1	REVISION 1		
2	Morin-type transition in 5C pyrrhotite		
3	C R S Haines ^{1*} , G I Lampronti ¹ , W T Klooster ² , S J Coles ² , S E Dutton ³ , and M A Carpenter ¹		
4	¹ Department of Earth Sciences, University of Cambridge, Cambridge, UK		
5	² UK National Crystallographic Service, Chemistry, Faculty of		
6	Natural and Environmental Sciences,		
7	University of Southampton, SO17 1BJ, UK		
8	³ Department of Physics, University of Cambridge, Cambridge, UK		
9	Abstract		

10 We report the discovery of a low temperature spin-flop transition in 5C pyrrhotite at ~155 K 11 that is similar to those seen in hematite at 260 K and FeS (troilite) at 440 K. The 5C crystal was 12 produced by annealing a 4C pyrrhotite crystal at 875 K, to produce a change in the vacancy-13 ordering scheme that developed during cooling. The 5C structure is confirmed by single crystal 14 x-ray diffraction and the stoichiometry and homogeneity by electron microprobe and SEM BSE 15 mapping. RUS, heat capacity and magnetisation measurements from room temperature down to 16 2 K are reported. The transition is marked by a steep change in elastic properties at the transition 17 temperature, a peak in the heat capacity and weak anomalies in measurements of magnetisation. 18 Magnetic hysteresis loops and comparison with the magnetic properties of 4C pyrrhotite suggest 19 that the transition involves a change in orientation of moments between two different 20 antiferromagnetic structures, perpendicular to the crystallographic *c*-axis at high temperatures 21 and parallel to the crystallographic *c*-axis at low temperatures. The proposed structures are

- 22 consistent with a group theoretical treatment that also predicts a first order transition between the
- 23 magnetic structures.

24 I. INTRODUCTION

25 The pyrrhotite system comprises a number of related structures having compositions Fe_{1-x}S 26 where 0 < x < 0.125. The best known of these structures are troilite (FeS) and 4C pyrrhotite 27 (Fe₇S₈). 4C is so called because it is a superstructure with a 4 cell repeat along the c-axis with 28 respect to the parent NiAs structure (Bertaut 1953; Powell 2004). Other commensurate 29 superstructures have been reported as 3C (Fleet 1971, Nakano et al 1979), 5C (de Villiers et al 30 2009, Elliot 2010, Liles and de Villiers 2012) and 6C (Koto et al 1975, de Villiers and Liles 31 2010), while some are incommensurate (Nakazawa and Morimoto 1971, Morimoto et al 1975, 32 Yamamoto and Nakazawa 1982, Izaola et al 2007). Ferrimagnetic pyrrhotite (4C) is an important 33 carrier of magnetic remanence on earth and possibly also on Mars (e.g. Martin-Hernandez et. al 34 2008; Rochette et. al 2005), but 4C structures are not unique and the other superstructure types 35 have different structure/property relationships which are much less well known. The key issue in 36 relation to the magnetic structures are how vacancies order on the cation sites as the 37 stoichiometry changes and how the distribution of vacancies then controls the magnetic 38 structures.

39 By adopting a group theoretical approach, Haines et al. (2019a) have shown that the 40 commensurate superstructures of pyrrhotite, which have different Fe/vacancy ordering schemes, 41 are all likely to undergo a magnetically driven spin reorientation transition with accompanying 42 small distortions of the crystal lattice at low temperatures. The prediction for the case of 5C 43 pyrrhotite is for an abrupt spin-flop transition similar to the Morin transition in hematite. The 44 present work focuses on the magnetic and elastic properties of a natural 4C crystal that 45 transformed unexpectedly to 5C crystal after being heated above its concomitant vacancy ordering and Néel temperature of 595 K. We report a newly identified magnetic transition at 46 47 ~155 K in the 5C crystal and propose that it is a spin-flop transition between two

48 antiferromagnetic structures, with an origin that is closely related to that of the magnetic 49 transitions already known in FeS at 440 K and 4C Fe₇S₈ at 35 K.

50 In general, 5C pyrrhotite is found to have a composition close to Fe_9S_{10} . However, Pósfai et. 51 al. (2000) reported a 5C polytype with composition Fe₇S₈. Kontny et. al. (2000) also presented 52 some x-ray diffraction and magnetisation measurements on multiphase samples, suggesting the 53 possibility of pyrrhotite with a 5C structure having the stoichiometry close to the Fe₇S₈ that is 54 usually associated with the 4C structure. De Villiers et al. (2009) proposed space group number 55 63 Cmce (formerly Cmca) as the correct crystallographic space group at room temperature. 56 Elliot (2010) had claimed $P2_1/c$ for the room temperature structure. Liles and de Villiers (2012) 57 subsequently proposed P_{2_1} for the structure at both room temperature and 120 K. There do not 58 appear to have been any studies of the magnetic structure.

In this work, we first establish the stoichiometry of the sample through electron microprobe analysis before going on to present a structural solution in the space group 63 *Cmce* based on single crystal diffraction data collected on a single crystal sample of 5C pyrrhotite at room temperature. We then present heat capacity, DC magnetisation and resonant ultrasound spectroscopy (RUS) measurements on a second single crystal over the range 2-300 K.

64 II. SAMPLE PREPARATION

The single crystal sample of pyrrhotite used in this study was cut from a cm-sized crystal in the mineral collection of the South Australia Museum that had originated from a mine in Mexico. Samples from the same parent crystal have been described in Haines et al (2019c). At the start, it was nearly in the shape of a rectangular parallelepiped with two pairs of parallel faces 1.7 mm and 1.5 mm apart and a third pair of non-parallel faces separated by between 1 mm and 1.5mm. The first experiment was to collect RUS spectra through the temperature interval 10 - 295 K to confirm that it was 4C pyrrhotite with a Besnus transition (Besnus and Meyer 1964) at
~35 K (see Haines et al. 2019b).

73 After removal from the RUS instrument, the crystal was heated slowly to 875 K in flowing 74 argon, with an oxygen trap, over four and a half days before cooling it back to room temperature 75 over three days. The sample was held between two alumina rods. The heating and cooling rates 76 through the 595 K transition were 2.2 K/hour. As a consequence of this treatment, the crystal 77 acquired a thin oxide layer that was removed mechanically, and also broke into two pieces. The 78 larger piece had a mass of 9.1±0.1 mg and was used for heat capacity, magnetisation and RUS 79 measurements. On completion of these measurements, the larger piece was mounted and 80 polished for chemical analysis. The smaller piece weighed 2.2 ± 0.1 mg and was broken up for 81 the single crystal diffraction measurements.

82 III. ELECTRON MICROPROBE ANALYSIS

Chemical analysis on the larger piece was carried out using a Cameca SX100 electron 83 84 microprobe in the Department of Earth Sciences, University of Cambridge (instrumental 85 conditions: 20 keV, 10 nA, 1µm beam diameter). With respect to 10 sulphur atoms, the average 86 of 20 analyses gave the number of Fe atoms as 9.21±0.05. Co, Cu, Mn, Ni and Zn were 87 measured for, but all were below the detection limit on all of the 20 analysis spots. We highlight 88 here that this shows a significant change in stoichiometry from the parent crystal described in 89 Haines et al. (2019c) in which electron microprobe analysis revealed a homogeneous and pure 90 4C crystal of stoichiometry 7.00(6) Fe atoms per 8 sulphur atoms. We suggest that this change 91 of stoichiometry is due to the loss of sulphur from the crystal rather than the addition of iron for 92 which no source or mechanism is present. To check for zoning or inhomogeneity in the 93 stoichiometry of the crystal a scanning electron microscope back-scattered electron (SEM-BSE) 94 analysis map of nearly the whole surface of the same piece used for the electron microprobe analysis was collected. The SEM-BSE map showed no sign of any chemical inhomogeneity above the level of detection. We therefore conclude that the transformation from stoichiometry $Fe_{7.00(6)}S_8$ to $Fe_{9.21(5)}S_{10}$ is complete and homogeneous. The images are included as supplemental information (SI 2020).

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100 IV. SINGLE CRYSTAL X-RAY DIFFRACTION

101 The single crystal used for X-ray diffraction was black and plate-shaped, with dimensions 102 $\sim 0.080 \times 0.040 \times 0.020 \text{ mm}^3$. It was mounted on a MITIGEN holder in perfluoroether oil on a Rigaku FRE+ diffractometer equipped with VHF Varimax confocal mirrors and an AFC12 103 104 goniometer and HyPix 6000HE detector. The temperature was 293 ± 2 K during scans of 0.5° 105 per frame for 1.3 s with Mo K radiation (Rotating-anode X-ray tube, 40 kV, 30 mA). The total 106 number of runs and images was based on the strategy calculation from the program CrysAlisPro (Rigaku, V1.171.39.46, 2018). The maximum resolution achieved was at 27.478° (0.77 Å). Cell 107 108 parameters were retrieved using the CrysAlisPro (Rigaku, V1.171.39.46, 2018) software and 109 refined using CrysAlisPro (Rigaku, V1.171.39.46, 2018) on 5787 reflections, 22% of the 110 observed reflections. Data reduction, scaling and absorption corrections were performed using CrysAlisPro (Rigaku, V1.171.39.46, 2018). The final completeness is 99.7 % out to 25° in Θ . A 111 112 multi-scan absorption correction was performed using CrysAlisPro 1.171.39.46 (Rigaku Oxford 113 Diffraction, 2018) using spherical harmonics as implemented in SCALE3 ABSPACK. The absorption coefficient of this material is 13.664 mm^{-1} at this wavelength (= 0.711Å) and the 114 115 minimum and maximum transmissions are 0.522 and 1.000.

Figure 1 shows a section of the (0*kl*) plane of the diffraction pattern, with 5 superlattice reflections clearly visible along the $c^*(k_z)$ of the parent NiAs cell. We looked for evidence of incommensuration by initially indexing the pattern with the NiAs unit cell and then using an

119	incommensurate k-vector to describe the remaining peaks. The k-vector which describes the
120	pattern has components of 0.500(3) along the planar hexagonal axes and 0.204(6) perpendicular
121	to the ab-plane of the parent structure. This is therefore commensurate within experimental
122	uncertainty.
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126 The structural solution and subsequent refinement were undertaken using the SHELXS 127 program (Sheldrick 1997), proceeding with 9 Fe and 8 S atomic positions. Neutral scattering 128 factors were used for all atoms. The appropriate 3-individual twin matrix describing a 120° 129 rotation along the c axis was introduced in the first stages of the refinement; since the three 130 individual fractions were found to be all close to 33%, this step was necessary to take the 131 discrepancy factor R1 below 0.2. Individual occupancy factors for the Fe atoms were refined 132 with an overall isotropic thermal parameter to detect the partially vacant sites. The Fe sites that 133 refined to occupancies close to 1.0 were then fixed at unity, and the thermal displacement 134 parameters were then refined independently. Lastly, the ADPs were refined anisotropically, 135 together with the occupancies of the four partially vacant Fe atoms. Because of an unreasonably 136 prolate ADP for the atom at (0,0,0) Fe9 the thermal displacement parameters were restrained to 137 be similar to those of the neighbour Fe sites. We here highlight that this is not the same position 138 found to be vacant in the structural refinement performed by De Villiers et al. (2009), 139 corresponding to Fe4 in our model. Details of the structural solutions are given in the 140 accompanying cif file.

The refinement converged with a discrepancy factor R1 = 0.0485 for 1675 observed reflections and R1 = 0.0604 for all 2084 measured reflections and 120 parameters. The goodness of fit parameter was S = 1.476 and wR2 was 0.3431. No extinction correction parameter was used as it was found to converge to zero. The largest residual electron density amounted to 1.32 e/Å. Interestingly, the structural model reported by De Villiers et al. (2009) also differs in the way the Fe planes stack along the [001] direction (see Figure 2).

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The composition, calculated from the refinement of the structure, is $Fe_{9.22(1)}S_{10}$, in excellent agreement with the results of the electron microprobe analysis. Details of the refinement and the comparison to that of De Villiers et al. (2009) are given in Table 1. A cif file containing detailed information about data collection and structural refinement is supplied separately.

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156 V. HEAT CAPACITY

Heat capacity measurements were carried out in a Quantum Design PPMS. The 9.1mg sample was fixed to the measurement platform of the heat capacity puck with Apiezon N grease. The puck (plus grease) was measured through the temperature interval 2-300 K with and without the sample to determine the background contribution needed to obtain the heat capacity of the sample alone. At low temperatures the heat capacity of the sample was larger than the background but at temperatures above ~30K the background was larger.

163 Figure 3a shows a peak in the heat capacity of the crystal near 155 K. Broad features at 215 K 164 and 280 K are artefacts due to the Apiezon grease. A closer view of the peak (Fig. 3b) reveals it 165 to be at 154 K with an apparent onset temperature of \sim 147 K and a tail up to \sim 157 K. Repeat 166 measurements in a field of 9 T applied parallel to the [100] direction of the crystal (H//a of the NiAs cell) yielded a peak at 148 K (Fig. 3b). The shape of the peak was unchanged, but the 167 transition had clearly been shifted down in temperature by ~6 K. There is no evidence in the data 168 for the Besnus transition at 35 K, which confirms that the crystal had fully transformed from the 169 170 4C structure. The peak is asymmetric. There is a shoulder on the low temperature side suggesting that there are two contributions to the peak. This is somewhat similar to the heatcapacity anomaly recently observed in 4C pyrrhotite (Haines *et al.* 2019b).

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174 VI. MAGNETISATION

175 Magnetisation was measured using a Quantum Design MPMS. Zero field cooled (ZFC) data 176 were collected by cooling the crystal in zero applied field to 2 K before applying a 10 mT field 177 (H/a) and measuring the moment in a continuous heating sweep up to 300 K at 3K/min. Field 178 cooled (FC) data were collected by cooling the crystal to 2 K in an applied field of 5 T before 179 removing the 5T field and applying a 10 mT field (H/a) during measurements of the moment in 180 a continuous heating sweep up to 300 K at 3K/min. There are two distinct features observable in 181 both the ZFC and FC magnetisation curves, at ~ 120 and ~ 155 K. The feature at ~ 155 K cannot 182 be explained by any obvious impurity and matches up well with the transition seen in the heat 183 capacity of the 5C crystal. The feature at \sim 120 K is almost certainly from the Verwey transition 184 of magnetite, revealing the presence of a small amount of magnetite as an impurity phase. This is 185 most likely remaining from incomplete removal of the thin oxidised layer, following the heat 186 treatment at 875 K. The same crystal was used for both magnetisation and heat capacity 187 measurements, and the lack of any anomalies in the heat capacity at ~120 K supports the 188 conclusion that the proportion of magnetite was very small.

In order to put the absolute values of magnetisation in context, results for the 5C crystal and for a 4C crystal cut from the parent crystal temperature are compared in Figure 4b. 4C pyrrhotite is ferrimagnetic, due to an uncompensated moment arising from the fact that one in eight Fe sites are vacant. The measured moment of the 4C crystal at room temperature is a factor of ~100 larger than that of the 5C crystal, suggesting that the magnetic ground state of 5C pyrrhotite is a fully compensated antiferromagnet. Furthermore, this puts an upper limit on the amount of 4C

195 pyrrhotite remaining in the sample of approximately 1%. In the ZFC data the step-like feature at 196 155 K takes place between 148 K and 155 K, with a midpoint of 152 K. The start stop and mid-197 points of the step in the FC data are the same, though there is a small kink at the mid-point that is 198 not observed in the ZFC data.

199 Figure 5a contains hysteresis (M-H) loops measured between +7 and -7 T at two 200 temperatures well above and two temperatures well below the magnetic transition. The values of 201 M never exceed $\sim 2 \text{ Am}^2/\text{kg}$ and are smaller at all fields by a factor of at least ten in comparison 202 with the 4C structure (Powell et. al 2004; Charilou et. al 2015). They also show no indication of 203 saturation at any temperature, consistent with antiferromagnetic structures. Above the transition, 204 the magnetisation at 7 T (1.8 Am^2/kg) is more than 4 times larger than it is below the transition 205 (0.4 Am²/kg) at the same field. Below 150 K, [100] is a relatively hard direction, but becomes 206 relatively soft above ~150 K. The difference remains marked even at 140 and 160 K, but there is 207 a metamagnetic transition at 150 K from the low temperature magnetic structure, at low fields, to 208 the high temperature structure at high fields (Fig. 5b). This is permissive of a first order spin-flop 209 transition between two antiferromagnetic structures that have their moments aligned in the plane 210 perpendicular to the *c*-axis above the transition point and parallel to the *c*-axis below it. In this 211 scenario, small openings of the curves at low fields (Fig. 5c) and the small hysteresis seen in the 212 magnetisation curves at all temperatures are attributed to the magnetite impurity identified on the 213 basis of the magnetic anomaly at \sim 120 K (Fig. 3a). In addition, the (near) saturation moment of 214 these two impurities (at a field of 1.5 T) would limit their presence in the sample to similar 215 values. If the entire signal at 300 K and 1.5 T were attributed to magnetite (this is around 0.5 216 Am^2/kg) this would equate to 46 µg of magnetite or 0.5% of the total mass, using a value of 90 217 Am²/kg for saturation magnetisation of magnetite (Hunt et al. 2103 and references therein). In a

similar way we can constrain the 4C pyrrhotite to be less than 2.5% of the total mass (using 20 Am^2/kg as the value of the magnetisation from Hunt et al. (1995) and Haines et al (2019b).

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221 VII. RESONANT ULTRASOUND SPECTROSCOPY

222 Details of the instrument used to collect RUS data at low temperatures have been given 223 elsewhere (McKnight et. al 2007). The sample sits lightly between two piezoelectric transducers 224 in the RUS head, which is attached to the end of a stick and lowered into an Orange helium flow 225 cryostat. In the present study, resonance spectra were collected in two automated cooling and 226 heating sequences with steps of 30 K on cooling from room temperature to 10 K, and of between 227 1K and 10K on heating from 10 K to 295 K. A settle time of 1200s was included to allow for 228 thermal equilibration at each set point. Individual spectra contained 65,000 data points in the 229 frequency range 100 kHz - 2 MHz, with a driving voltage of 25 V. One sequence was for a 230 different piece of the original large 4C crystal and the second was for the 9.1 mg fragment of the 231 5C crystal. Analysis of elastic and anelastic properties was undertaken by fitting selected peaks 232 with an asymmetric Lorentzian function to give peak frequency, f, and width at half-maximum 233 height, Δf . The natural resonances of a mm sized sample involve primarily shearing motions and the variations of f^2 therefore reflect variations of different combinations of predominantly shear 234 235 elastic constants, in this case of the single crystal. Acoustic dissipation is expressed in terms of the inverse mechanical quality factor, Q^{-1} , which, in a RUS experiment is usually specified as 236 237 $\Delta f/f$.

Figure 6 contains a stack of spectra from the heating sequence of the 5C crystal. The y-axis is amplitude, but the spectra have been offset in proportion to the temperatures at which they were collected. The axis has then been relabelled as temperature in order to allow ready visualisation of the principal features of the elastic behaviour. All the resonances show the same pattern of 242 reducing frequency (elastic softening) as the transition point is approached from above, followed 243 by abrupt increases (elastic stiffening) at the expected transition temperature of ~155 K. In other 244 words, all the shear elastic constants of the crystal evolve in more or less the same manner. Blue curves are fits to individual peaks, and Figure 7a shows the variations of f^2 and O^{-1} for a peak 245 246 near 650 kHz. With falling temperature there is a trend of elastic softening by a few percent 247 towards the transition point, followed by $\sim 10\%$ stiffening in a narrow temperature interval 248 between 150 and 160 K. This temperature interval corresponds almost exactly with the 249 temperature interval over which the anomaly in heat capacity extends (Fig. 3). The peaks generally remain quite sharp at all temperatures, with values of Q^{-1} in the vicinity of 2×10^{-3} apart 250 251 perhaps from a slight increase in the vicinity of the transition point. The variation of f^2 252 normalised to the room temperature value for three different peaks is shown in Figure 7b. Whilst 253 the overall temperature dependence is similar the peaks behave slightly differently, as would be 254 expected for a single crystal sample in which each resonance has a distinct contribution from the 255 different elastic constants.

Variations of f^2 and Q^{-1} data from fits to a resonance peak near 620 kHz in spectra collected 256 257 from the piece of original the 4C crystal are shown in Figure 7c. The crystal was cut from the 258 same original sample as the piece that was converted to 5C. There are very obvious differences 259 from the variations in elastic and anelastic properties that accompany the magnetic transition 260 shown in Figure 7. In particular, softening by a few % with falling temperature is associated with a peak in O^{-1} at ~270 K. This turns into a trend of gradual stiffening followed by steep softening 261 ahead of the Besnus transition, which is marked by a minimum at ~36 K. f^2 values then recover 262 steeply and there is a distinct peak in Q^{-1} immediately below the transition. The maximum 263 change in f^2 at the frequency of this resonance, ~620 kHz, is ~6%. Changes in elastic and 264

anelastic properties accompanying the Besnus transition in 4C pyrrhotite are described in detailelsewhere (Haines et al 2019b).

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268 VIII. DISCUSSION AND CONCLUSIONS

269 We have shown that 4C pyrrhotite can be transformed to 5C by annealing above the 270 concomitant Néel and vacancy ordering temperature. On the basis of single crystal x-ray 271 diffraction at room temperature we confirm that a model in space group *Cmce* can successfully 272 explain the diffraction data. The low temperature structure has previously been refined 273 successfully using a model in space group $P2_1$ by Liles and deVilliers (2012). The Besnus 274 transition is no longer observed and a new Morin-like spin-flop transition occurs at ~155 K 275 instead. The magnetic properties of the 5C crystal are incompatible with impurities of 4C 276 pyrrhotite or magnetite above the level of 1%. In addition, the heat capacity anomalies 277 characteristic of the low temperature magnetic phase transitions in these two minerals were not 278 observed. The 4C crystal from which the piece used in this study was cut showed no impurities 279 above the level of detection in either powder XRD or electron microprobe measurements 280 (Haines et al. 2019b). Therefore, it is difficult to imagine any impurity other than the magnetite 281 formed at the surface or 'untransformed' 4C pyrrhotite. We have carried out electron microprobe 282 analysis on the 5C crystal and the sample is monophase with a high degree of homogeneity. The 283 impurity levels are on the border of detectability. The stoichiometry is homogeneous and works 284 out to be $Fe_{9,2}S_{10}$ in good agreement with the single crystal refinement.

The most significant structural difference in comparison with the 4C structure is the change in vacancy ordering scheme which develops during cooling. Without specifying the ordering in detail, it is possible to assess the changes in structure and properties from the perspective of symmetry. If the reference structure at high temperatures is taken to be that of NiAs in space 289 group $P6_3/mmc$, the order parameter which gives rise to the 5C superstructure is most likely to 290 have the symmetry of the active representation U1(1/2,0,1/5) (Haines et al 2019a). A likely 291 candidate for the magnetic order parameter has the symmetry of irreducible representation $m\Gamma_5^+$. 292 The list of possible magnetic space groups that can result from this combination includes the 293 previously reported crystallographic space groups *Cmce*, $P2_1/c$ and $P2_1$ (Haines et. al 2019a). 294 New magnetic and elasticity data have been interpreted here on the basis that 5C pyrrhotite is 295 antiferromagnetic and that the magnetic transition at ~ 155 K is from one antiferromagnetic 296 structure to another. In all of the reported space groups the moments can be aligned either in the 297 plane perpendicular to the crystallographic c-axis of the parent or parallel to the c-axis. Across 298 the Fe_{1-x}S phase diagram (Schwarz and Vaughan 1972) the 590 K transition is to a state in which 299 the moments are in the plane perpendicular to the crystallographic *c*-axis. The low temperature 300 spin-flop transition observed in troilite and the low temperature evolution of 4C pyrrhotite 301 involve a rotation of the moments towards the *c*-axis.

Haines et. al (2019a) predicted that the magnetic transition in 5C would be an abrupt spinflop transition. This is because for none of the experimentally reported structures is it possible to have distortions which allow both $m\Gamma_4^+$ and $m\Gamma_5^+$ to have non-zero values. In other words, the moments are strictly confined to either the plane perpendicular to the *c*-axis or to the direction parallel to the *c*-axis. *Cmce* was reported for a crystal at room temperature with the P2₁ structure being solved using data collected at 120 K. These two structures do not have a group-subgroup relationship so that a *Cmce* – P2₁ transition is necessarily first order.

The spin-flop transition in 5C pyrrhotite is significantly different from the Besnus transition in 4C pyrrhotite, as is seen clearly in differences between the two sets of elastic and anelastic data in Figure 7. As a function of temperature, the spin-flop transition is characterised by a steep increase in all the shear elastic constants over a narrow temperature interval, consistent with first

313 order character. The low temperature structure is stiffer than the high temperature structure and 314 has the same level of acoustic loss. In contrast, the Besnus transition occurs over a wider 315 temperature interval, consistent with a second order transition between magnetic structures that 316 do have a group-subgroup relationship. There is also a distinct peak in the acoustic loss that 317 could relate to the development of a ferroelastic microstructure. Thus, the transition in the 5C 318 crystal appears to be closely analogous to the magnetic transition to that seen in FeS at 440 K, 319 which involves the same change in orientation of moments (Harihara and Murakami 1958, 320 Andresen 1960, Sparks et. al 1960 and 1962, Andresen et. al 1967, Horwood et al 1976).

321 This combination of symmetry analysis and experimental data on the magnetoelastic 322 behaviour shows that 5C pyrrhotite fits well into the wider picture for the diverse structures 323 suggested for the system $Fe_{1-x}S$ by Haines et (2019a). The al stable 324 antiferromagnetic/ferromagnetic structure which appears across the solid solution at ~590 K has 325 moments aligned perpendicular to the crystallographic *c*-axis. At transition temperatures which 326 reduce from ~440 K with increasing concentration of vacancies, this gives way to another 327 magnetic structure in which the preferred orientation of the moments is 90° away from the c-328 axis. Details of the precise magnetic structure and possibilities for thermodynamically 329 continuous pathways between the two magnetic states then depend on details of the vacancy 330 ordering at different stoichiometries, as expressed in terms of order parameters with irreducible 331 representations of the form U(1/2,0,1/x) where x is 3, 4, 5, 6 or irrational, corresponding to the 332 3C, 4C, 5C, 6C and incommensurate phases of pyrrhotite. All these phases, apart from 3C, are 333 expected to show one magnetic transition below room temperature. In 3C pyrrhotite the 334 transition is expected to be above room temperature.

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336 IX. IMPLICATIONS

337 The discovery of a low temperature spin-flop transition in the 5C superstructure of pyrrhotite 338 must be seen within the context of the whole $Fe_{1-x}S$ family. In this work we have shown that the 339 behaviour of the 5C superstructure is consistent with a comprehensive group theoretical 340 framework described in Haines et al. (2019a). The result fits with the 3C superstructure being 341 ferrimagnetic at room temperature and, if the crystallographic space group of P3₁21 (Nakano et 342 al. 1979; Keller-Besrest et al. 1983) is correct, with the magnetic moments aligned parallel to 343 the *c*-axis. A low temperature spin-reorientation transition like the Besnus transition must also 344 be expected in the 6C superstructure. Thus, it seems that the spin flop transitions are all closely 345 related and fundamentally due to the same underlying thermodynamic driving force.

An additional consideration relates to the fact that the 5C crystal studied in this work was the result of thermally cycling a 4C crystal through the concomitant vacancy ordering and ferrimagnetic ordering temperature, at 595 K, and then up to 875 K before cooling back to room temperature. To our knowledge, this is the first time that 5C pyrrhotite has been synthesised in this way.

Finally, our improved understanding of the nature of the vacancy ordering and related magnetic structures in the $Fe_{1-x}S$ system means that an understanding of the microscopic mechanisms may now be within reach for an important group of mineral sulphides. Further experiments and calculations should focus on the changes in electronic structure behind the changing magnetic anisotropy and its dependence on temperature, pressure and vacancy order.

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413			
414	Figure captions		
415			
416			
417	Figure 1. Part of the $(0kl)$ precession image of the room temperature single crystal X-ray		
418	diffraction pattern, showing the five-fold superlattice reflections of the 5C structure. The data		
419	has been reduced using the cell from the <i>Cmce</i> structure and the peaks are labelled with the (<i>hkl</i>)		
420	of the <i>Cmce</i> structure. The direction of the b^* and c^* are given in red on the left. The unit cell of		
421	the Cmce structure is shown by the grey grid. The projection of the hexagonal parent cell onto		
422	the <i>Cmce</i> $(0kl)$ plane is shown in blue. The five-fold modulation in the <i>l</i> direction can be seen		
423	and the peaks characteristic of this are labelled with their (<i>hkl</i>) values.		
424			
425			
426	Figure 2. Structural model of pyrrhotite 5C viewed along [100] from this work (top) and from de		
427	Villiers et al. (2009) (bottom). Color code: yellow = S sites; brown = Fe sites (partial occupancy		
428	is shown by half-grey half-brown); pink = partially Ni replaced Fe site (De Villiers et al., 2009);		
429	blue = vacant metal site from De Villiers et al. (2009) (see main text for details).		
430			
431	Figure 3. (a) Heat capacity of 5C crystal measured between 2 and 300 K in zero field,		
432	showing the magnetic transition near 155 K. Anomalies at \sim 215 and \sim 280 K are artefacts. (b)		

433 Higher resolution view of the peak at 154 K in zero field, which shifts to 148 K when measured 434 in a 9 T field (H//a).

435

Figure 4. Magnetisation, *M*, of 5C crystal. (a) ZFC and FC data, measured in a field of 10 mT (*H*//*a*). The anomaly at ~120 K is due to the presence of a small amount of magnetite (labelled as T_V for the Verwey transition) as impurity phase. The previously unknown transition is labelled as T_x . (b) Comparison with ZFC data for 4C and 5 C crystals (*H*//*a*), showing a difference in magnetisation of a factor of ~100.

441

Figure 5. (a) Field and temperature dependence of the magnetisation of the 5C sample. The field was applied parallel to the [100] direction of the parent NiAs structure unit cell (H//a). The structure at 200 and 300 K has marked differences in magnetic anisotropy from the structure at 5 and 100 K. Neither show any indication of saturation at high fields. (b) Field and temperature dependence of magnetisation of the 5C structure close to the magnetic transition at ~155 K (H//a). (c) Close in view of the low field region from (b), showing slight opening of the loops that is attributed to the presence of a small amount of magnetite.

449

Figure 6. RUS spectra collected from the 9.1 mg piece of the 5C crystal in a heating sequence from 10 to 295 K. The y-axis should be amplitude in volts, but the spectra have been offset in proportion to the temperatures at which they were collected and the axis relabelled as temperature. There is an abrupt change in frequencies of all the individual resonance peaks at the expected transition temperature, ~155 K. Fits to selected peaks are shown in blue.

455

Figure 7. (a) Temperature dependence of f^2 and Q^{-1} for the single crystal of 5C pyrrhotite from fitting of the resonance peak near 650 kHz and (b) f^2 for three resonance peaks at 660 kHz, 780 kHz and 840 kHz normalised to the value at room temperature. (c) Temperature dependence of f^2 and Q^{-1} for a single resonance with frequency near 620 kHz from a 4C pyrrhotite crystal cut

- 460 from the same original sample as was used to produce the 5C crystal. The Besnus transition is
- 461 marked by a minimum if f^2 at ~36 K. Crosses = Q^{-1} , right axis; filled circles = f^2 , left axis.

462

463 Tables

464

465Table 1. Crystal data and structure refinement details for 5C pyrrhotite.

This work	De Villiers et al. (2009)
Fe _{9.22(1)} S ₁₀	$Fe_{8.79}Ni_{0.118}S_{10}$
6682.51	6588.86
293(2)	293(2)
0.71073	0.71073
Cmce (formerly Cmca)	Cmce (formerly Cmca)
a = 6.890(2) Å	a = 6.893(3) Å
b = 11.9436(4) Å	<i>b</i> = 11.939(3) Å
c = 28.7916(10) Å	c = 28.635(15) Å
$\alpha = 90^{\circ}$	$\alpha = 90^{\circ}$
$\beta = 90^{\circ}$	$\beta = 90^{\circ}$
$\gamma = 90^{\circ}$	$\gamma = 90^{\circ}$
2371.36(13) Å ³	2356.4(15) Å ³
8, 4.689 g/cm ³	8, 4.643 g/cm ³
12.681	12.516
3358	3154
0.08x 0.04 x 0.02 mm ³	0.14 x 0.06 x 0.03 mm ³
2.12-31.82°	2.85-26.34°
$-9 \le h \le 9, -16 \le k \le 15, -40$	$-8 \le h \le 7, -8 \le k \le 14, -33 \le$
$\leq l \leq 42$	$l \leq 29$
29278/2084 [R(int) =	6041/1264 [R(int) = 0.0322]
0.0322]	
99.7%	99.2%
Full-matrix least-squares on	Full-matrix least-squares on
F^2	F^2
2084 / 1 / 120	1264 / 0 / 109
1.477	1.225
R1 = 0.0485, wR2 = 0.3231	R1 = 0.0716, wR2 = 0.1513
R1 = 0.0604, wR2 = 0.3431	R1 = 0.0937, wR2 = 0.1627
-	0.00003(2)
	1 500 1 0 105 5 8 -3
1.445 and $-1.346 e^{-1} A^{-3}$	$1.533 \text{ and } -2.125 \text{ e} \cdot \text{A}^{-3}$
	This work Fe _{9.22(1)} S ₁₀ 6682.51 293(2) 0.71073 <i>Cmce</i> (formerly <i>Cmca</i>) a = 6.890(2)Å b = 11.9436(4)Å c = 28.7916(10)Å $a = 90^{\circ}$ $\beta = 90^{\circ}$ $\gamma = 90^{\circ}$ 2371.36(13)Å ³ 8,4.689 g/cm ³ 12.681 3358 0.08x 0.04 x 0.02 mm ³ 2.12-31.82° $-9 \le h \le 9, -16 \le k \le 15, -40$ $\le l \le 42$ 29278/2084 [<i>R</i> (<i>int</i>) = 0.0322] 99.7% Full-matrix least-squares on F^2 2084 / 1 / 120 1.477 <i>R</i> 1 = 0.0485, <i>wR</i> 2 = 0.3231 <i>R</i> 1 = 0.0604, <i>wR</i> 2 = 0.3431 -

Figure 1



Figure 2





Figure 3a

a



Figure 3b



Figure 4a



Figure 4b



Figure 5a



Figure 5b



Figure 5c



Figure 6



Figure 7a



Figure 7b



Figure 7c

