PRE-TERRESTRIAL OXIDATION PRODUCTS OF IRON MINERALS IN CARBONACEOUS METEORITES IDENTIFIED IN MÖSSBAUER SPECTRA Duncan S. Fisher and Roger G. Burns, Department of Earth, Atmospheric and Planetary Sciences, Massachusetts Institute of Technology, Cambridge, MA 02139

Introduction. The occurrence of ferric-bearing assemblages, comprising phyllosilicates, oxide hydroxides and magnetite, in carbonaceous chondrites (hereafter abbreviated as CC) indicates that these meteorites underwent pre-terrestrial sub-aqueous oxidation reactions. Although several analytical studies of CC, have been reported [e.g.,1,2], the poor crystallinity of matrix phases makes it very difficult to estimate the modal mineralogy and proportions of Fe3+ and Fe2+ in these meteorites [2]. Early studies of Mössbauer spectra of meteorites at room temperature demonstrated that this technique may provide quantitative information on iron-bearing phases in CC [e.g.,3,4]. In a recent study of ureilites, for example, major proportions of metallic Fe and nanophase goethite were identified in these carbonaceous achondrites [5]. The metallic Fe occurs mostly as sub-microscopic inclusions in forsteritic rims where they have formed by carbon-induced reduction of Fe²⁺ cations in olivine grains in contact with the carbonaceous matrix [6]. The cryptocrystallinity of the metallic Fe inclusions renders the metal extremely vulnerable to oxidation so that nanophase ferric oxides (i.e. "rust") occur in all ureilites, including specimens collected as falls. The vulnerability of olivines in ureilites to such redox reactions suggested that other pre-terrestrial oxidation processes might be elucidated in CC, the olivines in which have undergone reactions producing serpentines, magnetite and poorly crystalline phases such as ferrihydrite and tochilinite [e.g.,2,7,8]. Reported here are results of a Mössbauer spectral study of a suite of CC demonstrating that a variety of ferrous and ferric-bearing phases may be distinguished in different classes of this meteorite-type.

Carbonaceous Chondrite Specimens. Samples of several CC catalogued as falls were obtained from the Harvard Mineralogical Museum Collection, and included: Orgueil (C1); Murchison, Murray, Cold Bokkeveld and Renazzo (C2); Allende (CV3); Warrenton (CO3); Karoonda (C5); and the find Coolidge (C4). Specimens acquired from the Antarctic Meteorite Collection, including some powdered (P) samples, comprised: ALH 83100(P) and MAC 88107 (C2); ALHA 83108 and LEW 85332 (C3); EET 87256(P) and ALHA 85002(P) (C4); EET 87860(P) (C5); and LEW 87009(P) (C6). Experimental details for measuring

Mössbauer spectra of meteorites are described elsewhere [9].

Results. Since room temperature Mössbauer spectra of several <u>CC</u> have been published previously [3,4,10-12] and were duplicated in this study, attention is focussed here mainly on spectra measured at

4.2 K of several <u>CC</u> petrologic types.

(1) The relatively simple 295 K spectrum of Allende ($\underline{\text{CV3}}$) [4,11] consists of Fe²⁺ and Fe³⁺ quadrupole doublets alone representing mainly olivine and phyllosilicates, respectively, and with %Fe³⁺/total Fe = approx. 10%. The 4.2 K spectrum of Allende (Fig.1) is dominated by peaks from magnetically ordered Fe²⁺ ions in olivine. No magnetic ferric phase is present in Allende.

(2) The spectrum of ALH 83100 (C2) at 295 K also consists of Fe²⁺ and Fe³⁺ doublets alone, but with comparable intensities. In the 4.2 K spectrum of ALH 83100, however (Fig.2), the Fe²⁺ doublet of ferroan serpentine remains while magnetic ordering of Fe³⁺ has occurred, indicating the presence of the Fe³⁺ serpentine, cronstedtite [12,13], possessing a magnetic ordering temperature of ≤ 8 K [14]. Tochilinite, which is particularly abundant in these C2 CC [2], does not magnetically order at 4.2K, by analogy with

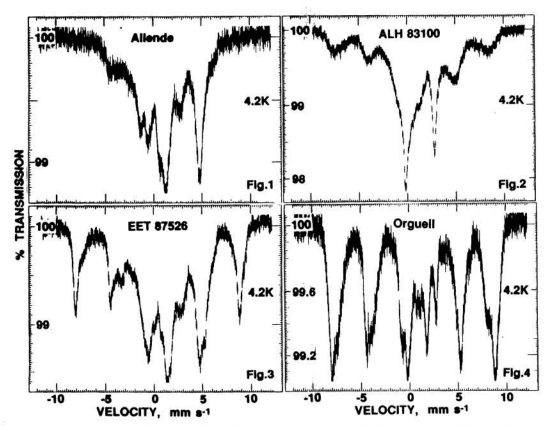
related mackinawite [15], and is recognized by peaks centered near zero mm s-1.

(3) The 295 K spectrum of EET 87526 (C4) contains the Fe²⁺ olivine doublet and a much weaker Fe³⁺ doublet than Allende, as well as the characteristic magnetic hyperfine spectrum of magnetite which constitutes about 20% of the iron modal mineralogy. The relative intensities of paired-peaks in the magnetite spectrum are not in the ratio 1:2, however, indicating that this oxide in EET 87526 and in most CC is an oxidized magnetite departing from Fe²⁺Fe³⁺₂O₄ stoichiometry. The 4.2 K spectrum of EET 87526 (Fig.3) is dominated by magnetite and magnetically-ordered Fe²⁺ in olivine. Similar 295 K and 4.2 K spectra were obtained for other C4, C5 and C6 specimens studied (i.e. Karoonda, ALHA 85002, EETA 87860, LEW 87009). Coolidge (found in 1937) contains no magnetite peaks in the 295K Mössbauer spectrum. Instead, its 4.2K spectrum consists of broadened sextet peaks attributed to terrestrial nanophase goethite.

(4) The spectrum of Orgueil (C1) at 295 K resembles previously published spectra of this CC [3,16], and consists of an intense Fe³⁺ doublet, a very weak Fe²⁺ doublet and the magnetic hyperfine profile of non-stoichiometric magnetite. At 4.2 K (Fig.4), the Fe³⁺ doublet has collapsed to a sextet, the peaks of which flank and broaden the magnetite peaks. This behavior is indicative of ferrihydrite [16], recently identified [8] as the the superparamagnetic phase proposed in Orgueil [17,18]. The spectra of Orgueil are thus consistent with the occurrence of magnetite, ferrihydrite, Fe³⁺ serpentine (cronstedtite) and Fe²⁺ serpentine

in the matrix [8].

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(5) The Mössbauer spectra of other <u>CC</u> not illustrated here revealed that Renazzo has major metallic Fe, minor magnetite, and Fe²⁺ and Fe³⁺ doublets of comparable intensities which do not order magnetically at 4.2 K. Metallic Fe but no magnetite occurs in ALHA 83108 (<u>C3</u>) and LEW 85332 (<u>CO3</u>) together with nanophase goethite, apparently formed by oxidative weathering in Antarctica, by analogy with ordinary chondrites [9]

Ureilite Mössbauer Spectra. Following previous discoveries [5] by Mössbauer spectroscopy of high modal proportions of metallic Fe and the presence of ferric oxides in all ureilites studied, as well as examples of pigeonite Fe²⁺ exceeding olivine Fe²⁺, additional ureilites were acquired for further spectral measurements at 4.2K to confirm these trends.

(1) Major amounts of metallic iron were determined in Goalpara, ALHA 82106, ALHA 82130, and LEW 85440. Minor amounts of Fe occur in MAC 88177 and EETA 83309, in which carbonaceous matrix is either negligible or localized in discrete areas not surrounding olivine grains.

(2) The modal proportion of Fe²⁺ olivine exceeds Fe²⁺ pigeonite in Goalpara, EETA 83225, LEW 85440 and MAC 88177. The Fe²⁺ modal proportions are reversed in ALHA 82106 and ALHA 82130, with pyroxene > olivine. Pyroxene is a very minor constituent of EETA 83309.

(3) Each of these ureilites contains significant to high proportions of ferric oxides, including Goalpara, the freshness of which led to the speculation that it was recovered soon after an unrecorded *fall* [19]. The FeOOH content is least in MAC 88177, the ureilite with negligible carbonaceous matrix.

These observations confirm the mechanism of carbon-induced reduction of Fe²⁺ in olivine [6] and the vulnerability of the sub-microscopic metallic Fe inclusions to rusting, presumably upon exposure to Earth's atmosphere.

References. [1] H.Y.McSween & S.M.Richardson, GCA, 41, 1145 (1977); [2] H.Y.McSween, GCA, 51, 2469 (1987); [3] W.Herr & B.Skerra, in Meteorite Research (P.M.Millman, ed.), p.106 (1968); [4] H.Roy-Poulsen et al., Phys. Scripta, 23, 1113 (1981); [5] S.L.Martinez & R.G.Burns, LPS, XXI, 736 (1989); [6] J.L.Berkley et al., GCA, 40, 1429 (1976); ibid., 44, 1579 (1980); [7] M.E.Zolensky & I.D.R.Mackinnon, Am. Min., 71, 1201 (1986); [8] K.Tomeoka & P.R.Buseck, GCA, 52, 1627 (1988); [9] T.C.Solberg & R.G.Burns, Proc. 19th LPSC, 313 (1989); [10] F.W.Oliver, Planet. Space Sci., 26, 289 (1978); [11] F.W.Oliver et al., Meteoritics, 19, 75 (1984); [12] W.F.Müller et al., Tsch. Min. Pet. Mitt., 26, 293 (1979); [13] T.E.Bunch & S.Chang, GCA, 44, 1543 (1980); [14] J.M.D.Coey et al., Phys. Chem. Min., 7, 141 (1981); [15] D.J.Vaughan & M.S.Ridout, J. Inorg. Nucl. Chem., 33, 741 (1971); [16] E.Murad & U.Schwertmann, Am. Min., 65, 1044 (1980); [17] M.B.Madsen et al., Nature, 321, 501 (1986); [18] T.J. Wdowiak & D.G.Agresti, Nature, 311, 140 (1984); [19] J.T.Wasson et al., GCA, 40, 1449 (1976); [20] Research supported by NASA grant NAGW-2037.