# Discovery of a 3.46 billion-year-old impact crater in Western Australia

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#### Abstract

Current models for the origin and constituents of Earth are based primarily on compositions of meteorites that fell recently (within the past ~1 million years) on Earth. Here we report the discovery of a probable crater (~150x100x3 km in the original dimension), which was possibly created by the impact of a large (~10 km in diameter) asteroid on a >2.5 km-deep, oxygenated ocean-floor in Pilbara, Western Australia 3.46 billion-years (Ga) ago; that is, more than 1.2 Gyr earlier than the oldest known impact crater on Earth. This discovery was made mostly through a drone-assisted survey of the distributions of impact breccias and tsunami deposits. We have also discovered numerous micro-fragments ( $<1 \mu m$  to  $\tilde{-3}$  mm in sizes) of the asteroid (named here as "The Apex Asteroid") in tsunami deposits and submarine pillow lavas that were most likely generated by the asteroid impact. The micro-fragments are mostly comprised of titanite (CaTiSiO5) and rutile (TiO2), and frequently associated with native iron (Fe), carbon (C)-rich nanocrystals (e.g., SiC, graphite), coesite (high-pressure polymorph of SiO2) and a variety of alloys, such as iron-iridium (Ir) alloys with up to ~10 wt% Ir, iron-nickel (Ni)-cobalt (Co)-, aluminum (Al)-copper (Cu)-, Al-Si-, Si-C-, Fe-C- and C-Al-Ca-Cu-Fe alloys. The Apex Asteroid was much richer in titanium (Ti), calcium (Ca) and vanadium (V), and poorer in magnesium (Mg) and Fe compared to recently-fallen meteorites. Thermochemical analyses of the asteroid minerals suggest that: (1) the parental planetary body of the Apex Asteroid condensed from a solar-gas cluster that was hotter and having higher H2/H2O and H/O ratios compared to planetary bodies for the recently-fallen meteorites; (2) it may represent a major building-block of Earth; and (3) Earth and the Moon have the same chemical and isotopic compositions, but Ti atoms in the Earth's magma oceans partitioned into the Ti-rich core and Ti-poor mantle, while most Ti atoms in the lunar magma oceans condensed as the Ti-rich mantle.

Table S6. Calculated temperatures of appearances and disappearances of solid phases during cooling of the Apex solar-gas nebula. Initial  $\sum P = 10^4$  atm.

Compound	Name	Appearance T (K)	Disappearance T (K)	
TiC	Titanium carbide	1,830	812	
Ti <sub>5</sub> Si <sub>3</sub>	Titanium silicide	1.733	1.570	
TiO	Titanium monoxide	1.660	1.397	
V <sub>2</sub> C	Divanadium carbide	1.550	1.294	
Ca4Ti3O10	Yuksporite	1 397	1 315	
Ca <sub>2</sub> Al <sub>2</sub> SiO7	Gehlenite	1,378	992	
Ca <sub>2</sub> SiO <sub>4</sub>	Larnite	1,320	1,121	
NiSi	Nickel silicide	1,320	984	
FeSi	Ferro silicide	1,294	968	
CoSi	Cobalt silicide	1,290	949	
VC <sub>0.88</sub>	Vanadium carbide	1,287	690	
Cr <sub>5</sub> Si <sub>3</sub>	Chrome silicide	1,260	1,055	
SiC	Silicon carbide	1,240	987	
CaSiO <sub>3</sub>	Wollastonite	1,160	861	
Ca <sub>3</sub> MgSiO <sub>4</sub>	Merwinite	1,080	1,021	
Cr <sub>3</sub> C <sub>2</sub>	Chrome carbide	1,055	720	
C	Graphite	1,039	730	
CaMgSiO <sub>3</sub>	Monticellite	1,021	1,002	
MgAl <sub>2</sub> O <sub>4</sub>	Spinel	1,002		
CaAl <sub>2</sub> Si <sub>2</sub> O <sub>8</sub>	Anorthite	992	780	
NaAlSi <sub>3</sub> O <sub>8</sub>	Albite	987		
CaMgSiO <sub>3</sub>	Diopside	987	835	
Ni <sub>5</sub> P <sub>2</sub>	Pentanickel diphosphide	984	954	
Fe <sub>3</sub> P	Iron phosphide	979	960	
Cu	Native copper	979		
Fe	Native iron	969	780	
Co <sub>2</sub> P	Cobalt phosphide	960	436	
Ni <sub>3</sub> P <sub>2</sub>	Nickel phosphide	954		
Co	Native cobalt	949		
Ca <sub>3</sub> Al <sub>2</sub> Si <sub>3</sub> O <sub>12</sub>	Grossular	911	854	
MnSi	Manganese silicide	875	855	
CaTiSiO <sub>5</sub>	Titanite	873		
KAlSi <sub>3</sub> O <sub>8</sub>	Sanidine	845	447	
MgSiO <sub>3</sub>	Enstatite	835		
Ti <sub>2</sub> O <sub>3</sub>	Tistarite	832	795	
Ti <sub>4</sub> O <sub>7</sub>	Titanium suboxide	802	730	
V2O3	Vanadium trioxide	790		
SiO <sub>2</sub>	Quartz	780		
TiO <sub>2</sub>	Rutile	733		
MgCr <sub>2</sub> O <sub>4</sub>	Magnesiochromite	606		
FeCr <sub>2</sub> O <sub>4</sub>	Chromite	606		

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18	Current models for the origin and constituents of Earth are based primarily on
19	compositions of meteorites that fell recently (within the past ~1 million years) on Earth.
20	Here we report the discovery of a probable crater (~150x100x3 km in the original
21	dimension), which was possibly created by the impact of a large (~10 km in diameter)
22	asteroid on a >2.5 km-deep, oxygenated ocean-floor in Pilbara, Western Australia 3.46
23	billion-years (Ga) ago; that is, more than 1.2 Gyr earlier than the oldest known impact
24	crater on Earth. This discovery was made mostly through a drone-assisted survey of the

25	distributions of impact breccias and tsunami deposits. We have also discovered numerous
26	micro-fragments (<1 $\mu$ m to ~3 mm in sizes) of the asteroid (named here as "The Apex
27	Asteroid") in tsunami deposits and submarine pillow lavas that were most likely generated
28	by the asteroid impact. The micro-fragments are mostly comprised of titanite (CaTiSiO <sub>5</sub> )
29	and rutile (TiO <sub>2</sub> ), and frequently associated with native iron (Fe), carbon (C)-rich
30	nanocrystals (e.g., SiC, graphite), coesite (high-pressure polymorph of SiO <sub>2</sub> ) and a variety
31	of alloys, such as iron-iridium (Ir) alloys with up to ~10 wt% Ir, iron-nickel (Ni)-cobalt
32	(Co)-, aluminum (Al)-copper (Cu)-, Al-Si-, Si-C-, Fe-C- and C-Al-Ca-Cu-Fe alloys. The
33	Apex Asteroid was much richer in titanium (Ti), calcium (Ca) and vanadium (V), and
34	poorer in magnesium (Mg) and Fe compared to recently-fallen meteorites.
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36	body of the Apex Asteroid condensed from a solar-gas cluster that was hotter and having
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38	meteorites; (2) it may represent a major building-block of Earth; and (3) Earth and the
39	Moon have the same chemical and isotopic compositions, but Ti atoms in the Earth's
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Current models for the origin and constituents of Earth postulate that its core is composed
primarily of Fe-Ni alloys with minor Si and S, such as found in Fe- and/or stony-Fe meteorites;
whereas the mantle resembles the compositions of stony meteorites, such as CI-carbonaceous
chondrites<sup>1,2</sup>, ordinary chondrites<sup>3</sup> or enstatite chondrites<sup>4</sup>, which are composed primarily of MgFe-rich and Ti-Ca-poor silicates. These meteorites fell mostly within the last ~1 million year

48 (Myr) [5], and referred to here "modern meteorites". However, questions remain as to whether 49 the Earth's building blocks ~4.56 Ga ago are truly represented by "modern meteorites" [6,7]. For 50 example, recognizing the differences in some elemental ratios (e.g., Al/Si, Mg/Si, Fe/Mg) and 51 isotopic ratios of elements (e.g., O, Re and Nd) between the estimated "primitive Earth mantle" and various chondrites, Drake and Righter<sup>6</sup> have suggested "The 'building blocks' of the Earth 52 53 must instead be composed of unsampled 'Earth chondrite' or 'Earth achondrite'. The oldest 54 meteorite falls are the 2.7 Gyr-old iron micro-meteorites discovered in carbonate beds in Pilbara, 55 Australia [8]. The oldest-terrestrial-aged stony meteorites, recovered from ~466 Ma-old 56 limestone beds in Sweden and China, are found to be much significantly richer in Ti (i.e.,  $\sim 2$ 57 wt% vs. <0.3 wt%), and have distinctly different abundance ratios of various types of chondrites 58 and achondrites than any "modern meteorites" [9,10].

Impact history of the Moon<sup>11</sup> suggests frequent bombardments of asteroids on early Earth.
Evidence of asteroid impacts on early Earth, such as impact spherules (i.e., terrestrial rocks
vaporized by asteroid impacts and condensed as melts) and Ir-rich sedimentary rocks, has been
found in several sedimentary-rock formations ~3.47-3.25 Ga and ~2.7-2.5 Ga in ages [12-16].
However, no stony-meteorite fragments older than ~466 Ma [9,10] or impact crater older than
~2.2 Ga in ages [17,18] has been found on Earth.

In 2003, the NASA Astrobiology Program carried out a series of drillings in the Eastern
Pilbara district of Western Australia (Fig. 1) to recover some of the oldest geologic formations
for investigations aimed at understanding the evolution of life and environment on early Earth.
The first drilling (ABDP #1)<sup>19</sup> recovered a 264 m-long continuous drill core that encompasses
the youngest section of the Duffer Formation (submarine felsic volcanic formation of ~3.46 Ga
in age), a ~150 m section of the Marble Bar Chert Member (MBCM, chert/jasper unit) and a ~70

71 m earliest-section of the Apex Basalt (submarine komatiite/basalt pillow lavas) (Fig. 1d & Fig. 2a). At the 2017 Goldschmidt Conference, we<sup>20</sup> reported the discovery of numerous (>1,000) 72 73 mineral fragments (<1  $\mu$ m to ~3 mm in sizes), which we interpreted as fragments of a meteorite 74 (or asteroid), throughout the  $\sim$ 70 m section of the Apex Basalt in the ABDP #1 core. These 75 "meteoritic" mineral fragments are comprised mostly of titanite (CaTiSiO<sub>5</sub>) and rutile (TiO<sub>2</sub>). 76 Our report, however, met with some skeptics, because: (i) such Ti-rich minerals are extremely rare in "modern meteorites", (ii) meteoritic minerals have not been reported in a submarine lava 77 of any geologic age, and (iii) titanite and rutile are common hydrothermal-alteration products of 78 79 submarine basalts.



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Subsequently, we have continued various types of investigations of the "meteoritic grains" and their geological and geochemical environments (see "*Methods*") and concluded that the semi-circular structure, presently ~80x65 km (but originally ~150x100 km) in size and encompassing the ~3 km-thick Apex Basalt and the Mt. Edgar Granite Batholith (Figs. 1-a & -c), likely represents a crater created by impact of a large (~10 km in diameter) asteroid, which produced the "meteoritic" fragments discovered in the Apex Basalt (Figs. 2-a & -b). Below we present various types of observations and data that have lead us to the above conclusions.



#### 98

**Figure 2. History of the Apex Impact Crater.** (See A in *Methods*). (a). A stratigraphic column of the ABDP #1 drill core. (b). The sequence of major events in the impact crater. (I): Pre-impact sage. (II): An impact of a ~10 kmsized asteroid on the deep ocean floor, causing: brecciation and fragmentation of the asteroid, the MBCM and the Duffer Formation; generation of tsunami currents and deposition of tsunami deposits. (III): Mantle impact melting and eruption of the Apex Basalt magmas. (IV): Emplacement of the Mt. Edgar granite batholith and eruptions of the Panorama volcanic rocks. (c): Impact melting of an oceanic crust and the mantle by a 30 km-sized bolide (modified after Jones<sup>23</sup>).

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## 107 Impact breccias and tsunami deposits

108 The MBCM is a ~20-200 m-thick, steeply dipping and slightly overturned chert/jasper unit 109 that outcrops continuously  $\sim 20$  km along a strike in the western part of the proposed Apex 110 Impact Crater (Figs. 1 b-c; Fig. 3a). Previous researchers recognized an abundance of highly-111 unsorted "chaotic" sedimentary rocks that contained angular fragments of the MBCM and 112 occurring between the well-bedded MBCM and overlying Apex Basalt. These features were 113 described as "turbidites interbedded with debris-flow deposits" by DiMarco & Lowe<sup>24</sup>, or as 114 "mixed conglomerate beds, chert breccia beds and chert folded beds of the MBCM" and 115 interpreted as the product of high-density turbidity currents and mass-transport complexes in

- 116 channels-levees in deep-sea fans by Olivier et al.<sup>25</sup>. However, the virtual absence of round
- 117 fragments of the MBCM in the "chaotic zone" is difficult to explain by these detrital-sediments
- 118 models.



120 Figure 3. Asteroid impact structures in the Marble Bar Chert Member (MBCM). (a): An aerial view (looking 121 south from ~250 m above the ground) of the MBCM in a ~100x50 m area at the Marble Bar Pool. (b): A close-up 122 view of area (b) ( $\sim$ 50x40 m area) in Fig. (a). Note the abundance of fractures (up to  $\sim$ 5 m wide) that are oblique to 123 the bedding planes, various-sized (up to  $\sim 10$  m) broken and rotated blocks and angular fragments of the MBCM, and 124 gray/black "muds" (i.e., tsunami sediments) that intruded into the fractures and gaps between the MBCM fragments. 125 (c): An outcrop photo of the impact-induced tsunami deposits, containing  $\sim 1$  cm to  $\sim 5$  m-sized fragments of the 126 MBCM. (d): A boulder of tsunami deposits, containing angular fragments of red-, black- and white cherts in the 127 matrix comprised mostly of powdered MBCM and the Duffer Formation.



- 130 MBCM around the Marble Bar Pool (Figs. 1 b-c), revealed an abundance of brecciations,
- 131 displacements, deformations, disturbances and impact metamorphism of the MBCM and the
- 132 Duffer Formation from  $<1 \mu m$  to  $\sim1 km$  in scales (Figs. 1b; Figs. 3 a-d; Extended Data (ED)
- 133 Figs. 1 a-k; ED Figs. 2 a-k). An entire section of the ~20-200 m-thick MBCM was broken into

various-sized, sharp-edged fragments and blocks, which were displaced by up to ~10 m-wide
fractures that cross-cut the MBCM bedding planes at high angles. Gaps between the blocks and
fragments were filled with black/gray-coloured "muds" that contained various-sized angular
fragments, pulverized chert and -volcanic rocks (from the Duffer Formation), and "inclusions"
and "intrusions" of basalts up to ~2 m-thick and ~10 m-long. Soft-sediment deformation features
of the bedded MBCM are common.

140 Through detailed mapping of a ~40x20 m area of the MBCM at the Marble Bar Pool, Oliver 141 & Cawood<sup>26</sup> recognized essentially the same features described above and interpreted them as 142 products of tectonic activity and the dewatering of partially-consolidated MBCM. However, we 143 interpret that aforementioned structural and textural features in the MBCM represent impact 144 breccias (in-site- and fallback breccias) that were created by shock waves from an asteroid 145 impact, because of the abundance of shock-metamorphic features in the MBCM and Duffer 146 Formation (ED Figs. 2 a-k) and of meteoritic minerals in the "mixed zone" (i.e., tsunami 147 deposits) and the Apex Basalt (ED Figs. 3 a-t; see later section). Pillow basalts with inclusions of 148 chert fragments are common (ED Figs. 1 b-k; ED Figs. 2 b-e). These features indicate that a 149 bolide impacted before the consolidation of the upper parts of the MBCM, and that eruptions of 150 the first >50 m of the ~3 km-thick Apex basalt began essentially concurrent to the depositions of 151 fragmented-, crashed- and powdered cherts. In areas in the eastern parts of the proposed Apex 152 Impact Crater where the MBCM did not form, boulders and fragments of the volcanic Duffer 153 Formation are mixed with powdered volcanic rocks in the tsunami deposits (ED Fig. 1k).

## 154 Meteoritic minerals in pillow basalts and tsunami deposits

155 *Titanite and rutile*: The most abundant "meteoritic" grains (>95 % of the total "foreign"

156 grains) in the Apex Basalt and tsunami deposits are Ca-Ti-Si-O-rich phases (referred to here as

157 "Ti-rich grains") that are comprised mostly of titanite and rutile (Figs. 4 a-h; ED Figs. 3 a-t). 158 These grains are virtually free of Fe (Supplementary Table 2), indicating that they were not 159 formed by alteration of Ti-rich iron-oxides by Ca-rich hydrothermal solutions (see reaction R-1 160 in *Method*), such as those found in some submarine basalts<sup>27, 28</sup>. Titanite crystals and basalts in the hydrothermally-altered zone of the ABDP #1 core are characterized by losses of essentially 161 162 all the Ca atoms by submarine hydrothermal fluids (ED Figs. 4 b, i, l, & m) (see reaction R-2 in 163 Method), indicating that the Ti-rich grains are exotic minerals (e.g., meteorite fragments) which 164 were incorporated in the pillow basalts and tsunami deposits before submarine hydrothermal 165 alteration.



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Figure 4. Meteoritic minerals in the Apex Basalt and tsunami deposits. (a): A microscope photo under reflected 168 light of the first meteoritic mineral fragment discovered in the Apex Basalt. Due to the reactions with oxygenated 169 submarined hydrothermal fluids, rutile (TiO<sub>2</sub>)-rich lamellae and haematitized (brown-coloured) surfaces have 170 became distinct. (b): A thin section photo (under reflected light) of a drill-core chip of the Apex Basalt, showing 171 abundant Ti-rich asteroid fragments (khaki colour) in the older basalt flows, but virtually absent in the younger 172 flows. (c): A composite EMPA maps of Ti (red; asteroid fragments) and Al (yellow; basalt) in area (c) in Fig. (b).

fragment with bent crystal structure, which was possibly created by a planetary collision. (f): A HAADF image of 175 the bent crystal lattice shown in (e). (g): An EPMA map of Ti in a Ti-rich fragment, showing the titanite-rich 176 matrices (blue), titanite- and rutile-rich lamellae (dark green) and rutile crystals (yellow & red). (h): An EPMA map 177 of Ti in Ti-rich melt spherules. (i): A thin section photo (under reflected light) of melted Ti-rich spherules. (j)-(n): 178 STEM photos of meteoritic minerals associated with Ti-rich fragments.

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180 The Ti-rich grains (Figs. 4 a-i; ED Figs. 3 a-t; Figs. g-j) are characterized by: (a). angular, 181 irregular and ragged shapes and the virtual absence of rounded grains, eliminating the possibility 182 of a detrital origin; (b). virtual absence of reaction rims; (c). large variation in the grain size (<1 183  $\mu$ m to ~3 mm); (d). heterogeneous distributions in the basalt lavas (Fig. 4b): highly abundant (~1 184 to ~5 volume %) in older basalt melts that crystallized probably in contact with seawater, but 185 virtually absent in younger lavas that infiltrated the older basalts and probably not in contact with 186 seawater; (e). secondary fragmentation of the grains, possibly by ablation by other crystals, 187 during incorporation/transportation in basalt lavas, as suggested by the jigsaw puzzle-like 188 matching of neighboring fragments (ED Fig. 3b); and (f), association with the minerals that have 189 been found in common chondrites and achondrites, such as iridium (Ir)-rich minerals, Fe-Ni-Co 190 alloys, Al-Si alloys, silicon carbide (SiC) and various forms of carbon (e.g., graphene, carbon 191 nanotubes) (Figs. 4 j-n; ED. Figs. 7-9). Characteristics (a)-(f) indicate that the Ti-rich grains did 192 not crystalize either as primary- or secondary (alteration) minerals of the basalt lavas or tsunami 193 deposits, but they were incorporated in the basalt lavas and tsunami deposits as foreign materials 194 from the ocean water. Because the solubilities of Ti-bearing minerals in aqueous solutions are 195 extremely low<sup>29</sup>, the Ti-rich compounds in the seawater must have existed as solid 196 suspensions/deposits (i.e., asteroid fragments). Characteristics (a)-(f) can be best explained by 197 the fragmentation of an outer zone of an asteroid body as it impacted to a deep-ocean floor. The 198 asteroid fragments, together with vaporized terrestrial rocks and chert fragments, were ejected 199 from the impact site on an ocean floor into the overlying ocean and atmosphere as impact clouds,

200 and then fell back to the ocean (see Fig. 2b). The Stokes' Law suggests that it would take ~4 201 days, ~400 days, and ~100 years, respectively, for titanite grains of 100  $\mu$ m, 10  $\mu$ m, and 1  $\mu$ m in 202 diameters to sink through a ~5 km-column of calm ocean water. However, tsunami waves would 203 have mixed and stirred them, especially inside the impact crater, to deposit widely heterogenous 204 Ti-rich grains and tsunami deposits. The tsunami waves within a large impact crater may have 205 lasted for more than several days [30], and eruptions of the Apex Basalt possibly began within a 206 few hours of the impact (see A-III in Method), therefore explaining why both the tsunami 207 deposits and at least the first 50 m section of the Apex Basalt contain various-sized Ti-rich grains 208 (Figs. 4 a-c; ED Figs. 3 a-t). It should also be noted that the maximum stability temperatures of titanite + rutile assemblage (>  $\sim$ 1,100 °C at  $\sim$ 1 atm)<sup>31</sup> are higher than the basalt lava temperatures 209 210  $(\sim 1,000 - 1,150^{\circ}C)^{32}$ , explaining why most of the titanite-rutile grains were not melted in the 211 basalt lavas. But some Ti-rich grains have melted to form Ti-rich spherules (Figs. 4 h-i; ED Figs. 212 5 a-d), possibly by a planetary collision of the parental body with another planet or by the heat in 213 the impact clouds.

214 The Ti-rich angular fragments are also characterized by: (g). exsolution textures with rutile 215  $(TiO_2)$ -rich lamellae and titanite (CaTiSiO<sub>5</sub>)-rich matrices (i.e., between lamellae) (Fig. 4g; ED 216 Figs. 3 a-t); (h). aggregates of nanometer-sized crystals of titanite and rutile that comprise the 217 lamella and matrices (Fig. 4g; Extended Data Figs. 3 f-t); and (i). distorted crystal lattice 218 structures (Figs. 4 e-f; ED Figs. 3 c-e). These features suggest the following sequence of events 219 for the parental planetary body of the Apex Asteroid: (1) condensation of nano-sized crystals of 220 Ti-rich crystals from the solar gases during the formation of the parental planet; (2) 221 reorganization of the nano-crystals of Ti-rich minerals to form larger (>10 µm in size) crystals 222 with rutile-rich lamellae and titanite-rich matrices during cooling of the parental body; and (3)

deformation of crystal lattice of the Ti-rich crystals by a collision of the parental body with
another planetary body. The Ti-rich spherules with rutile-rich lamellae and titanite-rich matrices
(Fig. 4 h-j; ED Figs. 5 a-d) were possibly formed by melting of angular Ti-rich fragments,
followed by slow cooling to develop the exsolution textures while in impact clouds. The basalticspherules (ED Figs. 5 e-h) and wiggled-shaped silica (ED Figs. 5 i-k) were probably condensed
in the impact cloud from vaporized volcanic rocks of the Duffer Formation and chert of the
MBCM, respectively.

Analyses of some rutile crystals by EELS (electron energy loss spectroscopy) indicate the presence of both  $Ti^{3+}$  and  $Ti^{4+}$  (ED Figs. 6 a-c)<sup>33</sup>; some  $Ti^{3+}$  atoms may have been  $Ti^{2+}$  before entering in the Earth's atmosphere (see later "Discussions").

233 Metallic Fe and Fe-oxides: Metallic iron and Fe-oxide crystals are intimately associated 234 with the Ti-rich crystals in the pillow basalts (Fig. 4 j-k). Fe-oxides are also found as discrete 235 grains in the tsunami deposits (ED Figs. 7 l-p). They vary from euhedral crystals (hexagonal, 236 octahedral) to irregular shaped crystals. Their Fe/O atomic ratios (ED Figs. 7 l-p; ED Figs. 8 i-j; 237 Supplementary Table S3) suggest that the original grains were metallic Fe (Fe/O atomic ratios 238 >>1) and possibly wüstite (Fe/O = 0.95-0.88) as well. Some grains, especially at the margins, 239 exhibit lower Fe/O ratios (e.g., ~0.75 for magnetite and 0.67 for haematite; Supplementary 240 Table S3), possibly due to partial oxidation while in the high-temperature impact clouds in the 241 O<sub>2</sub>-rich ocean and atmosphere and/or by O<sub>2</sub>-rich submarine hydrothermal fluids. The abundant 242 fragments of red cherts (Fig. 3d; ED Fig. 7k) in the tsunami deposits are unequivocal evidence 243 that the formation of the primary hydrothermal haematite crystals in the MBCM took place in the 244 O<sub>2</sub>-rich deep oceans prior to the asteroid impact at ~3.46 Ga, rather than by recent ground water 245 (see A-I in Methods).

It has been debated whether the native iron and Fe-C alloys in Tertiary-aged basalt lavas in the Disko Island are meteorite origin or products of the assimilation of C-rich rocks (e.g., coal), causing the Fe<sup>2+</sup>O component in basalt melts to be reduced by carbon [34, 35]. However, C-rich sedimentary rocks (e.g., black shales) are absent in pre-3.5 Ga geologic formations in the Pilbara district, eliminating the possibility of a terrestrial origin for the Fe-metal and wüstite in the Apex Basalt.

*Iridium-rich minerals*: As many meteorites are Ir-rich, researchers<sup>12-16</sup> have used Ir-rich 252 253 sedimentary layers as evidence for the contemporaneous impacts of asteroids (or large 254 meteorites) somewhere on Earth. In the tsunami deposits and pillow basalts, Ir is enriched in 255 three groups of meteoritic minerals: Ir-Fe-(Ni) alloys, Al- and REE (rare earth element)-enriched 256 phosphates, and zircon (see Table S4 in Supplementary Information). Their Ir contents vary from 257 <0.05 % (i.e., detection limit) to ~10 wt.% (Fig. 5a; ED Figs. 8 c-k), which matches the highest 258 Ir content previously reported for meteorites (i.e., inclusions in the Ornance carbonaceous 259 chondrites<sup>36</sup>).

Other alloys: Submicron-sized crystals of Ni-Cr-Co alloys, spherules of Al-Cu with
exsolution textures (Fig. 4l), Al-Si alloys (Fig. 4m) and C-Al-Ca-Si-Cu-Fe alloys (ED Figs. 9 ko) are found in the pillow basalts. The Al-rich alloys are similar to those discovered in the
recently-fallen Khatyrka meteorite<sup>37, 38</sup> and those in the "Hypatia"<sup>39</sup>.

*Carbon-rich phases:* The C-rich phases associated with Ti-rich grains in the pillow basalts occur in various forms (Fig. 4n; ED Figs. 9 a-o): graphite, graphene, carbon nanostructures (nano tubes, nano spherules, nano whisker) and silicon carbide (SiC) crystals, which often contain 1-2 nanometer-sized native metal- or metal-oxide particles. These C-rich phases have been found in a variety of meteorites [40-44]. Synthesis of carbon nanostructures and silicon carbide are

269 commonly carried out in laboratories via a variety of vapor deposition methods, in which C-rich
270 gas (e.g., CO, hydrocarbons) is reduced by metal catalysts (e.g., Fe, Ni, Co, Pt) [45, 46].

271 Therefore, the question remains as to whether some (or all) of the C-rich phases were condensed

272 in the Solar Nebula or re-condensed from the vaporized carbonaceous matter of the asteroid in

impact clouds.

*Coesite:* Silica-rich micron-sized crystals in some basalt samples (Fig. 4j) were identified by
electron diffraction method as coesite, a high-pressure (>2 GP) polymorph of SiO<sub>2</sub> [47]. Because
of its intimate association with meteoritic minerals (e.g., TiO<sub>2</sub>, native Fe and Fe-Ni-Cr-Co
alloys), this coesite was possibly produced by the shock metamorphism of quartz, crystobalite or
tridymite in the parental planetary body upon collision with another planetary body or upon
impact of the asteroid to Earth.





Figure 5. Chemical compositions of the Apex Asteroid. (a): Iridium contents of some meteoritic grains in the Apex Basalt, compared to other meteorites. (b): Comparisons of the elemental abundances (normalized to Si) in the Apex Asteroid with those in CI-chondrite, H-enstatite chondrite and the primitive mantle [48]. (c): Ti/Si and Ca/Si ratios of the Apex Asteroid are compared with those of CAI minerals (perovskites, hibonites, fossaites and pyroxenes), carbonaceous chondrites, H-enstatite chondrites, Earth mantle and lunar basalts [48-51], indicating that the Apex Asteroids and lunar basalts have much higher Ti/Si ratios compared to the other materials.

289 The asteroid fragments discovered in the Apex Basalt and tsunami deposits are all micro 290 fragments (<1 µm - 3 mm in size). It is possible that many other minerals beside the Ti-rich 291 minerals, such as pyroxenes, olivines and feldspars, had existed in the Apex Asteroid, but these 292 mineral fragments had been vaporized in the impact clouds and/or melted in the Apex Basalt 293 lavas, if their melting temperatures were less than ~1,100°C. However, because the abundances 294 of other phases (e.g., carbon and metal compounds) are less than ~1 volume % of the Ti-rich 295 grains and of the absence of identifiable other meteoritic mineral phases, we have assumed that 296 the bulk composition of the Apex Asteroid may be approximated by that of the least-altered Ti-297 rich phases by observing. Based on EPMA analyses of 219 spots on Ti-rich grains 298 (Supplementary Table S2) and the estimated abundance ratio of  $(85\pm5)/(15\pm5)$  for the 299 matrix/lamellae, we have estimated the average compositions of the bulk Ti-rich grains (Table 1) 300 as  $(Ca_{.74\pm.18}, Fe_{0.03\pm0.03})$   $(Ti_{1.27\pm.17}, Al_{0.05\pm0.05}, V_{0.02\pm0.01}, Si_{0.81\pm.18})O_5$ , if we assume that all the Ti 301 atoms are Ti<sup>4+</sup>. Compared to all the stony meteorites investigated previously, the Apex Asteroid 302 is significantly enriched (>10 times) in Ti, Ca and V, significantly depleted (<1/10) in Mg, P, S, 303 Fe and Ni, and moderately to slightly depleted ( $\sim 1/10$  to  $\sim 9/10$ ) in Cr and Mn, when normalized 304 to Si (Fig. 5b). The Ti/Si - Ca/Si relationships of the Apex Asteroid resemble those of calcium-305 aluminum-rich inclusions (CAIs) (Fig. 5c). However, the Ti/Si ratios of the Apex Asteroid and 306 lunar basalts are much higher than those of the CAIs and "modern" meteorites, suggesting the 307 genetic relationships between the Apex Asteroid and the Moon (see later discussions).



309 Figure 6. Thermochemical analyses of formation conditions of the Apex Asteroid (See C-7 in Methods). (a): 310 Temperatures of the appearances and disappearances of minerals from "the Apex solar-gas cluster" with the initial 311 total gas pressure of  $10^{-4}$  atm (see Supplementary Table 6). The red and pink lines represent Ti-rich minerals; black and gray lines for C- and/or Fe-rich minerals; blue lines for silicides; yellow and khaki lines for Ca-Al-rich minerals; 312 313 green lines for Mg-rich silicates; and purple lines for phosphides. (b): Changes in the abundances of major gaseous 314 species in the Apex solar-gas cluster during the formations of phases in (a). (c): Stability relationships among some 315 redox-sensitive minerals as a function of H<sub>2</sub>/H<sub>2</sub>O ratio of the coexisting vapor phase. The blue line is for the Apex 316 solar-gas cluster from (b); the gray area represents the redox-temperature conditions for the "modern" meteorites. 317

- 318 Assuming that ~4.6 Ga ago, the Solar Nebula had the same elemental ratios as the current
- 319 Solar Wind and CI-carbonaceous chondrites, some researchers, such as Grossman<sup>52</sup> and
- 320 Lodders<sup>1</sup>, compute the condensation temperatures of meteoritic minerals by using

thermodynamic data. By adopting a similar thermochemical approach, we have computed the condensation-, transformation- and disappearance temperatures of solid phases and the abundance ratios of gaseous species during the cooling of a hypothetical solar-gas cluster that has the elemental ratios of the Apex Asteroid (Table 1), and the same H/C/Si ratios of the current Solar Photosphere and the initial total vapor pressure of 10<sup>-4</sup> atm as Lodders<sup>1</sup> (see C-7 in *Methods*).

327 The results of our computations (Figs. 6a-c; Supplementary Table 6) differ significantly from 328 those by Lodders<sup>1</sup>, including: (1). The first major minerals to condense were the CAI minerals at 329 1,680-1,520K in Lodders<sup>1</sup>, but Ti-carbides and -silicides at ~1,800K in this study. (2) Ti-bearing 330 minerals in Lodders' are all oxidized (Ti<sup>4+</sup>) compounds, but our calculations show that he first 331 Ti-rich minerals condensed Ti<sup>0</sup>-rich minerals, which successively transformed to compounds with higher valences,  $Ti^{2+}$ -rich,  $Ti^{3+}$ -and finally to  $Ti^{4+}$ -rich minerals at ~900-700K. (3). In our 332 333 system, C-bearing solid phases initially condensed as carbides (e.g., TiC, FeC, SiC) at ~1830-334 1250K, which were transformed to C (graphite) at ~1040 K and decomposed to CH<sub>4</sub> (methane) 335 at 730K. In contrast, no C-bearing solid phases condensed in Lodders system, except for 336  $CH_4$  7H<sub>2</sub>O (methane hydrate) at <400K. (4) Many native metals (and their alloys), such as Fe, Cu 337 and Co, mostly condensed at ~980-900K. These data agree well with the observed occurrences 338 of reduced compounds, such as graphite, carbon nanocrystals, FeC, Fe-rich alloys (Fe-Ir, Fe-Cu-Co-Ni) and  $Ti^{3+}$  in the Apex Asteroid. (5). Phosphorus (P) condensed as phosphides (P<sup>0</sup>) of 339 340 heavy metals (e.g., FeP, Ni<sub>3</sub>P<sub>2</sub>), rather than as phosphates ( $P^{5+}$ ). (6) H<sub>2</sub>/H<sub>2</sub>O ratios of the Apex solar-gas cluster remained nearly constant at  $10^{7\pm1}$  at ~2000-800 K. In contrast, H<sub>2</sub>/H<sub>2</sub>O ratios of 341 342 the current Solar Photosphere and the gas clusters in equilibrium with the "modern" meteorites would be  $\sim 10^{-2} - 10^{+6}$ , because the Fe in these meteorites occurs predominantly as Fe<sup>2+</sup>-rich 343

minerals (e.g.,  $Fe_2SiO_4$ ,  $FeSiO_3$ ) as well as native Fe, and Ti occurs essentially as  $Ti^{4+}$ ; that is, these H<sub>2</sub>/H<sub>2</sub>O values are much lower (i.e., more oxidized) that those of the Apex solar-gas cluster (Fig. 6c). The major reason for such large differences in the results of computations stems from the difference in the estimated abundance of oxygen atoms in the systems: the H/O atomic ratio of 4,300 in our system vs. 1,750 in Lodders<sup>1</sup>.

349 Most models for the origins of planets in the Solar system are based on an unspoken 350 assumption that the compositions of solar gases have remained the same as the current Solar 351 Winds. However, Barcall et al. [53] have estimated that at 4.57 Ga ago the Solar radius was 352  $\sim$ 87% of today, luminosity  $\sim$ 70%, temperature  $\sim$ 97%, hydrogen mass fraction  $\sim$ 50% and 353 neutrino fluxes <10% of today. These estimates suggest that during the planet formations ~4.6 354 Ga ago elemental and isotopic compositions of the clusters of gases emitted from the early Sun 355 varied greatly from the present values. That is, the parental planetesimal for the Apex asteroid 356 (termed here as "Planetesimal A") condensed from solar-gas clusters with different chemical and 357 isotopic compositions from those for the "modern meteorites". The proto Earth was possibly 358 created by the accumulations of various planetesimals, not by a single type of planetesimal. For 359 example, a 50% mixture of Planetesimals A and the parental planetesimal for the enstatite 360 chondrites would have yielded the Earth's bulk compositions as Ti = -10 wt%, Ca = -8 wt%, Mg = -5 wt%, Fe = -15 wt%, Ni = -0.8 wt%, Si = -8 wt%, Al = -0.5 wt%, S = -3 wt%, Si = -2 wt%, Si = -2361 362 15 wt% and  $O = \sim 33$  wt%. Impacts of other planetesimals to Planetesimal A during the Earth 363 formation would have ejected fragments of Planetesimal A into space; one of which returned to 364 Earth 3.46 Ga ago as the Apex Asteroid.

The current popular model for Earth and Moon formation postulates that the heat generated from the conversion of kinetic energy from planetesimal accumulation and from the radioactive

367 decay of short-lived isotopes caused the entire Earth to melt and form magma oceans, and that 368 the impact of Mars-size Theia ejected some of this magma ocean into space to form the Moon at 4.51 Ga [54]. Recently, Lock et al.<sup>55</sup> have proposed that the collision of Theia and the proto-369 370 Earth would have vaporized both planets and created a fast-spinning vaporized state (termed 371 "synestia") and that magma oceans formed on Earth and the Moon as the synestia cooled. Both 372 models for Moon's formation imply that the bulk compositions of Earth and the Moon are the 373 same, yet the present mantles differ significantly, most likely due to the partitions of some 374 elements into their respective cores. The changes in the mantle compositions would have been 375 more significant for Earth, because the mass fraction of its core is about 32% [48], compared to 376 less than 2% for the Moon [56, 57]. Under the reducing conditions of the initial magma oceans, Ti atoms could have existed as  $Ti^0$ ,  $Ti^{2+}$  and  $Ti^{3+}$ as well as  $Ti^{4+}$  (see Fig. 6c). They would have 377 378 reacted with Fe<sup>0</sup> in the magma oceans to form titanium-iron alloys and wüstite (FeO), such as:

379 
$$TiO_{2(m)} + 3Fe^{0}_{(m)} = TiFe_{(m)} + 2FeO_{(m)}$$
 (1)

380 Similarly, SiO<sub>2</sub> in the magma oceans would have formed FeSi and FeO by:

381 
$$\operatorname{SiO}_{2(m)} + 3\operatorname{Fe}^{0}_{(m)} = \operatorname{FeSi}_{(m)} + 2\operatorname{FeO}_{(m)}$$
 (2)

382 where subscript *m* refers to melt. The Ti-Fe and Si-Fe alloys would have sunk to the bottom of 383 the magma oceans to create a metallic core, while the FeO<sub>(m)</sub> concentrated in the mantle, reacted 384 mostly with SiO<sub>2(m)</sub> to form ferrous silicates. As a result, the remaining magma oceans on the 385 early Earth would have become Ti-poor, FeO-rich, and its redox state (H<sub>2</sub>/H<sub>2</sub>O) became below 386 that of the Fe/FeO buffer line. In contrast, the lunar mantle remained Ti-rich, retained near the initial Fe<sup>0</sup>/Fe<sup>2+</sup> ratio at about the Fe/FeO buffer line. This model explains why the lunar basalts 387 generally contain much higher amounts of Ti than terrestrial basalts (average 3 wt% vs. ~1 388 389 wt%)<sup>56,57</sup>. It also suggests that the reason the Earth's core has a lower density than pure Fe metal may not be due to the additions of H, Si, O and/or S to the core, as suggested in the current
 paradigm<sup>58</sup>, but likely due to the significant additions of Ti and Si.

392 Similar to the impact history of the Moon<sup>10</sup>, asteroid impacts on early Earth must have been

393 much more frequent than during the Proterozoic and Phanerozoic Eons. This study suggests why

impact craters older than ~2.2 Ga have not been found on Earth. Throughout the geologic

395 history, most of the impact craters must have been created on deep-ocean floors rather than on

396 land. Most of them subducted into the mantle, but some may have been thrusted up to the

397 continental crust before being subjected to erosion. Therefore, many more Archaean-aged impact

398 craters should be found in greenstone/granite terranes, such as those in the Pilbara Craton (Fig.

399 1a).

401 Table 1. EPMA analyses of Ti-rich grains in the Apex Basalt402

Element	Matrix		Lamellae		Bulk	
	(n = 39)	2σ	(n = 21)	2σ	wt. %	2σ
	wt.%		wt. %			
0	38.59	3.65	38.67	2.17	38.60	3.3
Na	0.12	0.51			0.1	0.5
Mg	0.57	1.14			0.48	0.9
Al	1.29	2.09			1.09	
Si	11.97	2.16	2.40	2.15	10.53	2.1
Р	0.01	0.01			0.01	
S	0.01	0.01			0.01	
K	0.01	0.02			0.01	
Ca	16.14	4.27			13.71	3.6
Ti	22.87	6.67	57.27	3.68	28.03	6.2
V	0.35	0.21			0.30	0.2
Cr	0.01	0.01	0.53	0.29	0.08	0.1
Mn	0.02	0.04			0.02	
Fe	1.39	1.78	0.65	0.25	1.3	
Со	0.10	0.01			0.1	
Ni	0.01	0.02			0.01	
Cu	0.05	0.02			.04	
Zn	0.01	0.02			0.01	
Total	93.51	3.47	99.52		94.4	

- 403 n: The number of spots analyzed, excluding the data on more than 100 spots
- 404 with mixed mineralogy and/or those altered by submarine hydrothermal fluids).
- 405
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540 Methods

541

## 542 A. Methods for unravelling the geologic history of the Apex Impact Crater

543 Glikson et al. [15] reported the discovery of two zones of impact-spherule-rich layers in the 544 Marble Bar Chert Member (MBCM) of the Duffer Formation at 57-58 m and 77 m depths in the 545 ABDP #1 drill core. Each zone contains multiple spherule-rich layers that are ~1-5 cm in 546 thickness and packed with nearly uniform-sized (1-2 mm diameters) spherules with Mg-Fe-Ca-547 rich core (possibly basaltic in composition) and secondary siderite-rich rims. The spherule-rich 548 layers are interbedded with diamictite layers, which were most likely formed by impact-549 generated tsunami currents and comprised of variable-sized (<1 mm to  $\sim10$  cm) angular chert 550 fragments. These impact spherules were correlated to the 3,472-3,470 Myr-old impact spherule 551 units in the Barberton Greenstone Belt in South Africa. Because of the uniformity in the spherule 552 sizes and the absence of accompanying meteorite fragments, these small spherules and tsunami 553 deposits were probably generated by asteroid impacts at >500 km away, much like those found 554 in the Lake Superior region from the 1.8 Ga Sudbury Impact [59, 60]. 555 In contrast, fragments of "meteoritic minerals" discovered by us in a ~70 m section (~190 to 556  $\sim$ 264 m depth interval) of the Apex Basalt in the ABDP #1 drill core greatly vary in size (<1  $\mu$ m 557 to  $\sim$  3mm), shape and abundance (Figs. 4 a-i; ED Figs. 3 a-t). The associated diamictite zone is 558 tens of meter thick, containing large fragments (up to ~10 m in size) (Figs. 3 a-d; ED Figs. 1 a-559 k). These observations have lead us to hypothesize that: the Apex Asteroid impacted to the ocean 560 floor near (within ~100 km of) the ABDP #1 site during the accumulation of the MBCM; and the 561 semi-circular igneous structure, presently ~80 km E-W and ~65 km N-S in size and centered at S 562 21°10' and E 120°0' (Fig. 1c), may represent the asteroid impact crater, termed here "the Apex 563 Impact Crater". This is one of the eight greenstone-granite complexes that characterize the 564 geology of the Eastern Pilbara Craton (Fig. 1a). As the rock formations in the "Apex Impact Crater" are generally ~60°-80° tilted (overturned), mostly due to the deformation associated with 565 566 the intrusion of granite batholiths, the initial dimensions were much larger, possibly ~150 km (E-

567 W) and ~100 km (N-S). Below are the proposed sequence of events in the Apex Impact Crater
568 (see Fig. 2b).

569 A-I: Geological and geochemical environments before the asteroid impact: Submarine 570 volcanic rocks of the Warrawoona Group, which includes the North Star Basalt-, McPhee 571 Basalt-, Mt. Ad Basalt-, and Duffer Formations (Fig. 1d), are more than 13 km in thickness and 572 accumulated without a subaerial-erosional surface during a ~55 Myr period (3,515-3,460 Ma 573 ago) [21]. Because of its great thickness and long period of submarine volcanism, we suggest 574 that the Warrawoona Group, comprised mostly of basalt and komatiite lavas and minor 575 dacite/rhyolite lavas, represented an oceanic crust that formed on a mid ocean ridge. The 576 occurrences of felsic igneous rocks (rhyolite and dacite) in mid ocean ridges were possibly 577 similar to those found on Iceland today; they may have formed by fractional crystallization of 578 basaltic magmas and/or by the partial melting of hydrothermally-altered oceanic crust [61]. The 579 occurrences of Cu-rich volcanogenic massive sulfide deposits (VMS; the Lenners Find- and the 580 Big Stubby deposits; see Fig. 1c) in the upper part of the Duffer Formation suggest that the 581 seawater depths were greater than ~2.5 km (i.e.,  $P_{H2O}$  > ~250 bars), because Cu-rich submarine 582 hydrothermal fluids, which are typically 300-400 °C in temperatures, must discharge on the 583 seafloor without boiling in order to form Cu-rich massive sulfide ores [62]. In fact, Samieyani<sup>63</sup> 584 obtained homogenization temperatures of 250-390 °C for fluid inclusions with no evidence of 585 boiling in barite and quartz crystals from the Big Stubby deposits near Marble Bar. 586 The MBCM (~20-200 m thick) is a low-grade Algoma-type Banded Iron Formation, 587 characterized by alternating microbands of white-coloured silica-rich chert, red-coloured haematite-rich jasper, and black-coloured magnetite-rich jasper<sup>19</sup>. Based on detailed 588 589 mineralogical and geochemical investigations utilizing a variety of electron microscopes, Hoashi

590 et al.<sup>19</sup> have recognized that the primary haematite crystals are typically submicron-sized ( $\sim 0.1$ -

591 0.6  $\mu$ m) euhedral crystals (ED. Figs. 7 g-j), which most likely formed by rapid mixing of Fe<sup>2+</sup>-

592 rich hydrothermal fluids with O<sub>2</sub>-rich cold deep-ocean water at  $T \ge 60^{\circ}C$  on or near the seafloor:

593 
$$2Fe^{2+}_{(hyd. fl.)} + 1/2O_{2(sw)} + 2H_2O \rightarrow Fe_2O_{3(hm)} + 4H^+$$
 (R-1)

If the ferric-iron oxides had formed at temperatures below ~60°C, they would have been goethite 594 (Fe(OH)<sub>3</sub>), rather than haematite<sup>19</sup>. Based on these observations, they have concluded that the 595 596 ocean-atmosphere system was already fully oxygenated ~3.5 Ga ago. However, based on an 597 interpretation of Pb isotope data on some basalt samples from the ABDP #1 core, Li et al.<sup>64</sup> 598 concluded that the additions of uranium (U) and O<sub>2</sub> to the Apex Basalt occurred by modern O<sub>2</sub>-599 rich groundwater, not by the 3.46 Ga seawater, implying that the haematite crystals did not form 600 3.46 Ga ago. Focusing on the larger (>10  $\mu$ m), subhedral secondary haematite crystals in the MBCM, Rasmussen et al.<sup>65</sup> also suggested that all haematite crystals, including the submicron-601 602 sized haematite crystals, formed though the oxidation of Fe<sup>2+</sup>-rich minerals, such as siderite 603 (FeCO<sub>3</sub>) and magnetite (Fe<sub>3</sub>O<sub>4</sub>) by recent groundwater. However, they have neglected the photographic evidence, presented by Hoashi et al.,<sup>19</sup> showing that siderite and magnetite crystals 604 605 in the haematite-rich red bands of the MBCM clearly grew from the submicron-sized primary 606 haematite crystals (ED Figs. 7 i-j), which were subsequently converted to large secondary 607 haematite crystals. Furthermore, various geochemical characteristics of the Apex Basalt in the 608 hydrothermally-altered zone, such as the negative Ce anomalies, the increased contents of Ba, U and  $Fe^{3+}$  and  $Fe^{3+}/\Sigma Fe$  and U/Th ratios (ED Figs. 7 b-f), indicate that the 3.46 Ga submarine 609 610 hydrothermal fluids developed from the  $O_2$ -,  $SO_4^2$ - and U rich seawater [66]. 611 A-II. The asteroid impact and associated events: The Apex Basalt, which overlies the

612 MBCM, is ~3 km-thick (Fig. 1d), comprised of submarine basalt/komatiite lavas with abundant

613 pillow structures (ED Fig. 1i). Recognizing that the MBCM in the studied drill core is highly 614 fractured, brecciated, and interfingering with basalts, especially in the upper section (i.e., deeper 615 in the drill core), we suggest (see Fig. 2b) that: (i) an asteroid body impacted to the ocean floor 616 near (within ~100 km of) the ABDP #1 site during the accumulation of the MBCM, and created 617 a large impact crater; (ii) shock waves generated by the impact shattered, brecciated, dislocated, 618 locally overturned the chert blocks, and generated large tsunami currents, especially in areas 619 inside the crater, which formed thick (~5-30 m) tsunami deposits; (iii) upon impact, the asteroid 620 fragmented into various-sized pieces, which were blown into the overlying ocean and 621 atmosphere, and subsequently fell back (i.e., fallback breccia) on the tsunami deposits and the 622 pillow basalts, primarily inside of the impact crater. The geometry of the impact crater was most 623 likely defined by the outcrop-areas and thickness (~3 km) of the Apex Basalt. Therefore, the 624 original impact crater was a semi-circular in shape,  $\sim$ 150 km (E-W) and  $\sim$ 100 km (N-S) in size, 625  $\sim$ 3 km deep from the surrounding ocean floor and  $\sim$ 5-6 km deep from the sea level. Size of the impactor, according to an equation for the ratio of crater/impactor sizes<sup>67</sup>, would have been  $\sim 10$ 626 627 km in diameter.

628 **A-III. Generation and eruptions of the Apex Basalt magmas:** The proposed Apex Impact Crater is about a half the size of the Ontong Java Plateau. Some researchers<sup>23,68</sup> have suggested 629 630 that the Ontong Java Plateau, the world's largest oceanic plateau, was created by magmas that 631 were generated by the impact of a  $\sim$ 30 km-diameter asteroid to a deep ocean floor  $\sim$ 120 Ma ago. 632 According to the simulations by Jones<sup>23</sup>, the impact of a ~30-km diameter asteroid created a 633 crater ~400 km in diameter, and caused mantle melting to depths of ~150 km almost 634 instantaneously (i.e.,  $< \sim 10$  minutes) by impact heating of the oceanic crust and by sudden 635 decompression melting of deeper mantle (Figs. 2 b-c). The degree of melting of the mantle

peridotite would have decreased with increasing depth: from ~100 % at <~20 km depths, >50%
at <~70 km depths and >1% at <~150 km depths. Melts formed from ~100 % melting of</li>
peridotite would be komatiitic<sup>69</sup>, while those from ~50 % and ~5% partial melting would
produce basaltic magmas and granitic magmas, respectively [70]. These depths of melting would
be shallower when the impact asteroids were smaller.

641 Analogous to the ascent speeds of kimberlite magmas<sup>71</sup>, the komatiite/basalt melts generated 642 by impact melting of the mantle could have risen with the speeds of  $\sim 10-30$  m/sec through the 643 mantle fractures created by the impact. Thus, within one hour of the impact, komatiite/basalt 644 lavas would have erupted into the impact crater to form the Apex Basalt. According to the simulations<sup>72</sup>, tsunami currents generated by impact of a ~10 km-diameter asteroid would have 645 646 risen more than a few kilometers in heights and lasted for more than a day, especially within the 647 impact crater. Such a prolonged tsunami event and the rapid eruptions of basalt/komatiite lavas 648 would explain the intimate associations of basalts and chert fragments in the tsunami deposits 649 and in the early phases of the Apex Basalt (ED Figs. 2 b-e). Since the Apex Asteroid impacted to 650 a deep ocean floor, and the "impact plumes" were possibly confined mostly within the ocean 651 (rather than dispersed into the atmosphere), large tsunami currents would have stirred fragments 652 of various sizes and densities in the water column inside the impact crater. This would explain 653 the extremely large size-variation in asteroid fragments in the Apex Basalt and tsunami deposits. Volume of the Apex Basalt lavas in the proposed Apex Impact Crater would be  $\sim 5 \times 10^5$  km<sup>3</sup>, 654 655 assuming an original dimension of ~150 km diameter and ~3 km depth; this is approximately the 656 same volume as a typical LIP (large igneous province) (> $10^5$  km<sup>3</sup>)[73].

A-IV. Generation and emplacements of the granitoid magmas: A large granite body
 typically occurs in the central parts of impact craters on Earth, such as the 2.0 Ga Vredfefort<sup>17</sup>

659 and ~60 Ma Chixulub Crater<sup>74</sup>. These granite bodies may represent the rebound of the lower 660 continental crust, as these asteroids impacted on a thick continental crust. Granitoid batholiths in 661 the Apex Impact Crater (i.e., the Mt. Edgar Granite Batholiths) are, however, similar to those in 662 the 1.8 Ga Sudbury Impact Crater<sup>75</sup>, as they intruded as granitic magmas through the earlier 663 mafic/ultramafic igneous bodies. Crystallization ages of the granite complex in "the Apex 664 Crater" range from ~3,448 to ~3,426 Ma [21], indicating that the first granite intrusive occurred 665 at  $\sim 10$  Myr after the impact event at  $\sim 3,460$  Ma, and that the granite intrusive events lasted for 666 ~20 Myr. Similarities in the ages and compositions between the Mt. Edgar granite complex and the Panorama Formations<sup>76</sup> suggest that they were co-magmatic; the Panorama Formation (felsic 667 668 volcanics) represents the eruptive phase of the granitic magmas that formed the Mt. Edgar 669 granite complex. We estimate that the felsic volcanic rocks (i.e., the Panorama Formation) were 670 accumulating at least 5 km in thickness on top of the Mt. Edgar granite complex (i.e., the 671 lithostatic pressures of >1.5 kb) in order for granite magmas to crystallize without acquiring 672 porphyritic textures<sup>77</sup>. Thus, toward the end of the accumulation of the thick Panorama 673 Formation, the entire eastern part of the East Pilbara Terrane, from the Mt. Edgar area to the 674 North Pole Dome area (~100 km W of Marble Bar) would have become subaerial at ~3,430 Ma. 675 Regional erosion continued for ~80 Myr until the region became under a shallow sea and 676 deposited the Strelley Pool Chert at ~3,350 Ma [66]. 677 The questions of whether the continental crust existed beneath the Warrawoona Group

igneous rocks and the origin of the granitic magmas for the Mt. Edgar Batholith have been
debated. Some researchers<sup>21,78-80</sup> have suggested the thick submarine volcanic rocks of the
Warrawoona Group accumulated on an older sialic continental crust, implying that the large
granitoid magmas that formed the Mt. Edgar batholith and the volcanic Panorama Formation

682 were produced by the melting/assimilation of the older continental crust by mantle-derived 683 basaltic magmas. However, the available isotopic- and chemical data on zircons and igneous 684 rocks in the Pilbara district [78-80] are inconclusive for the origin of the granitoid-magmas in the 685 Pilbara district. If the granitoid magmas had been generated via either by the fractional 686 crystallization of basaltic magmas or by the melting/assimilation of continental crust, we would 687 expect the volumes of granitic magmas to have been much smaller than those of basaltic 688 magmas. However, the volume of the Mt. Edgar granitoids and the Panorama volcanics are too 689 large to have been produced by the Apex Basalt magmas, either by fractional crystallization or 690 by assimilation of the continental crust. A more likely scenario for the origin of the large felsic 691 magmas in the Apex Impact Crater would have been by 1-50 % partial melting of the deep 692 mantle peridotites by the asteroid impact (see Fig. 2c).

## 693 B. Origin of the Ti-rich fragments in the Apex Basalt

Titanite and rutile are common low-temperature alteration minerals in submarine basalts [27, 28], where Ti-rich iron-oxides, such as ilmenite (FeTiO<sub>3</sub>) and titaniferrous magnetite, reacted with  $Ca^{2+}$ -rich hydrothermal fluids, such as:

 $3FeTiO_{3(s)} + 3SiO_{2(aq)} + 3Ca^{2+} + 3H_2O + 1/2O_{2(aq)} = 3CaTiSiO_5 + Fe_3O_{4(s)} + 6H^+ (R-2)$ Such reactions produce magnetite and/or haematite as well as titanite. Submarine hydrothermal fluids are products of seawater-rock interactions as seawater circulate through the underlying rocks that are heated by an intrusive igneous body [62]. Temperatures of hydrothermal fluids that discharge onto the seafloor near a heat source generally increase with time from ~50 °C to the maximum of ~200-450 °C, depending on the initial temperature and size of the intrusive, and then decrease (i.e., waning) to <50 °C. Low-temperature (T< ~100°C) submarine hydrothermal fluids are typically enriched in Ca compared to the seawater [62], which promotes formation oftitanite via R-2.

In contrast, high-temperature submarine hydrothermal fluids are typically poor in Ca [62].
Therefore, titanite does not form through R-2. Instead titanite would lose its Ca and form rutile
via:

$$CaTiSiO_{5(titanite)} + 2H^{+} = TiO_{2(rutile)} + Ca^{2+} + SiO_{2(aq,s)} + H_2O$$
(R-3)

The behaviors of Ca and Fe in titanite crystals and their host rocks in the ABDP #1 core (ED

Figs. 5 a-m), especially in the chert/basalt mixed zone (~190–215 m depth zone), clearly indicate

that the Ti-rich grains in the Apex Basalt and tsunami deposits were not produced by reaction R-

2, but they were incorporated in the tsunami deposits and basalts as foreign grains, most likely as

714 meteoritic minerals, and subsequently lost Ca via reaction R-3.

## 715 C. Approaches

To examine the validity of the above suggestions (A) and (B), we have carried out the following investigations: (i) field investigations of the impact structures and tsunami deposits; (ii) microscopic investigations of impact textures (e.g., shock-metamorphic features) in the MBCM and Duffer Formation; (iii) mineralogical and geochemical investigations of "meteoritic minerals" in the tsunami deposits, as well as in the Apex Basalt; (iv) geochemical investigations of the tsunami deposits and hydrothermally-altered Apex Basalt; and (v) thermochemical analyses of the stability and abundances of solid- and gaseous phases from solar-gas clusters.

C-1. *Field investigations*: Since Yr. 1999, we have carried out eleven field
excursions/surveys to study the geology of the East Pilbara Craton, focusing specifically on
banded iron formations, volcanogenic massive sulfide deposits, paleosols and the Marble Bar
Chert/Jasper. During the field work in the Marble Bar area in 2018, we utilized a drone (DJI
Mavic Pro) equipped with a high-resolution (4k) camera to investigate the distributions and
structures of highly fragmented MBCM and the impact-induced tsunami deposits.

729 C-2. *Petrography and optical microscopy*: Optical microscopy lends itself to characterize
 730 Ti-Ca-Si-O-rich meteorite fragments under a microscope, especially their khaki-colour and

731 angular shapes. Therefore, it is not difficult to recognize the Ti-rich fragments on the surfaces of 732 the drill core using a hand lens or a binocular microscope. We have examined the entire  $\sim 265$  m-733 long ABDP #1 core for the presence/absence of meteorite fragments and, confirmed the 734 ubiquitous presence of Ti-rich fragments throughout the Apex Basalt section, from ~190 to ~265 735 m depths in the core section. The occurrences of meteoritic minerals in the selected 54 core chips 736 were photographed through optical microscopes under normal light at The Pennsylvania State 737 University (PSU). Morphologies of meteoritic grains in approximately 30 specimens (core chips, 738 polished sections, thin sections) were further examined under high-resolution microscopes by 739 Drs. Janine Diez and Walter Engel of the GRL, Inc. in Bern, Switzerland.

740 C-3. *Bulk-rock chemical analyses*: The abundances of the major-, trace-, REE-, and 741 precious-elements in 70 chips of the volcanic rocks (basalt, komatiite, dacite) and 142 chips of 742 the Marble Bar Chert were analyzed by the Activation Analysis Ltd., Ontario, Canada, and by 743 the Mineral Constitution Laboratory of the Pennsylvania State University (see the data in 744 Supplementary Table 1). The bulk-rock chemical analyses have identified the zones of tsunami 745 deposits and hydrothermal alteration (ED. Figs. 5 a-f; 7 a-f).

C-4. Chemical analyses by X-ray fluorescence spectroscopy: Spot analyses for the
concentrations of O, Mg, Al, Si, K, Cr, S, Ca, Ti, V, Mn, Fe, Ni, Cu, Zn, Co, Ga, Ge, Ir and Pt
were carried out on 21 spots on of un-polished core surfaces of Sample 257.1m, 29
spots on 213.10m, and 54 spots on 236.5m using an X-ray fluorescence spectroscopy
(XGT-5000 by Horiba, Inc.) at the PSU Geosciences Dept with a beam size of 10 µm. In
addition, elemental distribution maps of 6 areas (ranging from 1x1 to 5x5 mm in sizes) were
carried out using a 100 µm-size beam (see the data in Supplementary Table 2).

753 C-5. Chemical analyses by electron microprobe analyzers (EPMA): WDS analyses were 754 performed with a CAMECA SX-Five electron microprobe at the PSU Materials Characterization 755 Laboratory. The beam was regulated at 15kV 30nA with a spot size of 1 µm. A total of 580 756 spots on 10 samples were collected: 12 spots on Sample 259.95, 189 spots on 257.1, 41 spots on 757 236.6, and 40 spots on 214.43. Additionally, 55 spots on 259.95, 10 spots on 569, 220 spots on 758 214.43, 2 spots on 213.74, 3 spots on 213.3, 4 spots on 211.22, and 4 spots on 211 were analyzed 759 using a field emission scanning electron microscopy (FESEM, JSM-7001F) at Tohoku 760 University. In addition, approximately 30 elemental maps (ranging from 10x10 µm to 5x5 mm in

- areal sizes) were constructed on the above samples. Samples for the EPMA analyses at PSU
- 762 were first coated with iridium to determine the contents of C, as well as O, Na, Mg, Al, Si, P,
- 763 S, K, Ca, Ti, V, Cr, Mn, Fe, Ni, Cu, and Zn. Subsequently, samples 259.95 and 214.43
- 764 were re-polished, re-coated with carbon, and analyzed for Ag, Au, Pt, Ir and Pb,
- as well as the above major and trace elements. Samples for the FESEM analyses at
- Tohoku were coated with carbon (see the data in Supplementary Table 3).

767 **C-6.** *High resolution electron microscopy*: Sample powders for HRTEM (high-resolution 768 TEM) analyses were isolated and collected from microscopy thin sections, sample blocks and from selected rock chips using a needle to lift material grains, crystal assemblages and lamellae 769 770 under a 8x-50x track stand stereo zoom parfocal trinocular microscope. The sample 771 powders were dispersed on 200 mesh Formvar/carbon coated copper or nickel grids and 772 examined in analytical microscopy at the University of Kentucky and NIOSH (JEOL 2100F 773 field emission transmission and scanning transmission microscope (TEM/STEM using an 774 analytical probe with 0.17 nm resolution) at 200 kV with Gatan Ultrascan 4k x 4k CCD camera. 775 Gatan Digital Micrograph software, Digiscan II, Gatan HAADF (high Angle Annular Dark 776 Filed) detector and Gatan 2000 Image Filter and at Wright Patterson Research Lab using a FEI 777 Titan Aberration-corrected (S)TEM. Energy dispersive spectroscopy (EDS) and electron energy 778 loss spectroscopy (EELS) analysis (Oxford Aztec EDS system, was used for elemental mapping 779 and EELS spot analysis and trace lines were obtained with a 1 nm probe, alpha of 12 mrad and 780 beta of 6 mrad; elemental intensity maps were obtained from core edge intensity acquired after 781 background subtraction using an integration window ranging from 10-30 eV. The oxidation state for Ti in select grains was determined by EELS and compared to the Ti<sup>3+</sup>/Ti<sup>4+</sup> ratio of synthetic 782 783 standard samples (the valence state of Ti was measured by means of EELS using a 200 keV 784 beam intensity and a Gatan PEELS 666 parallel-recording electron spectrometer (calibration 785 technique used intensity lines and spectral features of the L3.2 white lines of Ti (with a resolution of 0.75 eV) which provides a direct determination of the Ti<sup>3+</sup>/ Ti<sup>4+</sup> ratio in nano-sized 786 787 grains. A Thermo Scientific FEI Talos F200S/TEM was used at WPAF Lab for fast EDS 788 mapping of nanoscale grains to minimize beam/sample interactions in 85 powdered samples 789 from Ti-rich sections of 210, 211, 213, 214.43, 218 and, 236.6. Elemental maps generally took 790 one to two minutes to acquire with a sensitivity great enough to detect elemental concentrations 791 in 3 nm size particles. A cryo-stage was also used to prevent heating of temperature-sensitive

sample grains and was operated with a Gatan CT3500 single tilt system with a tilt capability of65 degrees.

794 **C-7.** *Thermochemical analyses*: The thermodynamic calculations of crystallization history 795 of minerals and gaseous species during cooling of a hypothetical solar-gas cluster were carried 796 out with the following assumptions: (i) the system (gases + solids) remained closed; (ii) the 797 mineral- and gaseous species are in chemical equilibrium at all time; (iii) the bulk composition of 798 the system had the elemental abundance ratios of the Apex Asteroid (Table 1) and the H/C/Si ratios of the Solar System estimated by Lodders<sup>1</sup>; and (iv) the total gas pressure was 10<sup>-4</sup> atm. 799 800 Previous researchers<sup>1,52</sup> have adopted the elemental ratios of the carbonaceous chondrites, which are similar to the current Solar Wind, and the total gas pressure of 10<sup>-4</sup> atm in their 801 802 thermodynamic computations of the crystallization history of the Solar System. We have 803 computed the condensation/transformation/disappearance temperatures of nearly 700 compounds 804 as a function of temperature using the software package Thermo-Calc [81] with the SSUB4 805 database [82] developed by means of the CALPHAD method [83,84]. In the CALPHAD method, 806 the Gibbs energy function of each individual phase is modeled as a function of temperature, 807 pressure, and composition. The equilibrium state of a system and its change with respect to 808 temperature, pressure, and composition are obtained by the minimization of the Gibbs energy of 809 the system, and the equilibrium state is represented by the co-existing phases and their respective 810 compositions and quantities. In the present computations, all phases are treated as stoichiometric 811 compounds except the gas phase which consists of over 300 species and is treated as ideal gas. 812 Thermodynamic calculations for the selected redox reactions (Fig. 6c) were carried out using the method in Ohmoto<sup>66</sup> and thermodynamic data in Supplementary Table 5. 813

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- 892 modelings. H.O. coordinated this research and wrote the entire paper. All the authors discussed
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- 894 Competing interests
- 895 The authors declared no competing interests.
- 896 Data availability
- 897 Methods, Extended Data and Supplementary Information are available in the online version of
- the paper. Source data are provided with this paper.
- 899 Additional information
- 900 Correspondence and requests for materials should be addressed to H.O.
- 901

**Extended Data for** 

"Discovery of a 3.46 billion-year-old impact crater in Pilbara, Western Australia" By: H. Ohmoto, U.M. Graham, Z-K. Liu, Y. Tsukamoto, Y. Watanabe, H. Hamasaki & A. Chorney



**ED Figure 1.** Asteroid impact structures in the Marble Bar Chert Member (MBCM). (a) & (b): Aerial (drone) photos of the shattered MBCM from ~250 m above the ground at immediately south of the Marble Bar Pool. Note that the fractures (as wide as ~5 m) run mostly oblique to the MBCM's bedding planes, and that the brecciated fragments (as large as ~10 m) were rotated and dislocated. The black fine-grained sediments (i.e., tsunami sediments) filled the fractures and gaps between the fragments. (c)-(j): Impact-induced tsunami deposits in the Marble Bar Pool area. Notice the various-sized fragments of the MBCM, soft-sediment deformation, incorporation of the Apex Basalt in the chert-rich sediments (b & h) and incorporation of chert fragments in basalts (i & j). (k): Variable sized, mostly rounded blocks and fragments of volcanic rocks of the Duffer Fm. in matrices comprised of fine-grained volcanic rocks at ~4 km SE of the Lennon's Find (Fig. 1c). The tsunami deposits (~15 m thick) in this area lack fragments of the MBCM, because it did not form in this area.



905 ED Figure 2. Impact-induced textures in the MBCM and the Duffer Formation. (a): Photos of selected drill 906 core sections from the ABDP #1 hole, showing the impacted rock textures (fractures, dislocations, veins, 907 fragments) in the MBCM (203.2 - 165.4 m depths) and the Duffer Formation (48.3 - 22.6 m). The yellow scale-908 bars represent 5 cm. (b)-(e): Thin section photos of the mixtures of basalt- and chert-fragments in the tsunami zone 909 of the ABDP drill core. "Ti" represents Ti-rich fragments. (f): A thin section photo of "crushed chert fragments 910 and powders" of the Marble Bar Chert. (g)-(i): HAADF photos of nano-scale stress-features in a chert sample 911 from 171 m depth. (j)-(k): Thin section photos (under reflected light) of Spinifex textures and a quartz-bearing 912 micro vein in the Duffer Formation.



**ED Figure 3. Morphologies, textures and compositions of Ti-rich asteroid fragments.** (a)-(e): Microscope photos (under reflected light) of Ti-rich fragments on unpolished drill-core surfaces. (f)-(l): BSE and elemental maps of a Ti-rich fragments (Ti-, Ca- and V-rich) in basalt (Al-, Fe and Mg-rich) from 257.1 m depth. (m) & (n): BSE and Ti-map of a Ti-rich fragment in basalt at 259.7 m. (o)-(t): BSE photos of Ti-rich fragments in Apex Basalt of the ABDP #1 core.



ED Figure 4. Submarine hydrothermal alteration of the basalt and the tsunami deposits in the ABDP #1

core. (a)-(f): Bulk-rock chemical compositions of the drill-core samples, indicating that the tsunami deposits and the lower part of the Apex Basalt exhibit the characteristics of rocks that were altered by high-temperature (>150°C) submarine hydrothermal fluids, including the losses of Ca and Na and additions of K, Zn & Fe. (g)-(i): EPMA maps of Ti-rich fragments in hydrothermally-altered basalt and tsunami deposits, showing the losses of Ca. (m): EPMA spot analyses of Ti-rich grains, indicating that the matrices of unaltered grains are comprised of titanite or titanitesilica mixtures, the lamellae of mixtures of titanite and rutile, and the altered grains are essentially Ca-free.



**ED. Figure 5. Meteoritic and terrestrial materials that possibly formed in the impact clouds.** (a)-(d): Titaniterich spherules with rutile lamellae, which probably formed by the melting of Ti-rich angular fragments in the impact clouds. (e)-(g): Microscope photos of spherules of basalt melts with inclusions of Ti-rich meteoritic grains. (h): A HAADF photo of basaltic melt spherules with Ti-rich fragments. (i)-(k): Products of condensation of SiO<sub>2</sub>-rich vapor, which was probably generated by the asteroid impact to the MBCM, in the impact clouds.



**ED Figure 6.** Oxidation state of Ti in the Ti-rich fragments. (a) & (b): EELS analyses of Ti atoms in "rutile" grains have revealed the presence of  $Ti^{3+}$  as well as  $Ti^{4+}$  in the forms of  $Ti_4O_7$  and  $Ti_5O_9$  molecules. (c): Relationships between the  $Ti^{4+}/\Sigma Ti$  ratios and  $L_2/L_3$  ratios (after Stoyanov et al.<sup>33</sup>).



ED Figure 7. Involvements of O<sub>2</sub>-rich deep-ocean water in: the formation of primary haematite in the MBCM; the alteration of the basalt and tsunami deposits; and the alteration of meteoritic minerals. (a)-(f): Evidence for the involvement of O<sub>2</sub>- and SO<sub>4</sub><sup>2</sup>-rich ocean water in the alteration of the Apex Basalt and tsunami zone: negative Ce anomalies, increased Ba and U contents and increased  $Fe^{3+}/\SigmaFe$  and U/Th ratios of rocks in the hydrothermal alteration zone. (g)-(o): Microscopic and SEM images of sub-micron-sized primary haematite crystals in the MBCM (from Hoashi et al.<sup>19</sup>). (k): Fragmented red cherts in the tsunami deposits, clearly indicating that the haematite crystals in the MBCM formed prior to the asteroid impact. (l)-(p): SEM photos of meteoritic Fe-oxide crystals in the tsunami deposits, showing that they are more than 10 times larger than the primary Bhydrothermal haematite crystals (h-j) in size. Their original Fe/O ratios (~0.90) are similar to those of witsite (FeO); the lower Fe/O ratios (e.g., 0.82) in some crystals (Supplementary Table 3) were probably due to the reactions with O<sub>2</sub>-rich atmosphere and ocean at 3.46 Ga ago.



**ED Figure 8.** Iridium-rich Fe metal. (a)-(h): EPMA analyses of a ~50  $\mu$ m-sized Fe-rich grain. (i): Variations in the Fe/O ratios along Profile A. (j): Variations in the Fe/O ratios along Profile B. (k): Concentration profiles of Fe, O, Ir, Si, and Al along Profile B. Variations in the Fe/O ratios in (i) and (j) suggest that the original phase in this part of the grain was metallic iron (Fe/O >>1) and possibly wüstite (FeO), and that the decreases in the Fe/O ratios less than 1 were due to the oxidation of Fe-metal and wüstite to magnetite and haematite during/after the passage through an oxygenated atmosphere and ocean.



ED Figure 9. Carbon-rich phases from the Apex Asteroid. (a)-(j): Various carbon-bearing nanocrystals (graphene, tubular C, graphite and silicon carbide). (k)-(m): C/Al/Ca/Si/Cu alloys. Photos under HRTEM and HAADF (see C-6 in *Methods*).



**E.D.Figure 10. Thermochemical analyses of the formation conditions of the Apex** Asteroid at  $\sum P = 10^{-10}$  atm. (a): Temperatures for the condensation and transformation of minerals from a solar gas cluster ("the Apex gas cluster"). (b): Abundances of the major gaseous species in the Apex gas cluster. (c): Stability relationships among some redox-sensitive minerals as a function of H<sub>2</sub>/H<sub>2</sub>O ratio of the coexisting vapor phase and temperature. Superimposed is the change in H<sub>2</sub>/H<sub>2</sub>O ratio of the Apex gas cluster of (b).