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The phase diagram of the Fe-P binary system at 3 GPa and implications for phosphorus in the lunar core

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Abstract

Phosphorus is a potential candidate in the metallic core of the Moon. The phase diagram of the Fe-P binary system was investigated at the pressure of 3 GPa and temperatures of up to 1600 °C. Up to 3.0 wt% and 10.4 wt% phosphorus can dissolve in the solid iron and liquid Fe-P phases at 1100 °C and 3 GPa, respectively. The eutectic temperature on the iron-rich side was determined as 1085 °C at 3 GPa. The solubility of phosphorus in the iron decreases from \sim 1.4 wt% at 1100 °C to ~ 0.7 wt% at 1500 °C and 3 GPa. Structure of the solid iron in the quenched sample is the body-center cubic, corresponding to α -Fe phase. Extending the phosphorus solubility in the solid iron to the present lunar core conditions yields a maximum phosphorus concentration in a fully crystallized iron core of 0.85 ± 0.15 wt%. If there are Ni and C in the core, the value would be depressed to 0.4 ± 0.1 wt%. In addition, based on a simple siderophile mass balance between the bulk Moon (BM) and bulk silicate Moon (BSM) and a modeled phosphorus partition coefficient, $D_{\text{P-Moon}}^{\text{core/mantle}}$ (40–200) for the lunar magma ocean, a bulk silicate Earth-like P content (82 ± 8 ppm) in the initial Moon yields a lunar core with <0.3 wt% P. Some other potential light elements such as S and C could reduce the P content in the lunar core. Furthermore, the partition coefficient of phosphorus in the iron and liquid melt $(D_P^{SM/LM})$ was found to be 0.10 ± 0.03 at 3 GPa. Taking the sulfur into account, the $D_{\rm P}^{\rm SM/LM}$ increase to 0.18 ± 0.02 at 5 GPa in the S-rich liquid metal (~8 wt%). In the case of a solid lunar inner core and S-bearing liquid outer core, their P contents were assessed to be less than 0.09 ± 0.01 wt% and 0.51 ± 0.01 wt%, respectively, when the lunar core's storage of P is <0.3 wt%. The moderate phosphorus solubility in the solid iron, combined with the assumption of abundant phosphorus in the bulk Moon, indicates that the phosphorus concentration in the lunar core could higher than previously thought.

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Keywords: Fe-P binary system; High pressure; Phase diagram; Lunar core

1. INTRODUCTION

Based on the geophysical evidence from lunar seismology (Lognonné and Johnson, 2007; Weber et al., 2011) and laser ranging (Williams et al., 2001), Moon is supposed

https://doi.org/10.1016/j.gca.2019.03.037 0016-7037/© 2019 Elsevier Ltd. All rights reserved. to have a small metallic core, even with a solid inner and liquid outer core. Like the chemical composition model for the Earth's core, some light elements, such as carbon and sulfur, are suggested to alloy with iron in the lunar core. To explain geophysical observation, the concentration of sulfur in the lunar outer core has been constrained in different models (Weber et al., 2011 - <6 wt% S; Jing et al., 2014 - 3.6 wt% S; Antonangeli et al., 2015 6–11 wt%; Kuwabara et al., 2016). Models for metal/silicate partitioning of siderophile elements in the lunar magma ocean

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(LMO) also support a sulfur-rich core (Rai and van Westrenen, 2014, 6-8 wt% sulfur). However, the sulfur may not be the dominant light element in the lunar core. Steenstra et al. (2017) suggest a carbon-dominant core with 0.6-4.8 wt%, including relative lower sulfur concentration (<0.16 wt%). Recent studies have experimentally determined plausible ranges of S in the bulk silicate Moon which suggest the estimated S concentration in the lunar mantle could not meet the requirement for a lunar core that contains several wt% S (Bombardieri et al., 2005; Ding et al., 2017; Steenstra et al., 2018). According to the phase equilibria experiments on the Fe-Ni-S-C system at the temperature and pressure conditions relevant to the lunar interior, Righter et al. (2017) proposed a low S- and C content core (0.5 wt% C, 0.375 wt% S) which provides a simple heat source and mechanism for a lunar core dynamo (core crystallization). That composition would result in 3.8-11.8 wt% S and 1.7–2.4 wt% C in the fluid outer core with \sim 1.2 wt% C and ~ 0.02 wt% S in the solid Fe inner core.

Due to the depletion relative to CI chondrites and bulk silicate Earth (BSE), phosphorus is believed to be enriched in the iron-nickel core of the Moon by the core-mantle differentiation (Righter and Drake, 1996; Righter, 2002). Depending on the P/Nd ratio in meteorites, the phosphorus content in the bulk silicate Moon (C_{BSM}^{P}) was estimated to be 20 ppm, with a phosphorus content of 43 ppm in the bulk Moon (C^P_{BM}) (O'Neill, 1991). Approximately 0.1 wt% P may present in the lunar core by the core-mantle differentiation, which means that 55 wt% of the total phosphorus budget is in the core (Drake, 1987; Sha, 2000). Righter (2002) estimated the P concentration in the lunar mantle by different Moon-forming impact models, resulting in 8-56 ppm. However, this value is still in debate according to the theory that the abundance of major elements are same on the Moon and Earth (95 ± 15 ppm P from Ringwood, 1991). In the estimation of the bulk silicate Moon (BSM) by Hauri et al. (2015), 0.0206 wt% P₂O₅, equal to 90 ppm P, is present in the lunar mantle. The bulk silicate Moon enriched in crustal components model (the cBSM model)

proposed by Togashi et al. (2017) suggested that up to 0.04 wt% P₂O₅, equal to 174 ppm P, was present in the initial silicate Moon after the giant impact, much higher than the 20 ppm estimated previously by O'Neill (1991). Because of the core-mantle differentiation, the C_{BSM}^{P} is the key that controls the P budget in the lunar core thought the C_{BSM}^{P} is still unclear. In addition, the Fe-P phase diagram may provide valuable information to constrain the concentration of phosphorus in the lunar core.

Saklatwalla (1908) firstly published the phase relationship of the Fe-P system (0-24 wt% P) at ambient pressure conditions. Then Konostantinow (1910) and Haughton (1927) measured the Fe-P binary phase diagram up to 24 and 32 wt% P, respectively. The phase diagram at ambient pressure was reassessed by Zaitsev et al. (1995) using thermodynamic analyses. Stewart and Schmidt (2007) conducted Fe-P-S and Fe-P experiments at 23 GPa using multi-anvil presses. Compared with the value of 2.6 wt% at ambient pressure shown in Fig. 1(a), the solubility of phosphorus in α-Fe increases to 4 wt% at 23 GPa (Fig. 1 (b)). Up to $\sim 9 \text{ wt\%}$ phosphorus is soluble in the Fe-P melt at 1275 °C and 23 GPa (Fig. 1b). Even though the phosphorus solubility in the iron is appreciable, Stewart and Schmidt (2007) considered that phosphorus would not form a distinct phase within terrestrial planet cores due to the expectation of its minor concentration in planetary cores. However, to determine the phosphorus content in the Moon's core, the previous Fe-P phase diagram is insufficient and only contains a few experiment data conducted at 23 GPa (Stewart and Schmidt, 2007), which is inappropriate for the lunar core conditions at 4.5 GPa (coremantle-boundary) to 5.3 GPa (inner core) (Kuskov and Kronrod, 1998; Garcia et al., 2011). A very recent study investigated the pressure dependence of the liquidus and solidus temperatures on the iron-rich side of the Fe-P binary system by using in-situ ultrasonics at 2 and 4 GPa (Chantel et al., 2018). But, they provide little chemical composition information for the Fe-P binary system. Here we investigated the Fe-P phase diagram at 3 GPa and



Fig. 1. Fe-P phase diagrams at ambient pressure (a) (Baker, 1992; Zaitsev et al., 1995) and 23 GPa (b) (Stewart and Schmidt, 2007). Circle and square markers represent the experimental data.

determined the phosphorus solubility in the iron phase. Combined with the phosphorus partitioning model between the lunar mantle and core, we further constrained the phosphorus content in the lunar core.

2. METHODS

Stoichiometric amounts of red phosphorus (99.9 wt%, Alfa Aesar) and iron powder (99.99 wt%, Alfa Aesar) were mechanically mixed and ground under ethanol in an agate mortar to synthesize FeP by solid-state reaction. After airdrving of the mixture, it was pre-compressed and loaded into a 6 mm length hexagonal boron nitride capsule (h-BN, ϕ 4 mm) and then sealed and stored in a desiccator. FeP was synthesized at 1 GPa and 1000 °C for 3 hours in a multi-anvil apparatus. These P-T conditions are the same as those used by Gu et al. (2011). The recovered sample was identified by powder X-ray diffraction to be a pure FeP phase. We then use FeP and pure iron powder as the starting materials. Synthesized FeP is used in this study because it is representative of protophosphorus and widely distributed in meteorites. A similar starting material was also used in the Fe-FeS studies by Fei et al. (1997, 2000).

All experiments were conducted on a DS 6×600 t cubic-anvil-type apparatus with WC anvils of 23.5 mm truncation edge length at the Institute of Geochemistry, Chinese Academy of Sciences, Guiyang, China. The synthe-

sized FeP powder was thoroughly mixed with a pure iron powder to prepare different starting materials with phosphorus contents varying from 5.8 to 27 wt%. Three different starting materials were put into a boron nitride capsule, as shown in Fig. 2. An Al₂O₃ sleeve thermal insulator and a graphite heater were used in the sample assembly. The pressure was calibrated by the quartz phase transition and NaCl melting curve at high pressures and high temperatures (Bose and Ganguly, 1995; Rybacki et al., 1998). The uncertainty for the pressure is ± 0.2 GPa. Temperatures were monitored by a W5%Re-W26%Re thermocouple which was placed close to the sample, with the uncertainty of \pm 5 °C. The temperature gradient from the sample center to the thermocouple was less than 10 °C, measured by two thermocouples in another experiment. All the runs were carried out from 1075 to 1600 °C with an interval of 50 or 100 °C. Samples were compressed up to 3 GPa and heated to the designed temperature with a heating rate of 60 °C/min, and then held at that temperature for 8 or 10 hours for each experiment. The runtime was chosen by different starting component and referred to the equilibrium time in the experiments by Stewart and Schmidt (2007). Run products were then quenched by shutting down the power. Several products (run# FP-1-1, FP-7-1) were ground into powders for X-ray diffraction analyses, and the others were mounted in epoxy, polished carefully and carbon-coated for quench texture and chemical composition analyses.



Fig. 2. Cross section of the sample assemblage used in this study.

Powder X-ray diffraction patterns of several quenched samples were collected using an X-ray diffractometer with Cu-K α radiation. The iron phosphide phases (FeP, Fe₂P, and Fe₃P) had lattice parameters consistent with previous results (Table 1). We also checked the quenched iron sample from the run of FP-7-2 (Fe_{94.2}P_{5.8}) by micro-focused Xray diffraction analyses, using a RIGAKU D/Max Rapid IIR micro-diffractometer operated at 40 kV and 250 mA. The X-ray wavelength of Cu-K α l generated from copper anode and graphite monochromatic is 1.54056 Å. The two-dimensional diffraction data were converted to onedimensional (20-I) data by Display software of the instrument. The result is shown in Fig. 3.

The chemical composition analyses were performed by using a JXA-8230 electron microprobe operated at 20 kV and 10 nA at the Testing Center of Shandong Bureau, China. Iron and phosphorus were analyzed by wavelength dispersive spectrometry (WDS) employing analytical standards of pure iron and apatite, respectively. The ratios of peak/background for phosphorus in the low-P iron are generally ~11 at our experimental conditions. Nitrogen, boron, and oxygen were also directly analyzed as potential contaminants, using GaN, B metal, and Al₂O₃ as standards. There are no significant signals during analyses with the peak and background counting times of 20 s and 10 s respectively. Heterogeneity in the quenched melt phase is common in quenched metal alloys. To ensure representative compositions, a fully focused beam was used for crystalline iron and iron phosphide and beam diameter of 20-50 µm was used for quenched melt domains. It is as the same as the method used by Stewart and Schmidt (2007) and Dasgupta et al. (2009). Usually, 10 to 15 points were analyzed for the melts and solid phases. Final results were reduced by a ZAF correction method (Armstrong, 1995) and are listed in Table 2.

3. RESULTS AND DISCUSSIONS

3.1. Fe-P phase diagram and previous studies

Backscattered electron images of the run products show unambiguous quench textures of the molten samples (Fig. 4). Below the melting temperatures, different mineral phases are present in solid xenomorphic-granular phases and have obvious grain boundaries. Quenched liquids have the similar dendritic growth textures as described by Stewart and Schmidt (2007) in the Fe-P-S system at 23 GPa and by Dasgupta and Walker (2008) in the Fe-C



Fig. 3. Micro-focused X-ray diffraction analyses to the run product of FP-7-2 (Fe_{94.2}P_{5.8}). X-ray beam size with diameter of 200 μ m was used to analyze the interest region. The inset shows a two-dimensional diffraction pattern which was collected by the detector. The background of the diffraction data remains. Only the peaks of body-center cubic structure of iron are found and labeled with corresponding crystal indices (hkl).

system at 2 GPa. In the phosphorus-rich runs, some samples undergo partial melting with small mineral chips scattered in the melts (Fig. 4g). The temperature of the Fe-P eutectic on the iron-rich side is constrained at ~1085 °C due to the Fe-P liquid which appears from 1075 to 1100 °C (Fig. 4a–f). The mixture of FeP and Fe₂P begins to melt at about 1400 °C (Fig. 4h, i). In the run of Fe_{81.8}P_{18.2}, crystalline Fe₂P + Fe₃P disappears and quenched Fe-P melt coexists with crystalline Fe₂P when T > 1185 °C.

With increasing temperature, the phosphorus content in the liquid Fe-P decreases from ~10.1 wt% at 1100 °C to ~9.0 wt% at 1500 °C. The solubility of phosphorus in the solid iron is 0.7–1.3 wt% depending on the temperatures (Table 2). Some crystalline solid iron phases containing ~2.8 wt% phosphorus are noted in the run below 1250 °C and coexist with low-P solid iron (Table 2). The runs on the iron-rich side at 2 GPa indicate that 0.7–1.0 wt% phosphorus could dissolve in the solid iron (Table 2). Compared with ideal iron phosphides, the experimental products of FeP, Fe₂P, and Fe₃P have a slight excess of iron and this phenomenon also occurs in the Fe-P system at 23 GPa (Stewart and Schmidt, 2007).

3.2. Pressure effects on the Fe-P eutectic

The Fe-P phase diagram at 3 GPa (Fig. 5a) is widely consistent with the topology of phase diagram at ambient pressure (Zaitsev et al., 1995) and 23 GPa (Stewart and

Table 1

Lattice parameters of iron phosphide phases produced in this study and previous works.

	Space group	а	b	с	$V(\text{\AA}^3)$	Ref.
FeP	Pnma	5.1862(3)	3.0965(2)	5.7868(3)	92.93(1)	This study
		5.1826(6)	3.0976(3)	5.7838(7)	92.85(1)	Gu et al. (2011)
Fe ₂ P	P-62m	5.8651(1)		3.4555(1)	102.94(1)	This study
		5.8764(3)		3.4493(1)	103.14	Dera et al. (2008)
		5.8650(10)		3.456(1)	102.95(3)	Scott et al. (2008)
Fe ₃ P	I-4	9.0998(2)		4.4615(2)	369.44(2)	This study
		9.099(2)		4.463(2)	369.5(2)	Scott et al. (2007)
		9.1003(1)		4.4685(1)	370.06(1)	Gu et al. (2014)

Table 2
Starting materials, experimental conditions, coexisting phases, and phase compositions in the run products of Fe-P system at 2 and 3 GPa.

Run #	t, h	T, °C	Starting materials (wt %)	Phases	Р	Fe	Total
P = 3 GPa							
FP-7-1	8	1075	$Fe_{94,2}P_{5,8}$	Fe ₃ P	15.16(7) ^a	85.50(49)	100.65(55)
				α-Fe	2.95(3)	98.40(25)	101.35(23)
					1.40(1)	98.46(27)	99.86(25)
			$Fe_{90}P_{10}$	Fe ₃ P	15.17(4)	85.46(27)	100.85(24)
			2010	α-Fe	2.92(2)	97.37(11)	100.28(9)
					1.32(5)	99.48(22)	100.80(20)
			$Fe_{86} P_{137}$	Fe ₃ P	15.22(9)	85.13(36)	100.34(36)
				α-Fe	1.13(1)	98.37(29)	99.50(29)
FP-7-2	8	1100	$Fe_{94} {}_{2}P_{58}$	Melt	10.10(33)	90.11(52)	100.21(27)
			212 210	α-Fe	2.94(3)	97.62(56)	100.56(58)
					1.35(1)	98.84(42)	100.19(41)
			$Fe_{90}P_{10}$	Melt	10.39(47)	90.34(51)	100.74(37)
			2010	α-Fe	3.05(2)	97.21(29)	100.26(29)
			$Fe_{86} {}_{3}P_{13} {}_{7}$	Fe ₃ P	15.21(3)	85.52(10)	100.73(10)
			000 12.7	Melt	8.83(26)	91.58(32)	100.41(23)
FP-1-1	10	1150	$Fe_{94,2}P_{5,8}$	Melt	9.98(30)	88.78(57)	98.75(50)
			212 210	α-Fe	1.32(2)	98.33(38)	99.65(40)
			$Fe_{81,8}P_{18,2}$	Fe ₃ P	14.91(3)	84.94(19)	99.85(18)
			01.0 10.2	Fe ₂ P	21.05(8)	79.42(20)	100.48(20)
			Fe ₇₃ P ₂₇	Fe ₂ P	21.04(8)	78.44(26)	100.38(33)
			- 15 21	FeP	34.40(21)	65.31(32)	98.42(50)
FP-7-3	8	1185	Feod 2P5 8	Melt	10.62(23)	88.24(29)	100.84(20)
				α-Fe	1.32(5)	98.77(27)	100.09(28)
			FeooP10	Melt	10.26(35)	90.17(14)	100.44(23)
			90- 10	α-Fe	2.40(32)	97 93(44)	100 33(22)
			Feel Pie 2	Melt	10.93(23)	88 81(33)	99 74(34)
				Fe ₂ P	21.13(13)	79.49(41)	100.61(46)
FP-1-2	10	1200	Feod 2P5 8	Melt	9 78(36)	89 45(81)	99 24(72)
	10	1200	2 074.22 3.8	α-Fe	1 26(2)	99.02(48)	100.28(51)
			Feel Pie 2	Melt	9 60(54)	89 35(87)	98 94(43)
			- 001.0- 10.2	Fe ₂ P	20.96(9)	79.63(18)	100.59(20)
			Fe72P27	Fe ₂ P	20.97(6)	79 59(35)	100 56(40)
			2 0/32 2/	FeP	3478(13)	65 9(13)	100 68(24)
FP-1-3	8	1225	Feel Pie 2	Melt	9 62(34)	89 31(36)	98 92(42)
	-			FeaP	20.72(9)	79 13(24)	99.85(28)
			Fe ₇₂ P ₂₇	Fe ₂ P	20.84(7)	78.71(50)	99.55(53)
			/ 5- 2/	FeP	34.30(17)	65.44(49)	99.74(61)
FP-2-1	8	1250	Feod 2P5 8	Melt	9.67(16)	89.38(40)	99.05(53)
			94.2 3.6	α-Fe	1.24(1)	97.92(11)	99.16(11)
			$Fe_{90}P_{10}$	Melt	10.03(38)	90.31(43)	100.34(16)
			-90 10	α-Fe	2.91(21)	96.76(29)	99.67(18)
			$Fe_{86} P_{137}$	Melt	9.46(25)	89.84(37)	99.29(45)
				Fe ₃ P	15.22(10)	85.37(26)	100.59(36)
FP-2-3	10	1325	$Fe_{94} {}_{2}P_{58}$	Melt	9.88(24)	90.73(14)	100.61(18)
			212 210	α-Fe	1.06(1)	99.13(17)	100.19(18)
			$Fe_{81,8}P_{18,2}$	Melt	11.08(47)	89.37(37)	100.44(17)
			01.0 10.2	Fe ₂ P	21.14(8)	79.59(41)	100.72(46)
			Fe ₇₃ P ₂₇	Melt	25.93(40)	74.91(57)	100.83(44)
			- 15 21	Fe ₂ P	21.18(5)	79.56(10)	100.75(7)
				FeP	34.4(35)	66.28(52)	100.68(48)
FP-3-1	8	1350	$Fe_{94} {}_{2}P_{58}$	Melt	9.53(14)	91.04(29)	100.57(19)
			212 210	α-Fe	1.14(2)	98.71(40)	99.84(39)
			$Fe_{90}P_{10}$	Melt	9.52(27)	90.21(29)	99.73(20)
			$Fe_{86,3}P_{13,7}$	Melt	12.87(38)	87.09(36)	99.97(26)
FP-3-2	8	1375	$Fe_{94,2}P_{5,8}$	Melt	10.36(46)	89.87(73)	100.22(36)
			74.2 5.6	α-Fe	1.31(2)	98.25(33)	99.57(32)
			$Fe_{81,8}P_{18,2}$	Melt	10.37(19)	89.86(31)	100.23(15)
			01.0 10.2	Fe ₂ P	21.19(10)	79.88(39)	101.08(44)
			$Fe_{73}P_{27}$	melt	26.28(34)	74.28(41)	100.56(40)
				Fe ₂ P	21.08(18)	79.17(17)	100.24(32)
				FeP	34.62(8)	65.66(33)	100.28(39)
					、 /	(continue	d on next page)

Table 2 (*continued*)

Run #	t, h	T, °C	Starting materials (wt %)	Phases	Р	Fe	Total
FP-3-3	10	1400	Fe _{94.2} P _{5.8}	Melt	9.42(15)	91.01(26)	100.43(22)
				α-Fe	1.05(1)	100.12(37)	101.17(37)
			$Fe_{90}P_{10}$	Melt	9.77(7)	91.04(18)	100.81(16)
			Fe _{86.3} P _{13.7}	Melt	13.58(21)	87.21(33)	100.79(23)
				Melt	9.39(18)	90.85(22)	100.24(3)
FP-4-1	10	1425	Fe _{94.2} P _{5.8}	Melt	9.13(17)	91.25(28)	100.25(26)
				α-Fe	0.96(2)	99.42(35)	100.38(34)
			Fe _{81.8} P _{18.2}	Melt	13.26(126)	86.50(91)	99.75(36)
				Fe ₂ P	20.91(6)	80.32(27)	101.23(31)
FP-4-3	8	1475	Fe _{94.2} P _{5.8}	Melt	8.60(22)	91.18(35)	99.78(33)
				α-Fe	0.71(00)	100.27(20)	100.97(19)
			Fe _{81.8} P _{18.2}	Melt	14.65(57)	84.97(57)	99.63(22)
				Fe ₂ P	20.91(14)	79.60(21)	100.51(32)
			Fe ₇₃ P ₂₇	Melt	26.33(40)	73.31(48)	99.65(20)
				Fe ₂ P	21.27(15)	79.27(18)	100.54(9)
				FeP	34.45(40)	66.25(35)	100.70(31)
FP-5-1	8	1500	$Fe_{94,2}P_{5,8}$	Melt	8.20(17)	90.45(23)	98.65(22)
				α-Fe	0.78(3)	98.96(24)	99.74(25)
FP-5-3	8	1550	$Fe_{94,2}P_{5,8}$	Melt	10.02(26)	89.71(27)	99.73(12)
				α-Fe	1.01 (2)	98.75(13)	99.76(13)
FP-6-1	8	1600	$Fe_{94,2}P_{5,8}$	Melt	9.61(13)	89.61(15)	99.22(13)
				α-Fe	0.73(2)	98.58(17)	99.31(15)
P = 2 GPa						. ,	
FP-9-1	10	1200	$Fe_{94,2}P_{5,8}$	Melt	11.14(37)	89.67(31)	100.81(23)
				α-Fe	1.07(3)	99.74(38)	100.80(36)
FP-9-2	6	1350	$Fe_{94,2}P_{5,8}$	Melt	9.80(42)	89.40(26)	99.20(25)
				α-Fe	0.81(2)	98.71(34)	99.52(34)
FP-9-3	6	1500	$Fe_{94,2}P_{5,8}$	Melt	9.52(28)	90.93(40)	100.45(33)
				α-Fe	0.82(5)	98.77(25)	99.59(28)

^a Values in wt%. Numbers in parentheses indicate standard error of the mean based on replicate analyses $(\delta_M = \delta/\sqrt{n})$.

Schmidt, 2007). Liquidus temperatures on the iron-rich side (Fe_{94.2}P_{5.8}) is 1375–1400 °C which is about 100 °C higher than the temperatures determined by in-situ ultrasonics at $1257\pm50~^\circ\mathrm{C}$ and 4 GPa with starting P content of 5 wt% (Chantel et al., 2018). The reason is possibly the large uncertainty (±50 °C) in their measured experimental temperatures. The eutectic temperature on the iron-rich side is 1085 °C, about 40 °C higher than the 1048 °C eutectic at ambient pressure and 190 °C lower than the 1275 °C eutectic at 23 GPa (Stewart and Schmidt, 2007). Among the Fe-C, Fe-P and Fe-S systems, Fe + Fe₃S has the lowest eutectic temperature of 1100 °C under the similar pressure (~21 GPa, Fei et al., 2000). Around 3 to 5 GPa, the Fe-P eutectic temperature on the iron-rich side is ~ 100 °C lower than that of the Fe-C system (Hirayama et al., 1993; Fei and Brosh, 2014) and ~100 °C higher than the Fe-S system (Usselman, 1975). Phosphorus shows a moderate contribution to reducing the melting temperature of the iron alloy, which is significant for the formation of a liquid outer core in the Earth and terrestrial planets.

The eutectic liquid contains $\sim 10.5 \text{ wt}\%$ P at ambient pressure and $\sim 9 \text{ wt}\%$ P at 23 GPa, which suggests the solubility of phosphorus in the Fe-P eutectic liquid decreases with increasing pressures (Fig. 1). There is about 10.1 wt % P dissolved in the eutectic liquid at 3 GPa. In contrast, pressure has a positive effect on the maximum phosphorus solubility in the solid iron. As much as 4 wt% P is detected in the crystalline solid iron at 23 GPa, more than the result of 2.6 wt% P at ambient pressure (Fig. 1a and b). The crystalline solid iron phase contains up to 2.9 wt% P at 3 GPa in the present study, slightly higher than the value at ambient pressure.

3.3. Low-P and high-P crystalline solid iron

There are two iron phases on the iron-rich side, α -Fe (body-center-cubic, *bcc* structure) and γ -Fe (face-center-cubic, *fcc* structure) in the Fe-P phase diagram at ambient pressure (Fig. 1a). With increasing phosphorus content in the iron, the α -Fe is more stable because the P is a *bcc*-stabilizer and is thought to narrow down the stable temperature range of the γ -phase iron by forming the so-called " γ -loop" (Pepperhoff and Acet, 2001).

We analyzed the iron phase in a region which is as the same area of Fig. 4f (element mapping was shown in Fig. 6a). The structure of the quenched iron is the bodycenter cubic (*bcc*-Fe) (Fig. 3). Therefore, the two different P-bearing solid irons are both α -Fe which stable region is consistent with previous studies (Okamoto, 1990; Perrot et al., 2002). According to the Fe-P phase diagram at ambient pressure (Fig. 1a), there must be a 2-phase loop between the α -Fe and γ -Fe, called $\alpha + \gamma$ region that was shown in Fig. 5(b). The γ -Fe is generally considered to be unquenchable (Pepperhoff and Acet, 2001). Stability of γ -iron in the

Fig. 4. Backscattered electron images of the run products of FP-7-1 (a, b, c), FP-7-2 (d, e, f), FP-7-3 (g), FP-2-3 (h) and FP-4-3 (i). Images of a (d), b(e) and c(f) indicate the quenched phase structure of $Fe_{86.3}P_{13.7}$, $Fe_{90}P_{10}$ and $Fe_{94.2}P_{5.8}$ respectively. The black phases disseminated in Fe and Fe₃P are holes created during polishing because Fe-P alloy is fragile.

Fig. 5. (a) is the phase diagram of Fe-P system at 3 GPa given by this study. (b) is a close-up phase diagram on the iron-rich side (0–16 wt% P). Pure iron melting temperature is taken from Liu and Bassett (1975). The phase stable range of γ -Fe is referred from the iron phase diagram by Acet et al. (2001). Righ-black, grey, and left-black circles represent liquid, solid + liquid, and solid minerals, respectively, in run products of different starting materials and different temperatures. Open symbols with error bars are P concentration in the iron, melts, and iron phosphide. Solidus and liquidus are determined by partial-melting temperatures and full-melting temperatures. The dashed lines in (b) are solidus curves for γ -Fe, γ + α iron, and α -Fe at ambient pressure reported by Okamoto (1990).

metallic meteorites also strongly depends on the chemical composition and cooling-rate (Uhlig, 1954). The solubility of P in γ -Fe has been measured by various indirect meth-

ods, such as thermal analysis, metallography and magnetic measurement (Haughton, 1927; Vogel, 1929; Roquet and Jegaden, 1951; Lorenz and Fabritius, 1962; Fisher et al.,

Fig. 6. Selected EPMA mapping analyses of the quenched sample of $Fe_{94.2}P_{5.8}$ at 1100 °C at 3 GPa. Image (b) is a close-up of the white square region in the image (a). Pictures shown from left to right are backscattered electron image, P-mapping image and Fe-mapping image, respectively. Low-P and High-P represent low P and high P concentration iron.

Fig. 7. Partition coefficient of phosphorus between solid metal and liquid metal varies with pressures (T = 1250-1550 °C). The inset is the measured $D_p^{\rm SM/LM}$ in Fe-P binary system from this study at the temperatures from 1100 to 1600 °C. Open triangles and stars are the results at 3 and 2 GPa, respectively. Solid squares are the $D_p^{\rm SM/LM}$ in Fe-Ni-S-P system from Stewart et al. (2009) with S content of 8–14 wt%. Solid triangle is the data calculated from S-free to 8 wt% S in the liquid metal for Fe-S-P system at ambient pressure (Righter and Drack, 2000). There is a linear trend for $D_p^{\rm SM/LM}$ as increasing pressures in Fe-S-P system. Grey area in the inset figure is the potential temperatures at the current lunar coremantle boundary. Trapezoid grey area is the possible $D_p^{\rm SM/LM}$ at various pressure and sulfur content situations.

1966). Thermodynamic calculations predict that the maximum solubility of phosphorus in solid γ -Fe is ~0.3 wt% at ~1150 °C and ambient pressure (Lorenz and Fabritius, 1962; Okamoto, 1990). On the other hand, according to the extrapolation from Fe-P-C and Fe-P-Ni experiments, the metastable solubility of P in γ -Fe at ambient pressure is estimated at 1.1, 0.6, 0.3 wt% at 1000, 900, 800 °C, respectively (Kaneko et al., 1965).

3.4. Phosphorus partition coefficient between the solid metal and liquid metal

Using the phosphorus concentration in the iron and melts with starting materials of $Fe_{94,8}P_{5,2}$, we calculate the phosphorus partition coefficient between the solid and liquid metal $(D_p^{\text{SM/LM}})$. The value of the $D_p^{\text{SM/LM}}$ decreases with increasing temperature (Fig. 7). At the modeled temperatures for the lunar core-mantle boundary, 1200-1550 °C, the $D_{\rm p}^{\rm SM/LM}$ is 0.10 \pm 0.03 at 3 GPa. It is consistent with the partition coefficient formula at ambient pressure given by Righter and Drake (2000). They indicate that the natural logarithm of $D_p^{\text{SM/LM}}$ has a linear relationship with the sulfur content in the melts: $\ln D_p^{\text{SM/LM}} = 16.19 \times (Xs)^2 - 2.42$, where Xs is sulfur mole fraction. Thus, the $D_{\rm p}^{\rm SM/LM}$ are 0.095 with sulfur-free and 0.115 with 8 wt% (Xs = 0.13). It is obvious that more phosphorus is into solid iron with the addition of S. For pressures, the $D_p^{SM/LM}$ at 3 GPa is slightly higher than the result at 2 GPa which suggests that more phosphorus would be dissolved in the solid iron at high pressures. In the study of Fe-S-P system at high pressures, $D_p^{\text{SM/LM}}$ increase from 0.23 ± 0.02 at 9 GPa, 0.32 ± 0.03 at 15 GPa to 0.58 ± 0.03 at 23 GPa (Stewart et al., 2009). The result at 23 GPa is beyond the trend because the sulfur content is 13 ± 1 wt%, higher than the lower pressure experiments (8–10 wt%).

4. PHOSPHORUS IN THE LUNAR CORE

4.1. Phosphorus budget in the lunar core by partitioning model

The Moon has a small metallic core that has been supported by numerous geochemical and geophysical studies (Kuskov and Kronrod, 1998; Righter, 2002; Weber et al., 2011; Williams et al., 2014). There are three factors controlling the phosphorus content in the lunar core (C_{core}^{P}) : phosphorus partition coefficient between the core and mantle $(D_{P-\text{Moon}}^{\text{core/mantle}})$, core mass, and the phosphorus content in the initial bulk Moon (C_{BM}^{P}) . Through a simple model of mass differentiation (core's mass is set at 2.5 wt% of the bulk Moon), the C_{core}^{P} could be calculated by the following equation:

$$C_{\text{core}}^{\text{P}} = \frac{C_{\text{BM}}^{\text{P}}}{1/D_{\text{P-Moon}}^{\text{core/mantle}} + 0.025(1 - 1/D_{\text{P-Moon}}^{\text{core/mantle}})}$$
(1)

On the one hand, according to the cosmochemical composition models suggested by Allègre et al. (1995) and McDonough (2003), the *D*core/mantle P-Earth of the Earth is calculated as 20–50, which has large uncertainty because of volatile-loss (Halliday and Wood, 2010; Righter, 2011).

Fig. 8. Phosphorus concentration in the lunare core, C_{core}^{P} as a function of P storage in the initial Moon and reciprocal of partition coefficient of phosphorus between core and mantle, $1/D_{P-Moon}^{core/mantle}$. The initial bulk Moon phosphorus content (C_{BM}^{P}) are plotted as curves varying from 50 to 300 ppm. Verticle dashed line is the lowest estimate of $D_{P-Moon}^{core/mantle}$ at 40. Light grey area is the result of P core storage with a BSE-like P content (82 ± 8 ppm) in the initial bulk Moon.

The Moon has a similar partition coefficient (39-69) to the Earth (Steenstra et al., 2016a). But, this value should be reexamined because it is based on the relatively low C_{BM}^{P} (43 ppm) suggested by O'Neill (1991) and Righter (2002). Their estimate for the C_{BM}^{P} is significantly influenced by the siderophile elements abundance of the impactor. On the other hand, the partition experiments for the phosphorus between the metal and silicate melt provide more constrains. Steenstra et al. (2015) extended the phosphorus partition coefficient formula by combining all available experimental metal-silicate partitioning data from a wide range of studies conducted at core conditions of the Moon. The formula proposed a modeled $\log D_{P-Moon}^{core/mantle}$ between 1.9 and 2.3 $(D_{P-Moon}^{core/mantle} = 79-200)$ corresponding to the metalsilicate equilibration of a sulfur-rich (11 wt% in the core) and nickel-bearing metallic core at the conditions of $P \approx$ 4 GPa, T = 2377 °C, $\Delta IW = -2.15$. However, there may be a lower temperature condition for the lunar magma ocean (LMO): T = 1900 °C, $\Delta IW = -2.2$, $X_S = X_C = 0$, where X_S and X_C are mole fraction of S and C in the melts (Righter, 2002, 2003; Righter et al., 2009; Rai and Van Westrenen, 2014). The high S content in the lunar core would result in lower modeled core formation temperatures and vice versa (Rai and van Westrenen, 2014; Steenstra et al., 2016a). The phosphorus partition coefficient will decrease with decreasing temperatures (Righter et al., 2010). It is also possible for an intermediate LMO. The metal-silicate equilibration in the Moon could also happen at the pressure of 3 GPa (Elkins-Tanton et al., 2011), corresponding to the pressure of a lunar magma ocean with a depth of ~650 km where deep moonquakes occur (>700 km, Nakamura et al. 1973). When we take the cosmochemical models and partition coefficient experimental models into account, the $D_{P-Moon}^{core/mantle}$ in this study is set at a range from 40 to 200. According to the Eq. (1), we plot the $C_{\text{core}}^{\text{P}}$ against the C_{BM}^{P} ranging from 50 to 300 ppm and different $D_{\text{P-Moon}}^{\text{core/mantle}}$ (40–200), as shown in the Fig. 8.

Previous estimates of $C_{\text{BSM}}^{\text{P}}$ and C_{BM}^{P} was 20 ppm and 43 ppm, respectively, which resulted in ~ 0.1 wt% P content in the lunar core (O'Neill, 1991). However, the phosphorus content in the bulk Moon (C_{BM}^{P}) is probably underestimated. More evidence from the elemental and isotopic studies indicates that the composition of the bulk Moon is more similar to the bulk silicate Earth (BSE) (Zhang et al., 2012; Dauphas et al., 2014; Taylor and Wieczorek, 2014). Therefore, using a BSE-like phosphorus concentration (82 ± 8 ppm, Wang et al., 2018) as the C_{BM}^{P} may be more appropriate for the initial silicate Moon. Such value would result in a slightly higher P content (C_{core}^{P} , 0.1-0.3 wt%) in the lunar core (Fig. 8). Moreover, the bulk silicate Moon enriched in crustal components model (the cBSM model) suggest that up to 0.04 wt% P₂O₅, equal to 174 ppm P, was present in the initial silicate Moon after the giant impact (Togashi et al., 2017). It would form a P-bearing core with 0.35-0.58 wt% phosphorus in the most optimistic scenario.

Here we used a moderate core mass (2.5 wt%) in the model to be consistent with previous studies (Kuskov and

Kronrod, 1998 2.5 \pm 2 wt%; Steenstra et al., 2016a 2.5 \pm 0.1 wt%). The smaller lunar core with \leq 1.5 wt% mass was also suggested by Williams et al. (2014) according to the geophysical measurements obtained by GRAIL. Lowering the core mass will lead to higher phosphorus concentration in the lunar core (Eq. (1)). In this case, the P core storage would be 0.45 wt% (core mass = 1.5 wt% of bulk Moon mass, $C_{BM}^{P} = 90$ ppm, and $D_{P-Moon}^{core/mantle} = 200$), larger than 0.30 wt% P with a 2.5 wt% core.

4.2. Phosphorus in a Fe-Ni-S-C lunar core

The thermal state of the lunar core has been discussed and described in different models depending on the geophysical and geochemical data. For the current temperature at the core-mantle boundary, there is a wide range from $\sim 1200^{\circ}$ C (Chacko and De Bremaecker, 1982; Ziethe et al., 2009; Scheinberg et al., 2015) to 1377 °C (Spohn et al., 2001), to as high as ~1500 °C (Zhang et al., 2013; Laneuville et al., 2014; Scheinberg et al., 2015). Pressure at the core-mantle boundary and inner core are speculated as 4.5 and 5.3 GPa, respectively (Kuskov and Kronrod, 1998; Garcia et al., 2011). To crystallize a solid iron core in the Moon, the upper limit to the core's temperature should be $\sim 1700^{\circ}$ C at 5.3 GPa, according to the iron phase diagram (Pepperhoff and Acet, 2001). As shown in the Fig. 5, the phosphorus solubility in the iron crystalline is ~ 0.7 wt% at 1500 °C and 3 GPa. At higher temperatures, T = 1600 °C, its solubility is slightly higher. Combined with the multiple effects from the temperatures and pressures on the phosphorus solubility in the iron, the maximum P content dissolved in the iron is assessed as 0.85 ± 0.15 wt% at the lunar core conditions (4.5–5.3 GPa and 1500 ~1700 °C).

The lunar core is assumed to have a complex chemical composition system such as Fe-Ni-S-C (Righter et al., 2017). Nickel is considered a major alloying element in the lunar core and has a wide concentration ranging from 9 to 30 wt% (Righter, 2002; Righter et al., 2017). In a scenario where the LMO equilibrated with the lunar core at pressures > 3 GPa, the Ni core storage will be 10–24 wt% (Steenstra et al., 2017). Carbon in the lunar core is reassessed to be ~ 1.2 wt% by Righter et al. (2017) but no more than 2.0 wt% according to the Fe-C phase diagram at 5 GPa (Fei and Brosh, 2014). Lunar core is also expected to have a higher C content range of $\sim 0.6-4.8$ wt%, given high C abundance in the BSM (Steenstra et al., 2017) which is also consistent with the high amount of C in the lunar mantle (Wetzel et al., 2015). As for bulk S content in the lunar core, there is still in debate whether the core is sulfur-rich (Weber et al., 2011 < 6 wt%) or sulfur-poor (Steenstra et al., 2017 < 0.16 wt%).

Experiments on the Fe-Ni-P system at ambient pressure suggest that nickel has a weak influence on the phosphorus solubility in γ -Fe (12 wt% Ni). Approximately 0.1 wt% P was reduced in γ -Fe-Ni at 1000, 900 and 800 °C (Kaneko et al., 1965). It is similar to the decrease of C content in the Fe-Ni-C melt at ambient pressure (Tsymbulov and Tsmekhman, 2001) and 2 GPa (Dasgupta and Walker,

2008). About 1 wt% carbon in the solid Fe-Ni(12 wt%)-P alloy could reduce 50% phosphorus concentration at 800-1000 °C by experiments (Kaneko et al., 1965). The C-bearing partitioning experiment shows that \sim 76% phosphorus is reduced in the liquid Fe-C metal at 5 GPa and 2000 °C (Jana and Walker, 1997). Sulfur in the iron-light element system may depress P concentration, similar to the sulfur depression on carbon in the Fe-Ni-S-C-P liquid in the case of high S content, 8-24 wt% (Tsuno and Dasgupta, 2015; Li et al., 2016). In the Fe-Ni-C-S system, the carbon concentration would be reduced about half in the melts with Xs = 0.1 (6 wt% sulfur) (Zhang et al., 2018). S mainly occurs as FeS with Fe under lunar core pressure conditions (Hsieh et al., 1987). If we assume that Fe₃C is similar to Fe₃P bound in the liquid, the addition of FeS bond will replace Fe₃P. It is easy to explain the increase of D_p^{SM/LM} along with increasing of S content (Fig. 7, Righter and Drake, 2000).

If we assume that lunar core is fully crystallized and the core is composed by solid iron and some light elements. The main light element should be carbon, with the concentration of ~1.2 wt% (Righter et al., 2017). Sulfur content should be less than $\sim 0.02 \text{ wt\%}$ due to the extremely lowsolubility in solid iron at lunar core pressures (Li et al., 2001). According to the Fe-P phase diagram in this study, the bulk phosphorus budget in the lunar core could be around 0.85 ± 0.15 wt% if phosphorus is saturated in the core. Considering the integrate influence from nickel and carbon, the phosphorus solubility in the core should be less than 0.4 ± 0.1 wt% ($0.5 \times P$ loss). It is a little close to the modeled bulk P storage in the core by core-mantle differentiation, using the BSE-like P content (82 ± 8 ppm, Wang et al., 2018) as C_{BM}^{P} (Fig. 8). However, because an inner core is expected for thermal-dynamical reasons (Wieczorek, 2006), and an estimated average core density is 5.2 ± 1.0 kg/cm³, lunar core model favors a liquid outer core (Garcia et al., 2011). About 60% volume of the core is liquid as implied by reanalyzes of Apollo-era seismic data (Weber et al., 2011). Righter et al. (2017) suggested the possibility of a C and S-poor core (0.5 wt% C, and 0.375 wt% S) which is consistent with all available geochemical and geophysical data and provides a simple heat source and mechanism for a lunar core dynamo. The inner core can be extensive 75-80% volume crystallized at 1300 °C with a relatively thin fluid outer core (Righter et al., 2017). Phosphorus content model for the lunar core (Fig. 8) indicates that phosphorus concentration must be less than ~ 0.3 wt % in the most optimistic scenario. Lunar core light element composition model suggested by Righter et al. (2017) results in 3.8-11.8 wt% S in the outer core. Following the trend of $D_{\rm p}^{\rm SM/LM}$ in Fig. 7, we speculate that the possible $D_{\rm p}^{\rm SM/LM}$ at the S-bearing lunar outer core *P*-*T* conditions is 0.18 ± 0.02 (Fig. 7). Using the core radius and density model proved by Antonangeli et al. (2015), the liquid outer core contains 49% of total core mass. Therefore, with the mass balance between solid inner core and liquid outer core, the P concentration is $0.09\pm0.01~\text{wt\%}$ in inner core and 0.51 ± 0.01 wt% in outer core ($C_{core}^{P} = 0.3$ wt%).

5. CONCLUSIONS

This work provides particular information of Fe-P binary phase diagram at 3 GPa and up to 1600 °C. The low-P iron observed in this study gives a new insight to constrain the phosphorus budget in the lunar core. With increasing temperature, the P solubility in solid iron decreases to ~0.7 wt% at 1500 °C and 3 GPa. The partition coefficient model for phosphorus between the metal and silicate melt is a function of pressure, temperature, and oxygen fugacity and has been well investigated at lunar core conditions (Righter et al., 2009, 2010; Steenstra et al., 2016a, 2016b, 2017). But the P budget in the lunar core relies on the P content of bulk Moon. Assuming the C_{BM}^{P} is like the P content of bulk silicate Earth, it would result in less than 0.3 wt % P in the lunar core. This relative low concentration is in agreement with an S- and C-poor core model suggested by Righter et al. (2017). Because of the complex composition model (Fe-Ni-C-S) for lunar core, the C_{core}^{P} should be smaller than the examined P solubility in the iron in this study. In the case of a full-crystallized core, the P core storage is expected less than $\sim 0.4 \text{ wt\%}$ due to the negative effect caused by nickel and carbon. The $D_p^{SM/LM}$ is also determined as 0.10 ± 0.03 at 3 GPa and 1250–1550 °C in the Fe-P binary system with S-free. If the sulfur is taken into account, $D_{\rm p}^{\rm SM/LM}$ increases to 0.18 \pm 0.02. In a scenario of solid inner core and liquid outer core, the P concentration is estimated as 0.09 ± 0.01 wt% in inner core and 0.51 \pm 0.01 wt% in outer core. To be compatible with the lunar mantle or bulk silicate Moon abundance of P, the core should be P-poor although the phosphorus has high solubility ($\sim 10 \text{ wt\%}$) in the liquid metal. Future works are needed to consider the maximum solubility of P in a complex system at lunar core conditions. We also note that titanium affects Fe-P bond in γ -iron (Kaneko et al., 1965), and there is perhaps an ilmenite-rich layer beyond lunar outer core (de Vries et al., 2010; Matsumoto et al., 2015). It is necessary to investigate the effect of titanium on the phosphorus partitioning in the lunar magma ocean.

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APPENDIX A. SUPPLEMENTARY MATERIAL

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