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2	"Fe ₅ S ₂ identified as a host for sulfur in Earth's core"
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Fe₅S₂ identified as a host for sulfur in Earth's core

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48 Abstract

49 Planetary habitability, as we experience on Earth, is linked to a functioning geodynamo 50 which is in part driven by the crystallization of the liquid iron-nickel-alloy core as a planet cools 51 over time. Cosmochemical considerations suggest that sulfur is a candidate light alloying 52 element in rocky planetary cores of varying sizes and oxidation states; such that, iron sulfide 53 phase relations at extreme conditions contribute to outer core thermochemical convection and inner core crystallization in a wide range of planetary bodies. Here we experimentally investigate 54 55 the structural properties of the Fe-S system and report the discovery of the sulfide, Fe₅S₂, 56 crystallizing in equilibrium with iron at Earth's outer core pressures and high temperatures. 57 Using single-crystal X-ray diffraction techniques, Fe_5S_2 was determined to adopt the complex 58 Ni₅As₂-type structure ($P6_{3}cm$, Z = 6). These results conclude that Fe₅S₂ is likely to crystallize at 59 the interface of Earth's core and mantle and will begin to crystallize during the freezing out of 60 Earth and Venus' core overtime. The increased metal-metal bonding measured in Fe₅S₂ compared to the other high *P*-*T* iron sulfides may contribute to signatures of higher conductivity 61 62 from regions of Fe₅S₂ crystallization. Fe₅S₂ could serve as a host for Ni and Si as has been

63 observed in the related meteoritic phase, perryite, (Fe, Ni)₈(P, Si)₃, adding intricacies to

64 elemental partitioning during inner core crystallization. The stability of Fe_5S_2 presented here is

65 key to understanding the role of sulfur in the multicomponent crystallization sequences that drive

66 the geodynamics and dictate the structures of Earth and rocky planetary cores.

67

68 Introduction

69 Earth and the other terrestrial planets along with rocky exoplanets that continue to be discovered are composed of silicate mantles and iron-rich metallic cores (e.g. Birch 1952; Scott 70 71 and Wasson 1975; Jephcoat and Olsen 1987; Mcdonough and Sun 1995; Rubie et al. 2011). The 72 likelihood of planetary habitability as we know it demands the presence of a geomagnetic field, 73 sustained by a convecting metallic liquid core, that protects the planet from harmful cosmic rays 74 (e.g. Buffet, 2000). In Earth's liquid outer core, the presence of light alloying elements 75 introduces complex pressure-temperature-dependent melting and crystallization thermodynamics 76 that drives core convection (Fearn and Loper, 1981; Stevenson 1981; 1988; Nimmo 2015). 77 While there remains no method to directly sample Earth's or any terrestrial planetary core, iron 78 meteorites are recognized as relics of the cores of planetesimal building blocks of terrestrial 79 planets, and have long provided insight into the cosmochemically abundant light elements 80 contributing to planetary core dynamics (e.g. Scott and Wasson 1975; Malvin et al. 1984). Of the 81 candidate core-alloying elements such as Si, O, S, C, and H, sulfur is present in nearly all iron meteorites, suggesting that sulfur is a core alloying element in rocky planets with varying sizes, 82 83 oxidation states, and formation histories (Scott and Wasson 1975; Jones and Drake, 1983; Kruijer et al. 2014). Examination of the structural properties of iron sulfides at high pressures 84

and temperatures (*P-T*) is therefore critical to ascertaining the chemistry and thermodynamics of
Earth, terrestrial and exoplanetary cores.

At present, the crystallographic assessment of iron sulfides at Earth and planetary core 87 conditions remains limited, even though properties such as density and elasticity are directly 88 89 related to the atomic arrangement of core-crystallizing phases and are critical for assessing the 90 seismic and dynamic observations of planetary cores. Previous work on Fe-rich systems indicate 91 that tetragonal Fe₃S (Fe₃P-type structure) is stable from Martian core conditions to Earth's outer 92 core conditions (Fei et al. 2000; Kamada et al. 2010; Ozawa et al. 2013; Mori et al. 2017), and an 93 orthorhombic Fe₂S (Co₂Si-type structure) phase becomes stable near Earth's inner-core boundary (ICB) conditions (Tateno et al. 2019). The presence of Si has also been observed to stabilize the 94 95 Fe₂(S, Si) chemistry over Fe₃S at Mercurian core-mantle-boundary (CMB) conditions (Tao and 96 Fei 2021). These reports were predominantly based powder X-ray diffraction studies at high 97 pressures and temperatures and chemical analyses of recovered samples. However, recent work 98 has demonstrated that the iron-sulfur phase diagram is more complex at high P-T, but that this 99 complexity requires a superior method for accurately characterizing crystal structures at extreme 100 conditions. Using single-crystal X-ray diffraction techniques on multigrain samples, Co₂P-type 101 Fe_2S (*Pnma*, Z = 4) was recently identified and characterized at 90 GPa (Zurkowski et al. *in* 102 *press*). It remains, however, that these methods have not been applied to iron-sulfide phases 103 synthesized at high temperatures above 90 GPa, leaving the role of sulfur in the crystallization 104 sequences occurring in Venus, Earth and larger rocky exoplanets open to further discovery. 105

107 **Results and Discussion**

108 In the current study, Fe-rich Fe-S compositions were probed using both single-crystal and 109 powder X-ray diffraction techniques in a laser-heated diamond anvil cell to 200 GPa and 3300 K. 110 At high temperatures between 120 and 200 GPa, spanning Earth's CMB to mid-outer-core 111 conditions, crystallization of hcp-Fe (Figure 1a) is observed in the diffraction patterns along with 112 diffuse streaks and sets of closely spaced reciprocal nodes (Figure 1a, Figure S1), suggesting a 113 complex atomic arrangement of the coexisting sulfide. Upon temperature quenching in this 114 pressure range, diffraction images were collected while rotating the DAC across a $+/-17-30^{\circ}$ 115 range (depending on the DAC configuration). Grains of a hexagonal lattice were identified in the 116 reciprocal space with indexed parameters: a = 5.979(3) Å, 11.088(6) Å at 140(2) GPa and 117 3070(180) K (Table S1). Assessment of the systematic absences for the structure factors reduced 118 for this hexagonal sulfide suggests a *P*6₃*cm* space group, and structural solution and positional 119 and displacement parameter refinement converged to the Ni₅As₂-type Fe₅S₂ (Z=6) (Table S2, S3; 120 Figure 2) (Oryshchyn et al. 2011). The Ni₅As₂-type Fe₅S₂ is derived from the Pb₅Sb₂ structure 121 (El-Boragy et al. 1970) where the Fe6 site (Table S3) is split about its position and given half 122 occupancy, changing its Wyckoff site from 6c to 12d (Oryshchyn et al. 2011) (Figure 2, Table 123 S3, Appendix A1). Within error, all sites are fully occupied, except the Fe6 site that is ascribed 124 half occupancy (Oryshchyn et al. 2011).

Additionally, two polytypes associated with stacking variations along the c direction were measured at lower temperatures and in stiff pressure media (Table S1). The Fe₅S₂ polytypism is accompanied by observations of diffuse scattering between reciprocal nodes along the c direction (Figure S1), indicating positional disorder as the atoms configure at high temperatures.

129 Analogous polytypes have been reported in related phases such as Pb₅Sb₂ (Saini et al. 1964).

130 X-ray diffraction from seven heating cycles performed between 100 and 200 GPa and to 131 3300 K provide insight into the Fe-rich Fe-S phase relations at outer core pressures and to high 132 temperatures. In each heating experiment, temperatures near melting were attained, and lattices 133 of Fe₅S₂ were indexed in the reciprocal space upon quenching. By further probing locations 134 across the thermal gradient of the laser heated spots, additional Fe-sulfides were characterized, 135 offering information on the lower temperature Fe-saturated sulfide crystal chemistries (Table S1, 136 S2). These include the Fe₃P-type Fe₃S (*I*-4, Z = 8) phase (Fei et al. 2000; Seagle et al. 2006; 137 Morard et al. 2007; Kamada et al. 2010; Kamada et al. 2012; Thompson et al. 2020), the C23 138 Fe₂S (Co₂P-type, *Pnma*, Z = 4) phase, and the C37 Fe₂S (Co₂Si-type, *Pnma*, Z=4) phase, in lower 139 temperature regions at 119 GPa, 131 GPa, and 140 GPa, respectively (Figure 3; Table S1, S2). 140 Notably, identification and crystal-structure analysis of Fe_5S_2 and Fe_3S in these Fe-saturated 141 experiments clarifies that Fe_5S_2 , not Fe_3S is stable on the solidus above 120 GPa, despite reports 142 from previous powder diffraction studies (e.g. Kamada et al. 2010; Kamada et al. 2012; Ozawa 143 et al. 2013; Mori et al. 2017). 144 Comparison of the interatomic distances and coordination environments in the Fe-145 sulfides observed in this study separates Fe_5S_2 as particularly unique among them. Within an interatomic distance of 2.7 Å (Table S4, Appendix A1-A4), C23 and C37 Fe₂S consist of Fe sites 146

147 coordinated only by S, Fe₃S marks an increase in Fe-Fe bonding with Fe surrounded by up to 9

148 Fe sites, and Fe_5S_2 exhibits Fe surrounded by up to 9 Fe sites with Fe-Fe bond lengths up to 25%

shorter in Fe₅S₂ compared to Fe₃S (Table S4). Following previous descriptions of related M_5X_2

- 150 phases (e.g. (Kjekshus et al. 1973; Oryshchyn et al. 2011), Fe_5S_2 can be viewed as an
- arrangement of 6 Fe sites and 3 S sites with the Fe1, Fe2, Fe4, and Fe6 sites in 13-fold

152	coordination, the Fe3 and Fe5 sites in 12-fold coordination, and the S sites in 10-fold
153	coordination (Figure 2c, Table S4). All sites are coordinated by both Fe and S sites.
154	The presented single-crystal derived structures of Fe ₅ S ₂ , Fe ₃ S, C23 Fe ₂ S, and C37 Fe ₂ S
155	were then used to interpret the changes in powder X-ray diffraction patterns collected during
156	heating between 110 and 200 GPa and to ~3300 K (Figure 4, Figure S4). C23 Fe ₂ S is observed
157	coexisting with Fe to ~125 GPa and up to ~2300 K. In combination with previous studies, $C23$
158	Fe ₂ S is stable at moderate temperatures between 25 and 125 GPa (Zurkowski et al. <i>in press</i> ,
159	Zurkowski et al. submitted). A pressure-induced C23–C37 Fe ₂ S transition is constrained between
160	125 and 135 GPa, in agreement with previous investigations of Fe ₂ S (Zurkowski et al.
161	submitted). C37 Fe ₂ S is stable between 130–200 GPa and to \sim 3000 K in this study, and previous
162	work suggests that C37 Fe ₂ S may remain stable to 306 GPa at high temperatures (Tateno et al.
163	2019). Tetragonal Fe ₃ S forms from the reaction of C23 Fe ₂ S + Fe between 2000–2400 K at ~115
164	GPa. The stability of tetragonal Fe ₃ S is constrained to pressures below ~120 GPa at high
165	temperatures in Fe-rich systems (Figure 4) (Seagle et al. 2006; Morard et al. 2008; Kamada et al.
166	2010; Kamada et al. 2012; Thompson et al, 2020). Fe ₅ S ₂ is observed coexisting with iron
167	between ~120–200 GPa and in the 2400–3300 K range, making it the relevant sulfide in the
168	melting and crystallization properties of Fe-rich metallic cores at these pressures (Figure 4).

Sulfur in Earth and planetary cores 169

The stability of Fe₅S₂ and the phase relations observed in this study present novel 170 171 constraints on the material properties of Fe-sulfides at conditions relevant to Earth's outer core, Venus' core, and exoplanetary cores of similar size and core-mass fraction (CMF). Namely, at 172 pressures ≥ 120 GPa, Ni₅As₂-type Fe₅S₂, not tetragonal Fe₃S as previously believed, is the 173

174	relevant Fe-rich sulfide. As Earth's outer core crystallizes over time, Fe ₅ S ₂ will eventually
175	crystallize over a large depth range up to the CMB. Tetragonal Fe ₃ S has been confirmed up to
176	120 GPa (Figure 4) (Seagle et al. 2006; Morard et al. 2008; Thompson et al., 2020), constraining
177	the Fe ₃ S + Fe \rightarrow Fe ₅ S ₂ + Fe transition to ~120 GPa. Rocky planetary bodies with sizes and CMFs
178	like that of Venus (CMB = 114 GPa) (Aitta 2012) would likely crystallize Fe_3S at the CMB,
179	while Fe_5S_2 will be crystallizing over much of the core's depth. C37 Fe ₂ S is reported to be stable
180	on the solidus above 306 GPa in Fe-rich systems (Tateno et al. 2019), suggesting a Fe ₅ S ₂ + Fe \rightarrow
181	Fe_2S + Fe transition in the 200–300 GPa range. C37 Fe ₂ S remains the candidate Fe-rich sulfide
182	at Earth's ICB, but for planets like Venus with a central pressure of 274 GPa (Aitta 2012), Fe_5S_2
183	may instead be the relevant Fe-rich sulfide, necessitating experimental determination of the
184	$Fe_5S_2 \rightarrow Fe_2S + Fe$ decomposition <i>P-T</i> boundary. Additional assessment of the volume and
185	melting-temperature changes across these newly determined sulfide transitions are critical for
186	matching the density and seismic profiles of Earth and rocky planetary cores. Furthermore, as
187	Fe ₅ S ₂ exhibits a complex crystalline arrangement with significant metal-metal bonding compared
188	to Fe_2S and Fe_3S at core pressures, the significant depth over which Fe_5S_2 crystallizes in and
189	Venus- to Earth-sized planetary cores is likely to contribute to higher conductivity signatures in
190	these regions.

191 Terrestrial core chemistries are multicomponent; Si and Ni are also important core-192 alloying elements in planets such as Earth, drawing attention to the potential stability of the 193 perryite (Fe, Ni)₈(Si, P)₃ structure in (Fe, Ni) – (Si, S)-rich cores. Perryite, is a mineral observed 194 in enstatite chondrites and aubrites that adopts a structure that is trigonal stacking variant of the 195 Ni₅As₂-type structure (Wasson and Wai, 1970; Okada et al. 1991) and iron phosphides tend to 196 adopt analogous structures to that observed in iron sulfides (i.e., Fei et al. 2000; Dera et al. 2008;

197 Gu et al. 2014; Gu et al. 2016; Zurkowski et al., in press). Fe₅S₂ could potentially serve as a host 198 for nickel and silicon in the perryite structure at Earth's outer core conditions. Further, Si has 199 recently been shown to dissolve into iron-sulfide phases and expand their stability fields with 200 pressure (Tao and Fei, 2021). The dissolution of silicon into Fe₅S₂ may expand its stability to 201 higher pressures, making it potentially relevant at Earth's ICB conditions. If this is the case, the 202 partitioning of Si and Ni between (Fe, Ni)₅(S, Si)₂ and (Fe, Ni, Si) and the density difference 203 between the inner-core-crystallizing phase and remaining light-element-rich liquid will be crucial 204 to investigations of the inner-core-density deficit and inner core seismic morphology.

205

206 Materials and Methods

Experiments targeting multigrain synthesis in the Fe-S system consisted of mixtures of Fe (99.9+%, <10 μ m, Alfa Aesar) and iron sulfide (FeS, 99.99%, Alfa Aesar) in Fe+12.5 wt% S (Fe₈₀S₂₀), Fe + 23 wt% S (Fe₆₇S₃₃) compositions. The Fe-FeS compositions were ground in ethanol in a pestle and mortar for 1 hour then mixed dry for a short interval to homogenize any density settling during alcohol evaporation.

Pressure was generated using BX-90-type (Kantor et al. 2012) diamond anvil cells (DAC) 212 213 with type 1 Boehler-Almax conical diamonds and seats, and Mao symmetric-type DACs with 214 Type 1 standard cut diamonds mounted on tungsten carbide of cubic-boron nitride (cBN) seats. 215 Diamond culets ranged from 150-50 µm in diameter. Foils of Fe₈₀S₂₀ or Fe₆₇S₃₃ were produced 216 by pressing the Fe–S starting powders between two ungasketed diamond anvils, then loaded 217 between pellets of KCl or SiO₂. Sample chambers ranged from 180-25 µm in diameter and were 218 drilled from pre-indented rhenium gaskets. Samples were dried at 100°C for 30 minutes prior to 219 pressurization.

220 Angle dispersive X-ray diffraction (XRD) experiments were conducted at Argonne 221 National Laboratory, sector 13 ID-D, of the Advanced Photon Source. At Sector 13 ID-D, a 2.5 222 µm x 3.54 µm full width, half maximum (FWHM) monochromatic X-ray beam tuned to 37 or 42 223 keV was utilized, and diffracted X-rays were collected with a CdTe 1M Pilatus detector. Sample-224 to-detector geometries were calibrated using a LaB_6 NIST standard and a single crystal of 225 enstatite was used for calibration of the rotational geometry of the X-ray beam and detector. 226 Double-sided laser heating and *in-situ* XRD collection took place at Sector 13 ID-D. Fibers 227 lasers shaped with $\sim 10 \,\mu m$ radius flat tops were aligned with the X-ray beam using the X-ray 228 fluorescence of the sample pressure media or the gasket (Prakapenka et al., 2008). During 229 heating, thermal emission from a 6 µm central region of the laser-heated spot was measured 230 spectroradiometrically and fit to a gray body approximation (Heinz and Jeanloz, 1987). A 3% 231 temperature correction was then applied to correct for axial gradients through the sample 232 (Campbell et al., 2007; 2009). The laser power on each side of the sample was adjusted to 233 maintain uniform double-sided heating and X-Ray diffraction and temperature measurements 234 were collected every ~200 K with 1s exposure times. Samples were typically quenched at high 235 temperatures within 30-45 minutes of heating or after a phase transition and suitable grain 236 growth was observed. Powder diffraction patterns were processed using Dioptas (Prescher and 237 Prakapenka, 2015) and CrysalisPro (Rigaku OD, 2018). Pressure was determined using the 238 equation of state of hcp-Fe (Dewaele et al. 2006). 239 Upon quenching, X-ray diffraction maps of the heated spot were collected across a 100

µm² square region in 3 µm steps. Map locations showing high intensity, spotty diffraction
 patterns were chosen for multigrain single-crystal type X-ray diffraction collection approach, as
 these features indicate high-temperature induced crystallite growth. At chosen map locations, X-

243	ray diffraction images were collected across ± 17 to $\pm 30^{\circ}$ rotational scans in 0.25°–0.5° steps with
244	1-4 s exposure times. The diffraction reflections were then mapped in the reciprocal space and
245	target grains were identified and separated from reflections associated with the pressure media,
246	iron, and diamond (Rigaku OD, 2018).
247	Target Fe-S lattices were then indexed and the peak intensities were integrated and reduced
248	using CrysalisPro (Rigaku OD, 2018). Absorption corrections and scaling factors were applied to
249	the structure factors in CrysalisPro using the multi-scan method via the Scale3 Abspack program
250	(Rigaku OD, 2018). The final structure factors and lattice geometries were then refined to known
251	structure models (El-Boragy et al. 1970; Oryshchyn et al. 2011) using SHELXL2014/7
252	(Sheldrick 2015). Reflections showing anomalous calculated versus measured structure factors;
253	likely due to overlap with diamond and other phases in the multigrain sample, resolution
254	limitations, and volume of crystal illuminated by the X-ray beam, were omitted. Structure
255	models were visualized using Vesta during the refinement procedure (Momma and Izumi 2011).
256	

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273 Figures





- a) View of a crystallite of the high-temperature Fe_5S_2 coexisting with recrystallized hcp-Fe after
- synthesis at 159(2) GPa and 3230(130) K (inset). The spotty rings corresponding to the hcp-Fe
- 278 (100) and (101) planes demonstrate that iron equilibrated at 159(2) GPa and 3230(130) K. b)
- 279 View of the reciprocal space for reflections detected in an experiment on the $Fe_{80}S_{20}$ composition
- quenched from 140(2) GPa and 3070(140) K. The reciprocal lattice, colored in black, is
- associated with a Fe_5S_2 grain whose lattice parameters are provided in the top left.



283 Figure 2. Crystal structure of Ni₅As₂-type Fe₅S₂

- 284 Crystal structure of Fe_5S_2 viewed along the a) c axis and the b) (100) plane. c) The Fe-
- coordination polyhedra observed in this atomic arrangement are provided. Fe1, Fe2, Fe4, and
- Fe6 are coordinated by 13 sites and Fe3 and Fe5 are coordinated by 12 sites. Each coordination
- 287 polyhedron consists of Fe and S sites.



b) **C23 Fe₂S** (Co₂P-type, *Pnma*, *Z* = 4)



c) C37 Fe_2S (Co₂Si-type, *Pnma*, Z = 4)



288

289 Figure 3. Crystal structures of Fe₃S and Fe₂S observed in this study

- a) Fe₃S adopts the Fe₃P-type structure (I-4, Z = 8) that is composed of three tetrahedrally
- 291 coordinated Fe-sites, each with increasing Fe–Fe bonding. b) The C23 Fe₂S structure (Co₂P-type,
- 292 Pnma, Z = 4) is made up of columns of FeS₄ tetrahedra and columns of FeS₅ square pyramids
- linked along edges in the *b* direction. c) The C37 Fe₂S structure (Co₂Si-type, *Pnma*, Z = 4) has
- the same site symmetries as the C23 structure, but is marked by a shortened a axis and elongated
- 295 b and c axes accompanied by the formation of a 5-fold dipyramid.





297 Figure 4. Iron-rich sulfide phase diagram to Earth's outer core pressures

At moderate temperatures up to 130 GPa, C23 Fe₂S is observed. At moderate temperatures, tetragonal Fe₃S (Fe₃P-type) is observed below 120 GPa. At moderate temperatures between 130 and 200 GPa, C37 Fe₂S is observed, and on the liquidus between ~115–200 GPa, Fe₅S₂ is the liquidus phase. A kink the Fe-S solidus curve (Mori et al. 2017), is presented to account for the change in phase relations and the high temperatures at which Fe₅S₂ was observed. The closed squares represent experiments conducted on the Fe₈₀S₂₀ starting material, and the open squares represent experiments conducted on the Fe₆₇S₃₃ starting material. Both show compatible results, likely indicating that Fe-rich regions of the Fe₆₇S₃₃ foils were probed.

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475 **Fe₅S₂ identified as a host for sulfur in Earth's core**

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488 Supplementary Information

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489	The Su	pplementary Information includes:
490	\triangleright	Supplementary Information Text
491	\triangleright	Table S1
492	\triangleright	Table S2
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494	\triangleright	Table S4
495	\triangleright	Figure S1
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509 Supplementary Information Text

510 Synthesis of Fe_5S_2 to 200 GPa

511 Fe_5S_2 was synthesized between 119(2) and 193(4) GPa with heating to temperatures 512 between 2400–3300 K. The formation of Fe₅S₂ phase was first recognized in the diffraction 513 patterns by the onset of diffuse streaks and sets of closely spaced reciprocal nodes in the powder 514 diffraction images, suggesting a complex atomic arrangement (Figure 1, S1). With sustained 515 high temperatures in the samples loaded in more hydrostatic pressure media (e.g. KCl vs. SiO₂), 516 large grain growth $(3 - 6 \mu m)$ was observed and the diffuse streaks became less apparent. Further 517 experiments are required to assess the role of hydrostaticity, kinetics, pressure, and temperature 518 on the disorder of this phase.

819 Recrystallization of hcp-Fe with the Fe_5S_2 crystallites at high temperatures was observed 520 in all experiments using the $Fe_{80}S_{20}$ and $Fe_{67}S_{33}$ starting powders as evidenced by the spotty 521 (100) and (101) hcp-Fe rings in Figure 1a. Fe-recrystallization with this high-temperature Fe-522 sulfide establishes that the probed sample locations were in a Fe-saturated phase-field (Figure 523 1a) and this phase is important to consider further in the context of Fe-rich planetary cores.

524 Upon temperature quenching, rotational single-crystal X-ray diffraction scans were performed across a 37–60° omega range, depending on the DAC configuration, and grains were 525 526 indexed in the reciprocal space to a hexagonal unit cell compatible with an Fe_5S_2 volume (Table 527 S1, Figure 1b). Three unit-cell polytypes were observed and indexed in our experiments and are 528 listed in Table S1. All indexed grains exhibit an *a* axial length of ~ 6 Å, while 3 differing *c* axial 529 lengths are observed: ~ 11, 26, and 73 Å. Diffraction mappings show that the polytypism is 530 accompanied by diffuse scattering suggesting positional disorder as the atomic arrangement 531 along the c axis configures at high temperatures (Figure S1). Analogous polytypism has also

been observed in the transition metal binary phase Pb_5As_2 (Saini et al. 1964). Decreased diffuse scattering from the Fe_5S_2 grains was observed after continued heating in the KCl pressure media, and grains with $a \sim 6$ Å, $c \sim 11$ Å were indexed, indicating that this configuration is the most positionally ordered arrangement relevant to these high *P*-*T* conditions. These measured unit cell parameters are compatible with 6 formula units per cell volume of Fe_5S_2 .

537 The structure of a grain Fe_5S_2 synthesized at 140(2) GPa and 3070(180) K was 538 determined based on 159 observed reflections at these extreme conditions (Table S2). 539 Assessment of the systematic absences for the structure factors reduced for Fe_5S_2 suggests a 540 *P*6₃*cm* space group. Structural solution and positional and displacement parameter refinement 541 converged on the Ni₅As₂ structure type (Table S2, 3; Figure 2) (Oryshchyn et al. 2011). The 542 Ni₅As₂ structure is a slight modification of the Pb₅Sb₂ structure (El-Boragy et al. 1970) where the 543 Fe6 site (Table S3) is split about its position and given half occupancy, changing its Wyckoff site 544 from 6c to 12d (Figure 2) (Oryshchyn et al. 2011). This structural modification was initiated 545 based on the large U_{22} parameter observed on the M6 site (M = metal) in Ni₅As₂ when 546 anisotropic displacement parameters were refined using a Pb₅As₂ starting model (Figure S2). In 547 the current dataset, a comparably large U_{22} value was observed on the Fe6 site resulting in an 548 oblong displacement ellipse when refining the data against the initial Pb₅As₂ structure model. 549 The Fe6 site was positionally disordered following Oryshchyn et al. (2011), and the resultant R_1 550 value was improved by around ~5%, rendering the Ni₅As₂ structure model more compatible with 551 the current Fe₅S₂ data (Table S2). Isotropic displacement parameters showing errors > 3σ were 552 fixed to a value equal to approximately the average displacement parameter value for Fe or S 553 (Table S2). The isotropic displacement parameter for the disordered Fe site was also fixed at an 554 average Fe_{Uiso} value (Table S2). The number of reflections collected at these extreme pressures

limits the number of statistically meaningful parameters to refine, and fixing displacement
parameters to reasonable values precludes overinterpretation of the current dataset. The high
isotropic displacement parameter observed on the Fe4 site may represent some vacancies on this
site, but the dataset is again deficient to precisely describe the site occupancy.

559 The experimental details for the solution and refinement of Fe_5S_2 to the Ni₅As₂ solution 560 model are provided in Table S2 and the atomic coordinates of Fe_5S_2 synthesized at 140(2) GPa 561 and 3070(180) K are provided in Table S3. In accordance with previous descriptions of related 562 M_5X_2 transitional-metal binary phases (e.g. (Kjekshus et al. 1973; Oryshchyn et al. 2011), Fe₅S₂ 563 can be viewed as an arrangement of 6 Fe sites and 3 S sites with the Fe1, Fe2, Fe4, and Fe6 sites 564 are in 13-fold coordination, the Fe3 and Fe5 sites are in 12-fold coordination, and the S sites are 565 in 10-fold coordination (Figure 2c). The measured bond lengths for the Fe-coordinated polyhedra 566 are provided in Table S4. All sites are coordinated by both Fe and S sites. A CIF file for Fe_5S_2 is provided in Appendix A1. 567

568 Final R_1 values $\approx 10\%$ attest to the less-than ideal quality of the multigrain dataset as a 569 possible result of the 1-2 megabar synthesis conditions and observed c axial disorder and 570 stacking complexities in Fe_5S_2 (Table S2). Previous studies of isomorphic Ni₅As₂ and related 571 Pb₅As₂ at ambient conditions have also reported similar quality of refinements despite obtaining 572 significantly more reflections in the absence of a DAC (e.g. Saini et al. 1964; El-Boragy et al. 573 1970; Kjekshus et al. 1973). Notable challenges regarding the refinement of the Fe₅S₂ structure 574 model at these extreme conditions are discussed here and compared with similar difficulties 575 reported in previous characterizations of this structure.

576 Thirteen violations of the $P6_3cm$ systematic absence condition: l = 2n for (0kl), were 577 flagged during the refinement of Fe₅S₂. The reflections associated with these systematic absence 578 violations were examined in the raw diffraction images and all show low, diffuse intensity. 579 Discrepancy over the presence or absence of weak reflections with (0kl), l=2n+1 has been 580 reported in previous investigations of Ni₅As₂ and Pb₅As₂ (e.g. Saini et al. 1964; El-Boragy et al. 581 1970; Kjekshus et al. 1973), suggesting that these studies may have also faced difficulties with 582 space group determination. Observations of these low-intensity reflections may be a result of 583 residual disorder in the stacking arrangement along the c direction, and longer heating cycles 584 may be required for the atoms to arrange into equilibrium positions. It is likewise possible that 585 varying synthesis methods for Ni₅As₂ and Pb₅As₂ in previous ambient condition studies affected 586 the c axial atomic arrangement. Ni₅As₂ and isomorphic Ni₅P₂ also exhibits a homogeneity range of ~71.25 – 72.7 atomic % As (Kjekshus et al. 1973; Litasov et al., 2019), and slight 587 588 modifications of this structure based on varying stacking arrangements result in structures such 589 as $Ni_{31}Si_{12}$ (Frank and Schubert 1971) and (Fe, $Ni_{8}(Si, P)_{3}$ (perryite) (Okada et al. 1991). 590 Attempts to refine the current Fe-sulfide phase with the $Ni_{31}Si_{12}$ or perryite structure models did 591 not improve the refinement results. Based on the presence of disorder, polytypism, anisotropic 592 vibrational motion, and nonstoichiometry in the related M_5X_2 phases, additional nuances to the 593 Fe_5S_2 structure model may be developed in future studies; however, the identification and 594 characterization of the Fe₅S₂ crystal structure determined here to 200 GPa is novel, and the 595 observations and challenges reported in this study align with that of previous analyses of Ni₅As₂ 596 and Pb₅Sb₂ that were not limited by microdiffraction in a DAC at extreme conditions.

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598 Other sulfides observed during heating to 200 GPa

599 Three experiments and 7 heating cycles were performed in this study between 100 and
600 200 GPa and to 3400 K, and the X-ray diffraction patterns collected upon heating provide insight

601 into the Fe-rich Fe-S phase relations at outer core pressures and to high temperatures. In each 602 heating experiment, near melting temperatures were targeted and grains of Fe₅S₂ were identified 603 upon quenching. Additional Fe-sulfides were also observed across the thermal gradient of the 604 heated spot, providing information on the lower temperature Fe-saturated phase relations in each 605 heating cycle. Upon quenching from high-temperature synthesis at 119(2) GPa, the other grains in the sample were identified and indexed to a tetragonal cell with a = 8.094(3) Å, c = 3.990(2)606 607 Å, compatible with 8 formula units of Fe₃S (Table S1). The structure was then solved and refined 608 to the Fe₃P-type structure (*I*-4, Z = 8) (referred to herein as *I*-4 Fe₃S) in agreement with previous 609 studies (Fei et al. 2000; Seagle et al. 2006; Morard et al. 2007; Kamada et al. 2010; Kamada et 610 al. 2012; Thompson et al. 2020). The resultant structure model for Fe₃S at these conditions is 611 shown in Figure (3a), a CIF file for Fe₃S is provided in Appendix A2, and the refinement details 612 are given in Table S2. The structure can be viewed as containing 3 tetrahedrally coordinated Fe 613 sites (Blanchard et al. 2008): one Fe-site is coordinated only by S atoms with an average bond 614 length of 2.083(9) Å, another Fe site is coordinated by 3 S atoms and 1 Fe atom with an average 615 bond length of 2.146(8) Å, and the third Fe site is coordinated by 2 S and 2 Fe atoms with an average bond length of 2.137(8) Å (Figure 3). The measured interatomic distances are 616 617 comparable to reports on other transition metal M₃X structures (e.g. Aronnson 1955; Rundqvist 1979). 618

After temperature quenching from 131(2) GPa and 3050(140) K, grains of Fe₃S were not observed across the heated spot. Instead, orthorhombic lattices were also identified with parameters a = 4.869(2) Å, b = 3.256(2) Å, c = 6.139(2) Å, compatible with 4 formula units of Fe₂S. Structural solution and refinement indicate that the Fe₂S grains adopt the *C*23 structure (Co₂P type, *Pnma*, Z = 4) in agreement with previous structural analyses at lower pressures 624 (Zurkowski et al. in press). Refinement details for the structure model are given in Table S2 and 625 a CIF file for C23 Fe₂S is provided in Appendix A3. The structure is composed of columns of 626 FeS₄ tetrahedra and columns of FeS₅ square pyramids linked along edges in the *b* direction 627 (Figure 3b). The average Fe–S bond lengths are 2.213(8) Å and 2.011(2) Å in the square 628 pyramids and tetrahedra, respectively. These values are comparable with that observed in C23 629 Fe₂S at 90 GPa (Zurkowski et al. *in press*).

630 Upon compression and heating at 140(2) GPa and 3070(180) K, orthorhombic grains were identified in the sample chamber with a = 4.667(2) Å, 3.289(1) Å, 6.186(4) Å. This unit 631 632 cell is similar to the C23 Fe₂S observed at 130 GPa, but exhibits a 4 % contraction of the *a* axial 633 length, a 1% extension of the b and c axial lengths, and a 2% volume decrease. Structural 634 solution and refinement using the measured structure factors establish that Fe₂S adopts the C37635 structure (Co₂Si-type, *Pnma*, Z=4) at these conditions (Figure 3c). A CIF file for C37 Fe₂S is 636 provided in Appendix A4 and the analysis details are given in Table S2. In agreement with the 637 proposed C23-C37 Fe₂S phase transition around 140 GPa that is accompanied 1.6% volume 638 decrease (Zurkowski et al., in press), this work confirms that the C23–C37 transition in Fe₂S 639 occurs between 130 and 140 GPa with a similar volume change. Inherent to the C23–C37 640 transition is coordination change from the 4-fold Fe1 coordination polyhedra in the C23 structure 641 to the 5-fold dipyramid polyhedra in the C37 structure (Figure 3b, c). Comparing the C23 and 642 C37 structure models determined at 130 and 140 GPa, respectively, a 10% contraction of the 643 interatomic distance involved in the coordination change is observed (Figure S3). The average Fe–S bond lengths measured in C37 Fe₂S are 2.196(2) Å and 2.165(3) Å in the square pyramids 644 645 and dipyramids, respectively. These values are comparable with C23 Fe₂S.

647 *Fe-rich sulfide phase relations at core-mantle boundary pressures*

648 The presented single-crystal derived structures of Fe₅S₂, Fe₃S, C23 Fe₂S and C37 Fe₂S were then used to inform the changes in diffraction patterns collected during heating between 649 650 110 and 200 GPa and to ~3300 K. Beginning at 112 GPa, diffraction signal from C23 Fe₂S 651 coexisting with Fe was first identified upon heating of the $Fe_{80}S_{20}$ starting material to ~2000 K. 652 With continued heating, peaks associated with tetragonal Fe₃S were observed over a limited 653 temperature range ($\lesssim 2400$ K) until diffuse scattering signal and Bragg reflections from Fe₅S₂ 654 first appeared. Crystallization of the Fe_5S_2 grains with iron occurred with continued heating to 655 119(2) GPa and 2840(180) K. Upon heating beginning at 120 GPa, C23 Fe₂S was observed 656 coexisting with Fe to 122(1) GPa and 2290(120) K, above which the onset of diffraction from 657 the Fe_5S_2 phase was identified and Fe_5S_2 crystallites formed coexisting with hcp-Fe to 131(2) 658 GPa and 3050(140) K. I-4 Fe₃S was not observed. With continued heating cycles between 133(1) 659 GPa and 194(2) GPa and up to 3300 K, C37 Fe₂S coexists with Fe at moderate temperatures and 660 a phase transition to Fe_5S_2 occurs at high temperatures. A phase diagram satisfying these 661 observations is proposed in Figure 4, and integrated XRD patterns for the heating cycles at 662 119(2) and 184(3) GPa and to 3300 K are shown in Figure S4 to demonstrate the identification 663 of C23 Fe₂S, Fe₃S, C37 Fe₂S, and Fe₅S₂ coexisting with hcp-Fe. The peak intensities from the 664 Fe_5S_2 lattices vary during heating up at high temperatures as a result of disorder, polytypism, and 665 the formation of large crystallites at high temperatures (Figure 1, S1). These structural 666 complexities render the powder diffraction patterns difficult to characterize without 667 incorporating single-crystal XRD techniques. 668

669 Discrepancies among the current and previous studies

670 Transitional metal binary compounds with metal-to-nonmetal ratios ranging from 2.33– 671 2.66 (~70–73 atm% metal) predominantly adopt complex trigonal or hexagonal structures with 672 considerable metal-metal bonding (Chen and Whitmire 2018 and references therein). Our 673 observations of the disorder, polytypism, and complex coordination environments inherent to 674 Fe_5S_2 demonstrate that it aligns with this systematic characterization. Interestingly, the Fe_5S_2 675 atomic arrangement is thermodynamically favored at the extreme conditions of Earth's outer 676 core over that of I-4 Fe₃S. These results are contrary to previous reports of Fe₃S stability on to 677 high temperatures in Fe-rich systems to 250 GPa (Kamada et al. 2010; Kamada et al. 2012; Ozawa et al. 2013; Mori et al. 2017). Few lines of reasoning may account for these 678 679 discrepancies. Interpretation of Fe_5S_2 in the integrated powder diffraction patterns is difficult due 680 to the variation in diffraction signal obtained from the Fe_5S_2 polytypes and the low intensity 681 scattering from the disordered sites during its formation. Fe₅S₂ also forms large crystallites, 682 limiting the orientations of the phase and rendering indexation of all diffraction angles for Fe_5S_2 683 quite challenging without rotating the sample. The implementation of single-crystal X-ray diffraction techniques at high pressures was critical in the current study to accurately 684 685 characterize the hexagonal unit cell geometry and complex structure of the Ni₅As₂-like Fe₅S₂. 686 Several studies have also reported chemically analyzed Fe₃S grains in samples recovered 687 from high temperatures in the 200-250 GPa range (Mori et al. 2017; Yokoo et al. 2019; Ozawa 688 et al. 2013), but the difference in Fe content between Fe₃S and Fe₅S₂ may be as few as 3%. This 689 value is generally within 3 sigma of the atomic percent error reported for chemical analyses of 690 samples recovered from these extreme conditions, posing another challenge for distinguishing 691 between the synthesis of Fe_3S and Fe_5S_2 . Furthermore, based on characterizations of the

692 isomorphic Ni₅As₂ and Ni₅P₂, these structures exhibit a homogeneity range with up to ~73 atm% 693 metal (Kjekshus et al. 1973; Litasov et al., 2019). For example, TEM analysis of a Fe–S sample 694 recovered from 236 GPa and 2980 K reveals sulfide grains with on average ~73 atm% Fe 695 (Ozawa et al. 2013). This value is within 0.3-1.25 atomic percent of the possible range of Fe₅S₂ 696 stoichiometries and 2 atomic percent less than an Fe₃S composition. The results from this 697 previous work as well as other chemical analyses of samples recovered from outer core pressures 698 and high temperatures therefore do not contradict the current results (Mori et al. 2017; Yokoo et 699 al. 2019; Ozawa et al. 2013) and may reveal a previous misinterpretation of Fe₅S₂ as Fe₃S based 700 on measured chemistries. 701

- **Table S1.** Unit cell parameters of Fe₅S₂, Fe₃S, and Fe₂S measured upon quenching from high *P*-
- 704 *T* synthesis in this study. Each cell was indexed in the reciprocal space, and the DAC opening,
- and number of reflections obtained for each lattice are listed. For select experiments, high-
- temperature synthesis was conducted without collecting X-ray diffraction data, and the synthesis
- 707 conditions for these experiments are listed as "high T, not measured."

			Room temperature collection 5					Synthes	sis conditions				
	Starting		Phase	Р	a	b		с	v	DAC opening	no. of reflections	Р	Τ
Sample name	material	medium		GPa		Å	Å	Å	Å^3	(°)		GPa	a K
C140_P2_map25	Fe80S20	KCl	Fe5S2	105(2)	6.020(4)			11.204(7)	351.6(5)	60	290	119(2)	2840(140)
C140_P2_map31	Fe80S20	KC1	Fe5S2	105(2)	6.050(2)			73.0(2)	2313(1)	60	134	119(2)	2840(140)
C140_P4_map10	Fe80S20	KCl	Fe5S2	118(1)	5.983(3)			11.078(6)	343.4(4)	60	489	140(2)	3070(180)
C140_P4_map14	Fe80S20	KCl	Fe5S2	118(1)	5.979(2)			11.084(4)	343.2(3)	60	379	140(2)	3070(180)
C140_P4_map14	Fe80S20	KCl	Fe5S2	118(1)	5.973(1)			11.094(2)	342.8(1)	60	276	140(2)	3070(180)
C129_P140_map71	Fe67S33	SiO2	Fe5S2	136.4(5)	5.957(4)			71.89(4)	2209(3)	34	84	159(2)	3230(150)
C129_P150_map9	Fe67S33	SiO2	Fe5S2	144.2(8)	5.939(3)			25.99(6)	794(2)	34	76	high T,	not measured
C129_P150_map9	Fe67S33	SiO2	Fe5S2	144.2(8)	5.923(1)			26.004(6)	789.9(3)	34	146	high T,	not measured
C137_P1_map25	Fe80S20	SiO2	Fe5S2	159(1)	5.890(3)			25.837(7)	776.2(5)	40	108	184(3)	3250(260)
C137_P1_map25	Fe80S20	SiO2	Fe5S2	159(1)	5.880(1)			25.789(3)	772.0(2)	40	108	184(3)	3250(260)
C137_P1_map25	Fe80S20	SiO2	Fe5S2	159(1)	5.876(2)			25.884(8)	774.0(4)	40	116	184(3)	3250(260)
C137_P1_map25	Fe80S20	SiO2	Fe5S2	159(1)	5.883(3)			25.85(1)	774.6(7)	40	126	184(3)	3250(260)
C137_P1	Fe80S20	SiO2	Fe5S2	159(1)	5.897(3)			25.90(1)	775.1(6)	40	116	184(3)	3250(260)
C137_P1	Fe80S20	SiO2	Fe5S2	159(1)	5.878(1)			25.784(3)	771.6(2)	40	149	184(3)	3250(260)
C137_P2_34	Fe80S20	SiO2	Fe5S2	179(1)	5.8400(7)		25.583(2)	755.7(1)	40	189	193(4)	3010(160)
C140_P1_map	Fe80S20	KC1	Fe3S	100(1)	8.156(3)			4.025(3)	267.8(3)	60	416	high T, 1	not measured
C140_P2_map25	Fe80S20	KCl	Fe3S	105(2)	8.094(3)			3.990(2)	261.4(2)	60	296	119(2)	2840(140)
C140_P3_map6	Fe80S20	KC1	C23Fe2S	111(1)	4.869(2)	3.256(2	2)	6.139(2)	97.3(1)	60	227	131(2)	3050(140)
C140_P4_map10	Fe80S20	KC1	C37 Fe2S	118(1)	4.677(2)	3.289(1)	6.186(4)	95.18(9)	60	188	140(2)	3070(180)

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725Table S2. Select experimental details for crystal structure synthesis and analysis of Fe_5S_2 , Fe_3S_3 ,**726**and the Fe_2S polymorphs.

Phase	Fe ₅ S ₂	Fe ₃ S	C23 Fe ₂ S	C37 Fe₂S	
Sample Name	C140_P4_map10	C140_P2_map25	C140_P3_map6	C140_P4_map10	
Synthesis					
Pressure (GPa)	140(2)	119(2)	131(2)	140(2)	
Synthesis Temperature (K)	3070(180)	2840(140)	2050(140)	3070(180)	
Symmetry	Hexagonal, $P6(3)cm$, Z=6	Tetragonal, <i>I</i> -4, Z=8	Orthorhombic, $Pnma, Z = 4$	Orthorhombic, $Pnma, Z = 4$	
<i>a</i> (Å)	5.979(2)	8.094(3)	4.869(3)	4.677(2)	
<i>b</i> (Å)			3.256(4)	3.289(1)	
<i>c</i> (Å)	11.087(6)	3.990(2)	6.139(2)	6.186(4)	
V (Å ³)	343.2(3)	226.4(2)	97.3(1)	95.18(9)	
Reduction					
No. of measured, independent and observed $[I > 2\sigma(I)]$ reflections	396, 309, 157	332, 314, 219	227, 208, 112	188, 158, 105	
$R_{\text{int,}} R_{\text{sigma}}$	0.092, 0.062	0.053, 0.072	0.012, 0.016	0.009, 0.014	
Refinement					
$R[F^2 > 2\sigma(F^2)], wR(F^2), S$	0.098, 0.255, 1.12	0.067, 0.164, 1.05	0.053, 0.149, 1.21	0.047, 0.139, 1.22	
No. of reflections	156	219	112	105	
No. of parameters	21	17	10	10	
$\Delta \rho_{max}, \Delta \rho_{min} (e \text{ Å}^{-3})$	2.14, -2.18	1.92, -1.68	1.86, -2.02	1.91, -2.21	

Table S3. Atomic coordinates of the Fe₅S₂ refinement model for the data collected at 140(2) GPa
and quenched from 3070(180) K.

Wycoff					
site	ATOM	X	У	Z	Uiso
2a	Fe1	0	0	0.963	0.021
	error			0.003	0.005
4 b	Fe2	1/3	2/3	0.089	0.046
	error			0.003	0.005
6c	Fe3	0.259	0	0.126	0.021
	error	0.002		0.001	
6c	Fe4	0.613	0	0.222	0.021
	error	0.002		0.001	0.003
6c	Fe5	0.284	0	0.330	0.027
	error	0.002		0.001	0.003
12d	Fe6*	0.651	0.056	0.426	0.021
	error	0.003	0.003	0.002	
2a	S1	0	0	0.220	0.026
	error			0.004	
4 b	S2	1/3	2/3	0.299	0.026
	error			0.002	
6c	S 3	0.673	0	0.030	0.026
	error	0.005		0.003	

*indicates half occupancy

748	Fable S4. Selected interatomic distances for the Fe sites measured in Fe ₅ S ₂ at 140 GPa.
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Atom	bonded to	#	distance (Å)
Fe1	-Fe3	3	2.39(2)
	–Fe5	3	2.22(3)
	-Fe6	3	2.313(17)
	- S 1	1	2.66(8)
	S 4	3	2.09(3)
Fe2	-Fe3	3	2.275(3)
	-Fe4	3	2.375(16)
	-Fe6	3	2.40(3)
		1	2.36(4)
	S 4	3	2.108(13)
Fe3	-Fe1	1	2.39(2)
	-Fe2	2	2.275(8)
	-Fe4	3	2.350(15)
	-Fe5	1	2.30(3)
	-Fe6	1	2.307(17)
	-S1	1	1.90(4)
	-S4	3	2.68(3)
Fe4	-Fe2	2	2.375(16)
	-Fe3	3	2.320(11)
	-Fe5	3	2.416(14)
	-Fe6	1	2.303(17)
	- S 1	1	2.326(11)
		2	2.044(16)
	-S4	1	2.10(3)
Fe5	-Fe1	1	2.22(3)
	-Fe3	1	2.30(3)
	-Fe4	3	2.416(14)
	-Fe6	3	2.135(15)
	- S 1	1	2.09(5)
	- S 3	2	2.184(10)
	S4	1	2.27(3)
Fe6	-Fe1	1	2.313(17)
	-Fe2	2	2.40(3)
	-Fe3	1	2.307(17)
	-Fe4	1	2.303(17)
	-Fe5	3	2.135(15)
	-S3	2	2.15(3)
	S4	3	2.17(3)

- **Figure S1.** Unwarped diffraction mappings of the (1*kl*), (0*kl*), and (-1*kl*) directions for an Fe₅S₂
- rts4 crystallite exemplifying the diffuse scattering signal collected at 119(2) and 2840(180) K. The
- (1kl) and (-1kl) mappings also show the presence of a polytype with more closely spaced
- reflections oriented $\sim 51^{\circ}$ from the mapped crystallite.



Figure S2. Fe₅As₂ structure model shown with anisotropic displacement ellipsoids using the
starting Pb₅As₂ structure model. In accordance with the Ni₅As₂ structural characterization
(Oryshchyn et al. 2011), these sites were disordered about their position and ascribed half
occupancy.



Figure S3. Comparison of the C23 and C37 structure models refined at 130 and 140 GPa,
respectively. In the C23 Fe₂S structure, the next nearest sulfur site to the Fe1 tetrahedral site is at
a 2.963(1) Å distance at 130 GPa (dotted line). A 10% contraction of this interatomic distance
and a coordination change is observed in the formation of C37 Fe₂S at 140 GPa (right).





- 811 Figure S4. X-ray diffraction patterns collected upon heating in an Fe₈₀S₂₀ starting composition at
- a) 119(2) GPa and b)184(3) GPa. The miller indices for C23 Fe₂S (blue), Fe₃S (burgundy), and
- 813 *C*37 Fe₂S (green) are provided, and the red "H" symbols represent the observations of the
- 814 formation of Fe₅S₂. The growth of large crystallites, along with the disorder and polytypism of
- 815 this phase make for challenging powder diffraction indexing, but the lattice parameters of Fe_5S_2
- 816 indexed in the reciprocal space after quenching from these high *P*-*T* conditions are provided.



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820 Supplementary Information References

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