the gas in which the grain formed or last equilibrated. Such a model will be the subject of a future paper.

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928

CONCLUSIONS

We developed an EELS method to measure the $Ti^{4+}/\Sigma Ti$ ratio of oxide materials at various spatial scales. Experimental data, supported by first-principles DFT calculations, confirm that use of the EELS Ti L_{2,3} edge onset is suitable to quantify Ti-oxidation states in heterogeneous materials. By coupling STEM-EDS and -EELS and applying them to measurement of a hibonite grain from a CTA CAI extracted from the NWA 5028 CR2 chondrite, we constrained the hibonite chemistry. The grain is zoned, with AI

and Ti anticorrelating, and Ti occurring as both Ti³⁺ and Ti⁴⁺. Further, applying our developed 934 935 quantitative EELS method at the atomic scale, we find that the Ti $^{3+}$ mainly occupies the M4 site, Ti $^{4+}$ is 936 distributed between the M2 and M4 sites with a preference for M4, and Mg occupies M3. From the 937 crystal chemistry, we infer that multiple reaction paths incorporated both Ti and Mg, and that coupled 938 substitution is not the only reaction incorporating these cations into the hibonite structure. The 939 formation of oxygen vacancies in a reducing nebular gas impacted the concentration of the reduced Ti. 940 We conclude that the observed zoning testifies to a complex history in which hibonite may have been 941 witness to large changes in nebular oxygen fugacity. However, the use of hibonite as an oxybarometer via measurement of its $Ti^{4+}/\Sigma Ti$ ratio alone is made complicated by its crystal chemistry, the solutes 942 that can compete with Ti, and the varied reaction paths for Ti incorporation. We show that the μ_{Ti} , μ_{Me} , 943 and fO_2 of the system, play an important role in the final Ti⁴⁺/ Σ Ti ratio recorded by the grain. 944 945 Depending on the redox conditions of the system, the fO_2 or the μ_{Mg} could prevail. Thus, careful 946 measurement of these variables and consideration of the reaction paths are essential. An approach 947 combining measurements and modelling of the kind we report here could allow for more widespread 948 use of hibonite (and other phases) as direct probes of the thermodynamic landscape of the early solar 949 nebula.

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Standard	Ti ⁴⁺ /ΣTi ratio	Space group	Ti Symmetry	RMT Ti a.u.	RMT O a.u.	RKMax (R _{mt} * K _{max})	Energy threshold (Ry)	Final Kmesh points	Final Kmesh point for supercell	Initial cif file
TiO ₂	1	P4 ₂ /mnm	D _{2h}	1.82	1.65	7	-6	2000	100	(Smyth et al. 1995)
CaTiO ₃	1	Pmna	C _i	1.94	1.76	7	-6	200	10	(Yamanaka et al. 2002)
FeTiO₃	1	R-3C	C _{3i}	1.99	1.77	7	-7	500	10	(Wechsler and Prewitt 1984)
Ti₄O ₇	0.7	P-1	C ₁	1.74	1.57	7	-6	100	6	(Afir et al. 1999)
Ti₃O₅	0.5	C2/m	Cs	1.63	1.48	7	-6	100	5	(Onoda 1998)
Ti ₂ O ₃	0	R3cH	C _{3i}	1.71	1.55	7	-6	100	10	(Newnham and Haan 2014)
TiO	(2+)	Fm-3m	O _h	1.96	1.77	7	-6	1000	100	(Loehman et al. 1969)
Hibonite	0.5 - 1	P6₃/mmc	C_{3v} and D_{3h}	1.74	1.63	7	-8	10	3	(Moore et al. 1989; Doyle et al. 2014)

1217Table 1: DOS parameters for the SCF calculations and optimization of the different species. Space group shown in Hermann-Mauguin1218notation; Ti symmetry shown in Schoenflies notation. RMT: muffin-tin radius, corresponding to the non-overlapping largest-atomic spheres in a1219spherically symmetric potential approximation. Ry; 1 Ry \approx 13.605 eV. Kmesh point = sampling points in the irreducible wedge of the first Brillouin1220zone of the material. See text for more details. Visualization of the structures is presented in Fig. S2

Elements	Wt%	At%	1 <i>o</i>	pfu
Mg	2.44	2.12	0.07	0.68
Si	0.16	0.12	0.02	0.04
AI	41.95	32.79	0.59	10.49
Cr	0.04	0.02	0.01	0.01
Са	5.98	3.14	0.10	1.00
Fe	0.10	0.04	0.05	0.01
Ti	4.99	2.20	0.08	0.70
Κ	0.01	0.00	0.01	0.00
Mn	0.00	0.00	0.00	0.00
V	0.38	0.16	0.06	0.05
Na	0.00	0.01	0.02	0.00
Р	0.01	0.00	0.02	0.00
Cl	0.00	0.00	0.00	0.00
0	45.06	59.40		19.00
Total	101.11	100.00	Sum cations	12.99
1				

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1227 Table 2: Composition of the hibonite grain as measured via EPMA before FIB sampling. Composition given in weight percent (Wt%) and 1228 atomic percent (at%) and in cation per formula unit (pfu) based on 19 O atoms.

1

2 Figure 1: Stack of ELNES spectra of the Ti-bearing oxides. Two sets of data were processed and averaged

- 3 for each sample. Spectra are ordered according to their $Ti^{4+}/\Sigma Ti$ ratio. White line integration windows
- 4 defined by Stoyanov et al, (2007) are indicated in red (i.e. L_3 and L_2 windows). The boundaries of the
- 5 single integration window for the edge onset method presented in this work are indicated in blue. See
- 6 text for discussion of window width and placement. Peak positions are given in Table S2.

Figure 2: ELNES spectra of the synthetic (ALL series) stacked in order of increasing $Ti^{4+}/\Sigma Ti$ ratio and natural (NWA 5028) hibonite. Two sets of data were processed and averaged for each sample. White line integration windows defined by Stoyanov et al. (2007) are indicated in red (i.e. L_3 and L_2 windows). The single integration window for the edge onset method presented in this work is indicated in blue. Peak positions are given in Table S2.

11 positions are given in Table S3.

Figure 3: The white-line ratio shown as a function of Ti oxidation state. a) Application of the white-line ratio method using the window positions and the background subtraction method detailed in Stoyanov et al, (2007). Both Ti-bearing oxides (blue crosses) and synthetic hibonites (red circles) are plotted. The fitted curve formula is given in the top left corner of the graphic. b) The L₂ edge intensity (after normalization; see text for details) plotted as a function of $Ti^{4+}/\Sigma Ti$. Both Ti-bearing oxides (blue crosses) and synthetic hibonites (red circles) are plotted. Large variations are revealed for samples with similar

18 $Ti^{4+}/\Sigma Ti$ close to 1 (i.e., CaTiO₃, TiO₂, FeTiO₃ and synthetic hibonites).

19 Figure 4: ELNES from experimental (dashed curves) measurements and DFT calculations (solid curves). a)

20 TELNES (DOS + Broadening) calculation of the previously listed Ti-bearing oxide standards compared to

21 their experimental spectra. b) TELNES (DOS + Broadening) calculation of two hibonite structures with 22 distinct Ti oxidation states (as indicated, %Ti⁴⁺=0.5-1) compared to the synthetic hibonite experimental

distinct Ti oxidation states (as indicated, %TI⁴⁺=0.5-1) compared to the synthetic hibonite experimental
 spectra. All spectra are normalized to their maxima. In the %TI⁴⁺= 1 case, Ti substitutes on the M4 site; in

24 the $\%Ti^{4+}=0.5$ case, Ti is distributed over the M2 and M4 sites.

Figure 5: The normalized L_3 intensity, single window, as shown in Figs. 1 and 2, as a function of the Ti^{4+} concentration. Synthetic hibonite with known Ti^{4+}/Ti^{3+} falls on the same correlation line. Blue crosses are Ti-bearing oxide standards. Red circles are the synthetic hibonite samples from Beckett et al. (1988). The red line corresponds to the quadratic expression fit to the data (equation shown at top). Dashed black curves represents the error dispersion. This correlation enables determination of the $Ti^{4+}/\Sigma Ti$ ratio (0.86±0.01) of the hibonite grain sampled in NWA 5028 (green lines). The green area represents the error on the determination of the oxidation state.

Figure 6. Microprobe data on a CTA CAI from the NWA 5028 CR2 chondrite. a) Backscattered electron (BSE) image of the CTA CAI showing where the hibonite grain was extracted (orange rectangle). The region of interest is located at the edge of a dark (in BSE contrast) rim that surrounds the CTA (e.g., Ramprasad et al., 2018). b) BSE image of the RS2 area (e.g., Ramprasad et al., 2018) showing the hibonite grain (indicated by orange arrow and outlined with a white dotted curve). c) Higher magnification BSE image of the hibonite grain. The grain exhibits sharp boundaries indicated by a dashed white curve. Surrounding phases are identified. The orange rectangle represents the location of the FIB transect.

Figure 7: Aberration-corrected STEM and diffraction data on hibonite extracted from NWA 5028. a) Low
magnification HAADF - STEM image showing hibonite grains and adjacent perovskite. Some curtaining
appears (relatively dark vertical lines) as a result of ion milling of material with different sputtering rates.
Areas from which diffraction and image data were acquired are indicated. b) Higher magnification
HAADF – STEM image revealing lattice fringes of the oriented hibonite. c) SAED pattern of hibonite from
NWA 5028, viewed down the [1120] zone axis. Measurement of the pattern along the vertical direction
reveals an interplanar spacing of 2.2 nm consistent with the c-axis of hibonite.

- 46 Figure 8: Quantified element maps acquired by TEM-EDS and presented in cation pfu (160×160 pixels).
- 47 This dataset was acquired from the region shown in Figure 7a. Only major elements are plotted.
- 48 Figure 9: Quantification of the STEM-EDS data for hibonite grain 1 (hib.1 see Fig.7a). Composition
- shown per formula unit assuming Al+Ca+Mg+Ti+Si+Fe+V+Cr=13. Each point represents a quantified pixel
 of the rebinned EDS map (10×10 pixels).

Figure 10: STEM-EELS data on hibonite grain 1. a) A map (40×40 pixels) of the Ti⁴⁺/ Σ Ti ratio (hib.1 - see Fig.7a) determined using EELS by the edge-onset method. Ti⁴⁺/ Σ Ti decreases from the interior (higher in the panel) to the grain boundary at the bottom. b) The chemical shift of the Ti L_{2,3} edge (before doublestep function removal) visible between the boundary of the grain (bottom part of the image) and the core.

- 56 Figure 11: Experimental and simulated STEM-HAADF images of hibonite. a) Atomic-resolution STEM-57 HAADF data from hibonite grain boundary (see Figure 7a). b) HAADF image showing the location of 58 atomic-scale measurements (Fig. 12). Atomic columns are visible in this [1120] projection. The map is 59 acquired near the grain boundary. c) High-contrast HAADF image showing the presence of brighter spots 60 on the face sharing M4 sites and weak contrast on the M3 site (white ellipses). d) image simulation 61 (QSTEM[®]) of a nominal hibonite compared to a model in the same orientation $[11\overline{2}0]$. The ball-and-stick 62 model shows 1:1 correspondence with the atomic position visible on the experimental HAADF image and 63 in the simulation. Each site is indicated as a color, e.g., Ca (red), O (black) and the Al sites which are listed 64 from M1 to M5. e) Image simulation with aberrations of a nominal hibonite. f) Image simulation of 65 hibonite containing 0.7 pfu of Ti and Mg substituting on M4 site and M3 site respectively. A model in the 66 same orientation is compared. The white ellipses circle the site of Ti substitution (M4) g) Image simulation of the same hibonite (0.7 pfu of Ti/Mg) with aberrations. The white ellipses show the replica 67 68 of the brighter spot. The space between the blocks are darker than in e) due to the lower Z value as a
- 69 result of Mg substituting into the M3 site.
- 70 Figure 12: Atomic-resolution EELS mapping of the local area shown by the orange box in Fig. 11b in the
- 71 $[11\overline{2}0]$ orientation. (a) HAADF reference image. (b) Ca $L_{2,3}$ atomic-resolution EELS map. (c) Ti $L_{2,3}$ atomic-
- 72 resolution EELS map. (d) $Ti^{4+}/\Sigma Ti$ map. Pixels containing low Ti $L_{2,3}$ intensity were masked to better reveal
- 73 the relationship with the Ti columns and the $Ti^{4+}/\Sigma Ti$ ratio. The map exhibits variation of $Ti^{4+}/\Sigma Ti$ of 0.4
- 74 with an error of ± 0.08 .
- Figure 13: Mg_{Excess} in cation per formula unit (Mg/Ti^{4+}) compared to the $Ti^{4+}/\Sigma Ti$ ratio determined by our single window approach in a hibonite grain of NWA5028. The superimposed orange arrows in the plot correspond to reaction 4 (vertical arrow) and to the reaction 7 (diagonal arrow). The vertical only impacts the Mg concentration, whereas the diagonal line corresponds to Ti^{3+} replacing Ti^{4+} and increasing the isolated Mg defect. The horizontal dashed line corresponds to Mg=Ti or an ideal 1:1 correlation between Mg^{2+} and Ti^{4+} . Error bars are plotted according to the EDS quantification accuracy and the error discussed in section 4.2.
- Figure 14: Schematic of the formation and evolution of the hibonite grain selected near the rim of the CTA CAI. Two pathways 1 and 2 are considered, with each consisting of formation and equilibration stages 1 and 2, respectively in the solar protoplanetary disk. See the text for discussion. Ultimately, the grain did not reach equilibrium and contains an element and Ti-oxidation-state gradient. The direction of the $Ti^{4+}/\Sigma Ti$ gradient is different from the direction of the elemental gradient (orange arrows vs white/blue or white/green arrows in the bottom row).
- 88



















Ti

200 nm





0.68

Figure 8

10.60

- 10.58

- 10.56

- 10.52

- 10.54 P

















Lost of the 1:1 Mg:Ti correlation