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Iron isotope fractionation in planetary crusts

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Abstract

We present new high precision iron isotope data ($\delta^{56}\text{Fe}$ vs. IRMM-014 in per mil) for four groups of achondrites: one lunar meteorite, 11 martian meteorites, 32 howardite–eucrite–diogenite meteorites (HEDs), and eight angrites. Angrite meteorites are the only planetary materials, other than Earth/Moon system, significantly enriched in the heavy isotopes of Fe compared to chondrites (by an average of $+0.12\text{‰}$ in $\delta^{56}\text{Fe}$). While the reason for such fractionation is not completely understood, it might be related to isotopic fractionation by volatilization during accretion or more likely magmatic differentiation in the angrite parent-body. We also report precise data on martian and HED meteorites, yielding an average $\delta^{56}\text{Fe}$ of $0.00 \pm 0.01\text{‰}$. Stannern-trend eucrites are isotopically heavier by $+0.05\text{‰}$ in $\delta^{56}\text{Fe}$ than other eucrites. We show that this difference can be ascribed to the enrichment of heavy iron isotopes in ilmenite during igneous differentiation. Preferential dissolution of isotopically heavy ilmenite during remelting of eucritic crust could have generated the heavy iron isotope composition of Stannern-trend eucrites. This supports the view that Stannern-trend eucrites are derived from main-group eucrite source magma by assimilation of previously formed asteroidal crust.

These new results show that iron isotopes are not only fractionated in terrestrial and lunar basalts, but also in two other differentiated planetary crusts. We suggest that igneous processes might be responsible for the iron isotope variations documented in planetary crusts.

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1. INTRODUCTION

Iron is the ninth most abundant element in the Solar System, and the second most abundant element in the Earth and other terrestrial planetary bodies. Iron can behave as a siderophile (metal-loving), lithophile (rock-loving) and chalcophile (sulfur-loving) element and is ubiquitous in Solar System planetary bodies. On Earth, it is a major element in the core, the mantle and the crust (Allègre et al., 1995; Rudnick and Gao, 2003). In the past decade, the development of high-resolution Multi-Collector Inductively-

Coupled-Plasma Mass-Spectrometers (MC-ICP-MS) has allowed measurements of iron isotope composition at high precision (Belshaw et al., 2000; Zhu et al., 2001; Weyer and Schwiders, 2003; Dauphas et al., 2009b; Millet et al., 2012). Following this improvement, small yet resolvable iron isotopic variations in igneous rocks have been discovered (Poitrasson et al., 2004; Weyer et al., 2005; Schoenberg and von Blanckenburg, 2006; Weyer and Ionov, 2007; Heimann et al., 2008; Teng et al., 2008, 2011; Dauphas et al., 2009a; Weyer and Seitz, 2012).

Mid-ocean ridge basalts (MORBs), oceanic island basalts (OIBs), and continental basalts are enriched in heavy iron isotopes by $\sim +0.1\text{‰}$ when compared to chondrites; while martian and 4-Vesta (HED) meteorites all have

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similar iron isotope compositions to chondrites (Poitrasson et al., 2004; Weyer et al., 2005; Anand et al., 2006; Schoenberg and von Blanckenburg, 2006; Craddock and Dauphas, 2011). Three mechanisms have been proposed to explain this difference: (1) *Planetary accretion*: Heavy iron isotopes were enriched on the Earth and Moon by evaporative kinetic isotope fractionation during the giant impact that formed the Moon (Poitrasson et al., 2004). In this context, one of the difficulties is to explain why potassium, which is highly volatile, is not isotopically fractionated in the Earth–Moon system (Humayun and Clayton, 1995). Poitrasson et al. (2004) proposed that Fe was evaporated from core material as metal while K was evaporated as oxides. In that case, Fe would be more volatile than K and could be isotopically more fractionated. However, lunar mare basalts (low-Ti vs. high-Ti) have large variable iron isotope compositions and the $\delta^{56}\text{Fe}$ value of the bulk Moon is not well known (Liu et al., 2010), which provides little constraint on the evaporative isotope fractionation hypothesis. (2) *Core–mantle segregation*: At ultra high-pressures (>100 GPa) relevant to terrestrial core–mantle boundary conditions, Polyakov (2009) suggested based on nuclear resonant inelastic X-ray scattering data that a detectable fractionation between metallic and silicate phases should be present, which could explain the heavy iron isotope composition of terrestrial silicate rocks. While the approach used by Polyakov (2009) is sound, the results are highly dependent on the high-energy tails of the phonon density of states, which are highly uncertain for high-pressure minerals. In contrast to this hypothesis, laboratory experiments at temperatures and pressures applicable to differentiation of parent bodies to achondrites have shown no iron isotope fractionation between metal and silicate minerals (Poitrasson et al., 2009; Hin et al., 2010). (3) *Crust formation*: It has been widely observed that iron isotopes could be fractionated during various magmatic differentiation processes of the terrestrial crust, such as partial melting, mineral frac-

tionation and fluids exsolution (Williams et al., 2004; Poitrasson and Freyrier, 2005; Weyer and Ionov, 2007; Teng et al., 2008; Schuessler et al., 2009). Island arc basalts also show iron isotope fractionation that may be related to the degree of partial melting (Dauphas et al., 2009a). During partial melting, Fe(III) is more incompatible in olivine and pyroxene than Fe(II) is. Both theoretical calculations and experimental determinations show that Fe(III)-bearing phases tend to be enriched in the heavy isotopes of iron compared to Fe(II)-bearing phases (Polyakov and Mineev, 2000; Schauble et al., 2001; Schuessler et al., 2007; Shahar et al., 2008). Such equilibrium isotope fractionation between Fe(III) and Fe(II) may explain, at least in part, the heavy iron isotope composition of MORBs and OIBs relative to that of chondrites and other planetary basalts (Dauphas et al., 2009a). Indeed, terrestrial basalts are formed under more oxidizing conditions than martian meteorites or HEDs (McCammon, 2005; Wadhwa, 2008).

Achondrites are samples from differentiated planetary bodies formed under a variety of conditions. Here, we report high-precision iron isotope compositions of several classes of achondrites from at least four different parent bodies, including a lunar meteorite, martian meteorites, howardite–eucrite–diogenite meteorites (HEDs) and angrites, and discuss the implications regarding the conditions relevant to the differentiation of the parent-bodies of these meteorites.

2. SAMPLES AND METHODS

2.1. Sample descriptions

All samples, their classifications and weathering conditions (if available) are listed in Table 1. Well-characterized terrestrial geostandards were analyzed to assess the quality of the measurements and to provide a basis for inter-laboratory comparisons. BCR-2 is a continental flood basalt

Table 1
Iron isotope compositions of Earth/Moon rocks and achondrites.

Sample	Type	Fall/find (weathering)	Chemistry	Fe wt. %	$\delta^{56}\text{Fe}$	95% c.i. ^a	$\delta^{57}\text{Fe}$	95% c.i. ^a	<i>n</i>
<i>Earth/Moon</i>									
AGV-1 #1	Andesite		Routine ^b		0.081	0.034	0.133	0.049	8
AGV-1 #2	Andesite		Routine ^b		0.077	0.045	0.105	0.058	8
AGV-1 #3	Andesite		Routine ^b		0.081	0.046	0.134	0.066	9
AGV-1 #4	Andesite		Routine ^c		0.106	0.031	0.157	0.039	11
AGV-1 (average)					0.089	0.019	0.138	0.025	
AGV-2 #1	Andesite		Routine ^b	4.6	0.094	0.038	0.137	0.057	9
AGV-2 #2	Andesite		Routine ^b		0.083	0.035	0.094	0.057	8
AGV-2 #3	Andesite		Routine ^c		0.100	0.031	0.167	0.040	10
AGV-2 (average)					0.093	0.020	0.141	0.028	
BCR-2 #1	Basalt		Routine ^b	11.0	0.063	0.032	0.080	0.046	9
BCR-2 #2	Basalt		Routine ^b	11.0	0.052	0.033	0.110	0.058	9
BCR-2 #3	Basalt		Routine ^c		0.089	0.031	0.110	0.039	11
BCR-2 (average)					0.069	0.018	0.100	0.026	
BHVO-2 #1	Basalt		Routine ^c		0.116	0.038	0.164	0.057	4

Table 1 (continued)

Sample	Type	Fall/find (weathering)	Chemistry	Fe wt.%	$\delta^{56}\text{Fe}$	95% c.i. ^a	$\delta^{57}\text{Fe}$	95% c.i. ^a	<i>n</i>
BHVO-2 #2-1	Basalt		Routine ^d	6.3	0.099	0.030	0.155	0.044	9
BHVO-2 #2-2	Basalt		Long ^d	7.9	0.096	0.030	0.160	0.053	9
BHVO-2 #2-3	Basalt		UTEVA ^d	8.7	0.101	0.030	0.166	0.044	9
BHVO-2 #3-1 (+V + Cr) ^j	Basalt		Routine ^d	6.7	0.108	0.030	0.161	0.044	9
BHVO-2 #3-2 (+V + Cr) ^j	Basalt		Long ^d	8.5	0.098	0.030	0.154	0.044	9
BHVO-2 #3-3 (+V + Cr) ^j	Basalt		UTEVA ^d	9.0	0.098	0.030	0.154	0.044	9
BHVO-2 (average)					0.102	0.012	0.159	0.018	
BIR-1 #1	Basalt		Routine ^b		0.044	0.030	0.070	0.049	9
BIR-1 #2	Basalt		Routine ^b	7.7	0.038	0.038	0.047	0.075	8
BIR-1 #3	Basalt		Routine ^b	7.7	0.020	0.032	0.019	0.059	8
BIR-1 #4	Basalt		Routine ^c		0.068	0.031	0.070	0.040	10
BIR-1 (average)					0.043	0.016	0.058	0.026	
NMNH 116852-1 #1-1	Basalt		Routine ^d	3.7	-0.148	0.035	-0.223	0.050	9
NMNH 116852-1 #1-2	Basalt		UTEVA ^d	6.4	-0.081	0.035	-0.134	0.050	9
NMNH 116852-1 (average)					-0.115	0.025	-0.179	0.035	
NMNH 116852-3 #1-1	Basalt		Routine ^d	5.6	0.061	0.027	0.096	0.042	9
NMNH 116852-3 #1-2	Basalt		Long ^d	7.6	0.062	0.027	0.088	0.042	9
NMNH 116852-3 #1-3	Basalt		UTEVA ^d	7.7	0.052	0.027	0.094	0.042	9
NMNH 116852-3 (average)					0.058	0.016	0.093	0.024	
NMNH 116852-11 #1-1	Basalt		Routine ^d	5.5	0.134	0.027	0.190	0.035	9
NMNH 116852-11 #1-2	Basalt		Long ^d	7.1	0.133	0.027	0.200	0.035	9
NMNH 116852-11 #1-3	Basalt		UTEVA ^d	7.3	0.121	0.027	0.169	0.035	9
NMNH 116852-11 (average)					0.129	0.016	0.186	0.020	
GSP-1 #1	Granodiorite		Routine ^b		0.126	0.038	0.174	0.057	9
GSP-1 #2	Granodiorite		Routine ^b		0.141	0.046	0.204	0.066	9
GSP-1 #3	Granodiorite		Routine ^c		0.156	0.031	0.170	0.040	10
GSP-1 (average)					0.143	0.021	0.178	0.029	
AC-E #1-1	Granite		Routine ^d	1.2	0.314	0.030	0.452	0.044	9
AC-E #1-2	Granite		Long ^d	1.6	0.330	0.033	0.480	0.063	9
AC-E #1-3	Granite		UTEVA ^d	1.7	0.298	0.030	0.452	0.044	9
AC-E (average)					0.313	0.018	0.457	0.028	
MAC88105	Lunar meteorite	Find (A/Be) ⁱ	Routine ^b	3.4 ^h	0.086	0.034	0.132	0.049	8
<i>Martian meteorites</i>									
ALHA77005	Shergottite	Find (A) ⁱ	Routine ^b	15.6 ^h	-0.008	0.033	-0.026	0.058	9
EETA79001	Shergottite	Find (A/Ae) ⁱ	Routine ^b	14.3 ^h	-0.003	0.038	-0.038	0.075	8
Los Angeles	Shergottite	Find	Routine ^b	16.6 ^h	0.008	0.032	0.010	0.046	9
NWA1669	Shergottite	Find	Routine ^b		-0.016	0.032	-0.030	0.046	9
NWA1950	Shergottite	Find	Routine ^b	16.8 ^h	0.010	0.032	0.023	0.046	9
NWA5029	Shergottite	Find	Routine ^b		0.016	0.032	0.012	0.046	9
SaU008	Shergottite	Find	Routine ^b		-0.023	0.038	-0.042	0.075	8
Zagami	Shergottite	Fall	Routine ^b	16.0	0.016	0.031	0.064	0.055	9
	<i>Wt. Ave. of Shergottite</i>				0.001	0.012	0.002	0.019	
Mil03346	Nakhlite	Find (B) ⁱ	Routine ^b	14.8 ^h	0.025	0.032	0.027	0.046	9
Nakhla	Nakhlite	Fall	Routine ^b	16.0 ^h	0.031	0.032	0.033	0.046	9
	<i>Wt. Ave. of Nakhlite</i>				0.028	0.023	0.030	0.033	
ALH84001	Orthopyroxenite	Find (A/B) ⁱ	Routine ^b	13.6 ^h	-0.033	0.032	-0.034	0.046	9
	<i>Wt. Ave. of Mars</i>				0.003	0.010	0.004	0.015	
<i>HED</i>									
ALHA77256	Diogenite	Find (A/B) ⁱ	Routine ^c	14.2 ^h	-0.008	0.034	0.013	0.049	6
GRO95555	Diogenite	Find (A/B) ⁱ	Routine ^b	12.7 ^h	0.019	0.034	0.042	0.049	8
MET00424	Diogenite	Find (B) ⁱ	Routine ^c	15.8 ^h	-0.034	0.034	-0.074	0.049	6
MET00436	Diogenite	Find (B/C) ⁱ	Routine ^c	16.5 ^h	-0.032	0.032	-0.077	0.045	8

(continued on next page)

Table 1 (continued)

Sample	Type	Fall/find (weathering)	Chemistry	Fe wt.%	$\delta^{56}\text{Fe}$	95% c.i. ^a	$\delta^{57}\text{Fe}$	95% c.i. ^a	<i>n</i>
Mil07001	Diogenite	Find (A/B) ⁱ	Routine ^c		−0.017	0.032	−0.036	0.045	8
NWA1461	Diogenite	Find	Routine ^c	7.5 ^h	0.027	0.038	0.009	0.057	4
NWA5480	Diogenite	Find	Routine ^b		0.015	0.034	0.040	0.049	8
Shalka	Diogenite	Fall	Routine ^c	12.7 ^h	−0.011	0.038	−0.034	0.057	4
Tatahouine	Diogenite	Fall	Routine ^b	9.0	0.013	0.030	0.036	0.049	9
	<i>Wt. Ave. of Diogenite</i>					−0.004	0.011	−0.011	0.016
Frankfort	Howardite	Fall	Routine ^b	12.4	−0.002	0.038	−0.015	0.057	9
Kapoeta	Howardite	Fall	Routine ^b	8.7	0.026	0.030	0.042	0.049	9
Petersburg	Howardite/polymict eucrite	Fall	Routine ^b	8.3	0.035	0.038	0.056	0.057	9
	<i>Wt. Ave. of Howardite</i>					0.021	0.020	0.029	0.031
NWA1240	Eucrite (Ungrouped)	Find	Routine ^d	15.7 ^h	0.016	0.031	−0.026	0.043	6
Pasamonte	Eucrite (Ungrouped)	Fall	Routine ^b	11.5	0.019	0.030	0.063	0.045	8
EET87548 #1	Eucrite (Cumulate)	Find (B/C) ⁱ	Routine ^d	15.1	−0.023	0.034	−0.029	0.051	9
EET87548 #2	Eucrite (Cumulate)	Find (B/C) ⁱ	Routine ^b	15.4	−0.017	0.034	−0.030	0.049	8
EET87548 #3	Eucrite (Cumulate)	Find (B/C) ⁱ	Routine ^d		−0.014	0.030	−0.024	0.059	9
EET87548 (average)					−0.018	0.019	−0.028	0.030	
Moore County #1	Eucrite (Cumulate)	Fall	Routine ^b		−0.004	0.038	−0.008	0.057	9
Moore County #2	Eucrite (Cumulate)	Fall	Routine ^b	7.4	0.010	0.030	0.040	0.049	9
Moore County (average)					0.005	0.024	0.020	0.037	
Serra de Magé #1	Eucrite (Cumulate)	Fall	Routine ^d	12.3	0.001	0.034	0.026	0.051	9
Serra de Magé #2-1	Eucrite (Cumulate)	Fall	Routine ^d	12.0	0.017	0.028	0.025	0.039	9
Serra de Magé #2-2	Eucrite (Cumulate)	Fall	Long ^d	16.1	0.019	0.028	0.032	0.039	9
Serra de Magé #2-3	Eucrite (Cumulate)	Fall	UTEVA ^d	16.6	0.006	0.028	0.009	0.039	9
Serra de Magé (average)					0.012	0.015	0.023	0.021	
	<i>Wt. Ave. of Eucrite (Cumulate)</i>					0.001	0.010	0.009	0.015
Cachari	Eucrite (MG) ^e	Find	Routine ^b	15.9 ^h	0.002	0.030	0.011	0.045	8
Camel Donga	Eucrite (MG) ^e	Find	Routine ^d	15.3	0.019	0.034	0.034	0.051	9
Jonzac #1	Eucrite (MG) ^e	Find	Routine ^b	15.0	0.025	0.035	0.031	0.057	8
Jonzac #2	Eucrite (MG) ^e	Fall	Routine ^b	11.5	0.007	0.034	0.051	0.049	8
Jonzac #3-1	Eucrite (MG) ^e	Fall	Routine ^d	9.8	0.008	0.032	0.028	0.045	9
Jonzac #3-2	Eucrite (MG) ^e	Fall	Long ^d	14.8	−0.006	0.032	−0.007	0.045	9
Jonzac #3-3	Eucrite (MG) ^e	Fall	UTEVA ^d	14.6	0.010	0.032	0.035	0.045	9
Jonzac (average)					0.008	0.015	0.026	0.021	
Juvinas #1	Eucrite (MG) ^e	Fall	Routine ^b		0.015	0.038	0.035	0.057	9
Juvinas #2-1	Eucrite (MG) ^e	Fall	Routine ^d	9.9	−0.025	0.028	−0.019	0.040	9
Juvinas #2-2	Eucrite (MG) ^e	Fall	Long ^d	14.0	−0.022	0.028	−0.024	0.040	9
Juvinas #2-3	Eucrite (MG) ^e	Fall	UTEVA ^d	13.6	−0.017	0.028	−0.021	0.040	9
Juvinas #3-1	Eucrite (MG) ^e	Fall	Routine ^d	9.9	−0.026	0.035	−0.038	0.050	9
Juvinas #3-2	Eucrite (MG) ^e	Fall	Long ^d	14.0	−0.023	0.035	−0.036	0.050	9
Juvinas #3-3	Eucrite (MG) ^e	Fall	UTEVA ^d	13.6	−0.037	0.035	−0.059	0.050	9
Juvinas (average)					−0.020	0.012	−0.024	0.017	
NWA049	Eucrite (MG) ^e	Find	Routine ^b	15.4 ^h	0.023	0.029	0.033	0.047	9
	<i>Wt. Ave. of Eucrite (MG)^e</i>					−0.004	0.008	0.001	0.012
Agoult	Eucrite (Residual)	Find	Routine ^b	15.2 ^h	0.033	0.034	0.039	0.049	8
Dag945	Eucrite (Residual)	Find (W1) ⁱ	Routine ^c	15.6 ^h	0.028	0.036	0.037	0.052	5
	<i>Wt. Ave. of Eucrite (Residual)</i>					0.031	0.025	0.038	0.036
Nuevo Laredo	Eucrite (NL) ^f	Find	Routine ^c	16.2 ^h	0.031	0.036	0.056	0.052	5
Sahara02501	Eucrite (NL) ^f	Find	Routine ^c	16.3	0.037	0.029	0.053	0.049	8
	<i>Wt. Ave. of Eucrite (NL)^f</i>					0.035	0.023	0.054	0.036
Bouvante #1	Eucrite (ST) ^g	Find	Routine ^d	15.1	0.049	0.034	0.096	0.051	9
Bouvante #2-1	Eucrite (ST) ^g	Find	Routine ^d	10.7	0.032	0.028	0.036	0.036	9
Bouvante #2-2	Eucrite (ST) ^g	Find	Long ^d	16.6	0.038	0.028	0.054	0.036	9
Bouvante #2-3	Eucrite (ST) ^g	Find	UTEVA ^d	15.9	0.026	0.028	0.054	0.036	9
Bouvante (average)					0.035	0.015	0.055	0.019	
NWA2061	Eucrite (ST) ^g	Find	Routine ^b	14.4	0.018	0.029	0.031	0.047	9
NWA4523 #1	Eucrite (ST) ^g	Find	Routine ^b	13.3 ^h	0.077	0.034	0.118	0.049	8
NWA4523 #2	Eucrite (ST) ^g	Find	Routine ^d		0.071	0.034	0.097	0.051	9

Table 1 (continued)

Sample	Type	Fall/find (weathering)	Chemistry	Fe wt. %	$\delta^{56}\text{Fe}$	95% c.i. ^a	$\delta^{57}\text{Fe}$	95% c.i. ^a	<i>n</i>
NWA4523 #3-1	Eucrite (ST) ^g	Find	Routine ^d		0.056	0.027	0.071	0.039	9
NWA4523 #3-2	Eucrite (ST) ^g	Find	Long ^d		0.052	0.027	0.095	0.039	9
NWA4523 #3-3	Eucrite (ST) ^g	Find	UTEVA ^d		0.060	0.027	0.095	0.039	9
NWA4523 (average)					0.061	0.013	0.093	0.019	
Pomozdino	Eucrite (ST) ^g	Find	Routine ^d	15.2	0.034	0.034	0.060	0.051	9
Stannern #1	Eucrite (ST) ^g	Fall	Routine ^b	14.2	0.072	0.035	0.095	0.057	8
Stannern #2	Eucrite (ST) ^g	Fall	Routine ^d	13.7	0.036	0.045	0.079	0.065	9
Stannern #3	Eucrite (ST) ^g	Fall	Routine ^d		0.049	0.045	0.102	0.065	9
Stannern #4	Eucrite (ST) ^g	Fall	Routine ^d	14.5	0.053	0.045	0.071	0.065	9
Stannern #5-1	Eucrite (ST) ^g	Fall	Routine ^d	12.0	0.047	0.026	0.078	0.032	9
Stannern #5-2	Eucrite (ST) ^g	Fall	Long ^d	12.5	0.035	0.026	0.073	0.032	9
Stannern #5-3	Eucrite (ST) ^g	Fall	UTEVA ^d	15.6	0.037	0.026	0.059	0.032	9
Stannern (average)					0.045	0.012	0.074	0.016	
Yamato75011	Eucrite (ST) ^g <i>Wt. Ave. of Eucrite (ST)^g</i>	Find (A) ⁱ	Routine ^b	15.2	0.043	0.029	0.092	0.047	9
					0.045	0.007	0.073	0.010	
<i>Angrite</i>									
D'Orbigny	Angrite	Find	Routine ^d	19.2 ^h	0.139	0.030	0.193	0.062	9
LEW86010	Angrite	Find(A/B) ⁱ	Routine ^b	14.2 ^h	0.139	0.043	0.179	0.059	9
NWA1670 #1	Angrite	Find	Routine ^b	14.4 ^h	0.167	0.033	0.253	0.058	9
NWA1670 #2	Angrite	Find	Routine ^d		0.125	0.036	0.210	0.058	9
NWA1670 #3-1	Angrite	Find	Routine ^d		0.148	0.030	0.225	0.038	9
NWA1670 #3-2	Angrite	Find	Long ^d		0.113	0.030	0.172	0.038	9
NWA1670 #3-3	Angrite	Find	UTEVA ^d		0.154	0.030	0.223	0.038	9
NWA1670 (average)					0.142	0.014	0.212	0.019	
NWA1296	Angrite	Find	Routine ^b	19.4 ^h	0.104	0.033	0.185	0.058	9
NWA6291	Angrite	Find	Routine ^d		0.052	0.030	0.116	0.062	9
NWA2999	Angrite	Find	Routine ^d	24.2 ^h	0.124	0.030	0.183	0.062	9
NWA4801	Angrite	Find	Routine ^d		0.135	0.030	0.219	0.062	9
Sahara99555 #1	Angrite	Find	Routine ^d	18.0 ^h	0.116	0.040	0.165	0.081	9
Sahara99555 #2	Angrite	Find	Routine ^b		0.107	0.033	0.136	0.058	9
Sahara99555 #3	Angrite	Find	Routine ^d		0.110	0.030	0.141	0.062	9
Sahara99555 (average)					0.110	0.019	0.144	0.038	
	<i>Wt. Ave. of Angrite</i>				0.123	0.009	0.192	0.014	

^a Uncertainties are 95% confidence intervals (see Dauphas et al. (2009b) for details).

^b Samples chemically prepared at the Washington University in St. Louis and measured at the University of Chicago.

^c Samples both chemically prepared and measured at Washington University in St. Louis.

^d Samples both chemically prepared and measured at the University of Chicago.

^e Eucrite (MG) = Eucrite main group.

^f Eucrite (NL) = Eucrite Nuevo Laredo Trend.

^g Eucrite (ST) = Eucrite Stannern Trend.

^h Reference data: MAC88105 (Jarosewich, 1990); NWA1950 (Gillet et al., 2005); Los Angeles (Warren et al., 2000); EETA79001, ALHA77005, Nakhla, ALH84001 (Lodders, 1998); Mil03346 (Day et al., 2006); GRO95555, MET00436, MET00424 (Barrat et al., 2008); NWA1461 (Warren et al., 2009); Shalka (McCarthy et al., 1972); ALHA77256 (Sack et al., 1991); NWA1240, NWA049 (Barrat et al., 2003); Cachari (Barrat et al., 2000); Agoult, Dag945 (Yamaguchi et al., 2009); Nuevo Laredo, NWA4523 (Barrat et al., 2007b); D'Orbigny, Sahara99555 (Mittlefehldt et al., 2002); LEW86010 (McKay et al., 1988); NWA1670 (Jambon et al., 2008); NWA1296 (Jambon et al., 2005); NWA2999 (Gellissen et al., 2007).

ⁱ Weathering conditions are from MetBase (version 7.1). Weathering index A, B or C represents "minor", "moderate" or "severe" rustiness, respectively. Letter "e" represents evaporite minerals visible. Weathering grade W0 (fresh) to W6 (most weathered) is also shown if available.

^j These samples have been added V and Cr to double-check the purification of iron during chemistry.

from the Colombia River, Oregon. BIR-1 is a basalt from Iceland. BHVO-2 is a Hawaiian lava basalt. AGV-1 and 2 are andesites from the Guano Valley, Oregon. GSP-1 is a granodiorite from Silver Plume, Colorado. AC-E is a granite from Ailsa Craig Island, Scotland. In addition,

three modern island arc basalts (IABs) from New Britain (NMNH 116852-1, 116852-3 and 116852-11) were also analyzed here to test the accuracy of the measurements (see Dauphas et al. (2009a) for a detailed discussion about IABs).

Lunar highland meteorite, MAC88105, is a polymict breccia. It is dominated by ferroan anorthosite lithology and contains small low-Ti basaltic clasts (Jolliff et al., 1991; Neal et al., 1991). The bulk composition of this lunar highland meteorites is different from those of highland rocks sampled by Apollo project (Koeberl et al., 1991; Lindstrom et al., 1991) and might represent a different feldspathic highlands terrane (Warren et al., 1989). Orbital remote sensing composition data shows that lunar highland meteorites are reasonably representative of lunar surface (Korotev et al., 2003).

The martian meteorites studied include eight shergottites, two nakhlites and ALH84001. Shergottites are basaltic or lherzolitic rocks. Basaltic shergottites (e.g., Zagami, EETA79001) mainly consist of clinopyroxene and plagioclase (Mittlefehldt et al., 1998). Lherzolitic shergottites (e.g., ALHA77005) are cumulates, consisting of magnesian olivine, clinopyroxene and chromite (Mittlefehldt et al., 1998). Nakhlites are clinopyroxenites made of augite and a small amount of olivine (Mittlefehldt et al., 1998). ALH84001 is a cumulate orthopyroxenite.

The HED meteorites reported in this study include 20 eucrites, three howardites and nine diogenites. Based on spectroscopic observations, HED meteorites are generally recognized as being derived from asteroid 4-Vesta (McCord et al., 1970; Drake, 2001). Diogenites are orthopyroxenites, consisting chiefly of ~90 vol % coarse-grained orthopyroxene, and accessory minerals including olivine, chromite, troilite, and metal (Mittlefehldt et al., 1998). Eucrites are basalts (non-cumulates) or cumulate gabbros. Basaltic eucrites contain pigeonite, plagioclase and a minor amount of silica, ilmenite and chromite (Duke and Silver, 1967). Basaltic eucrites can be subdivided based on geochemical characteristics: “Main Group” (MG), “Nuevo Laredo Trend” (NL), and “Stannern Trend” (ST) (Stolper, 1977; Reid and Barnard, 1979; Yamaguchi et al., 2009). These three groups are very similar in major element compositions, however they are different in trace element abundances. The Stannern Trend shows a significant increase in incompatible trace elements without changes in Mg# compared to the Main Group; while the Nuevo

Laredo Trend shows a slight increase in incompatible trace elements and a decrease in Mg# compared to the Main Group. Cumulate eucrites have mineral assemblages similar to basaltic eucrites, but they display cumulate textures and their pyroxenes are Mg-rich. Howardites are polymict breccias and are mixtures primarily of diogenite and eucrite clasts. In addition, we have analyzed Pasamonte and NWA1240, two ungrouped eucrites recently found to have oxygen isotope compositions distinct from all other HEDs (Scott et al., 2009). Pasamonte is a polymict breccia that displays highly unequilibrated basaltic clasts (Takeda et al., 1978), and possibly records fluid–rock interactions (Schwartz and McCallum, 2005). NWA1240 is an unbrecciated stone, with unique features. It displays a porphyritic texture consisting of skeletal hollow low-Ca pyroxene phenocrysts set in a variolitic (fan-spherulitic) mesostasis of fine elongate pyroxene and plagioclase crystals. Pyroxenes are highly unequilibrated and their compositions range from En₇₆ to En_{0.6}. Although its texture indicates a rock formed from a rapidly-cooled melt, its composition is more akin to cumulate eucrites and led Barrat et al. (2003) to propose an impact-melt origin. Nevertheless, these ungrouped eucrites are probably not from asteroid 4-Vesta, but from Vesta-like asteroid parent-bodies (Scott et al., 2009).

Angrites are among the oldest basaltic rocks in the Solar System (Baker et al., 2005; Nyquist et al., 2009; Dauphas and Chaussidon, 2011). Only 19 angrites are recognized. Eight representative ones were analyzed here: D’Orbigny, NWA1296 and Sahara99555 consist of mainly Al–Ti-diopside–hedenbergite, Ca-rich olivine, anorthite and spinel olivine; they are also similar in bulk chemistry (Jambon et al., 2005); LEW86010 consists of ~58 vol % Al–Ti-diopside–hedenbergite, 21 vol % plagioclase, 20 vol % olivine and minor amount of spinel, troilite, and Fe–Ni metal (Prinz et al., 1988); NWA1670 consists of large olivine xenocrysts, and fine-grained groundmass with pyroxene, anorthite and olivine (Jambon et al., 2008); NWA4801 is composed of Al–Ti-diopside–hedenbergite and pure anorthite (Irving and Kuehner, 2007); NWA2999 contains 64 vol % Ca-rich olivine, 23 vol % Al–Ti-diopside–hedenbergite, 4 vol % spinel, 1 vol % plagioclase and 8 vol % metal (Kuehner et al.,

Table 2

Comparison of the three iron purification chromatography protocols used in this study (see text for references).

	Routine column	Long column	UTEVA column
Column	Disposable Bio-rad poly prep polyethylene column (9 cm length)	Reusable Savillex PFA column (10.5 cm length, 0.62 cm diameter)	Pre-packaged Eichrom U/Teva cartridges
Resin	1 mL AG1-X8 200–400 mesh anion-exchange resin	3 mL AG1-X8 200–400 mesh anion-exchange resin	1 mL Eichrom U/Teva resin
Conditioning	10 mL 6 M HCl	4 mL 10 M HCl	6 mL 4 M HCl
Matrix elution	8 mL 6 M HCl (0.5 + 0.5 + 1 + 2 + 4 mL increments)	5 mL 10 M HCl (0.5 + 0.5 + 1 + 3 mL increments) 22 mL 4 M HCl (0.5 + 0.5 + 1 + 5 + 5 + 10 mL increments)	25 mL 4 M HCl (0.5 + 0.5 + 1 + 5 + 8 + 10 mL increments)
Iron collecting	9 mL 0.4 M HCl (0.5 + 0.5 + 1 + 3 + 4 mL increments)	8 mL 0.4 M HCl (0.5 + 0.5 + 1 + 2 + 4 mL increments)	12 mL 0.4 M HCl (0.5 + 0.5 + 1 + 5 + 5 mL increments)

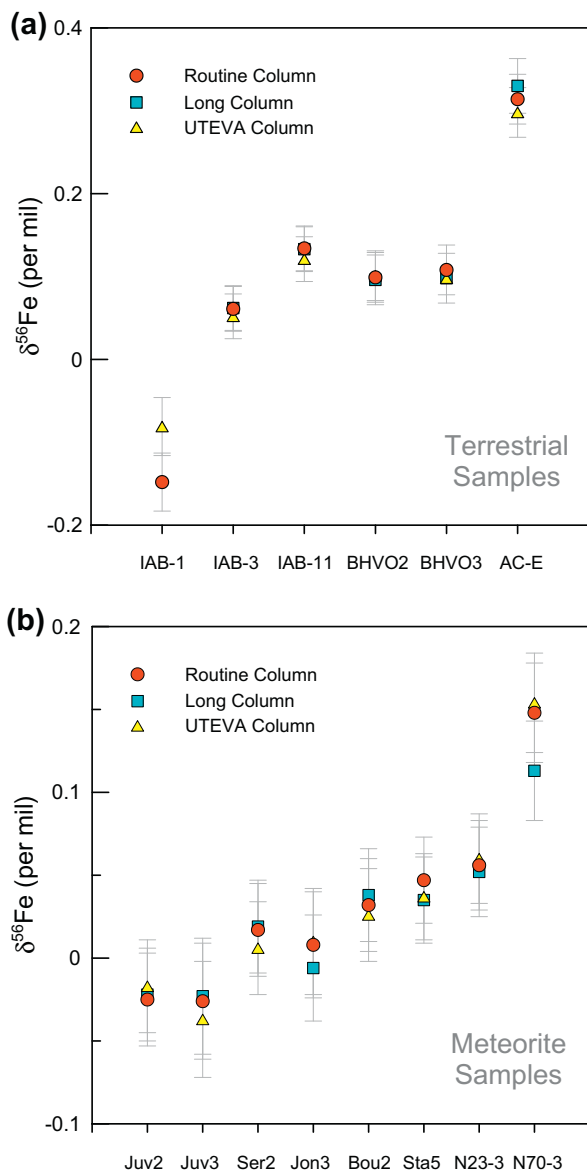


Fig. 1. Iron isotope results of the samples that were purified independently by three different purification chemistries, namely “Routine Columns”, “Long Columns” and “UTEVA Columns”. For a detailed description of these purification chemistry protocols, please see Table 2 and text. Top panel: terrestrial samples. IAB-1 = NMNH 116852-1, IAB-3 = NMNH 116852-3, IAB-11 = NMNH 116852-11, BHVO2 = BHVO-2 #2, BHVO3 = BHVO-2 #3, AC-E = AC-E #1. Bottom panel: meteorite samples. Juv2 = Juvinas #2, Juv3 = Juvinas #3, Ser2 = Serra de Magé #2, Jon3 = Jonzac #3, Bou2 = Bouvante #2, Sta5 = Stannern #5, N23-3 = NWA4523 #3, N70-3 = NWA1670 #3.

2006); NWA6291 is the most recently found angrite and is possibly paired with NWA2999 (Bouvier et al., 2011).

2.2. Analytical methods

Several hundred milligrams of meteorite was crushed in an agate mortar to ensure representative sampling. Be-

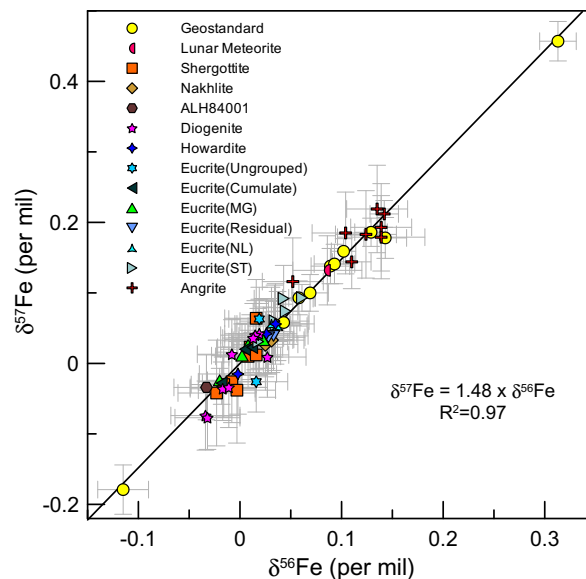


Fig. 2. Mass-dependent isotope fractionation of all samples in this study, shown in a $\delta^{56}\text{Fe}$ – $\delta^{57}\text{Fe}$ space. Eucrite (MG) = Eucrite Main Group, Eucrite (NL) = Eucrite Nuevo Laredo Trend, Eucrite (ST) = Eucrite Stannern Trend. Each point represents one meteorite/geostandard. For replicate measurements in Table 1, only the average is plotted.

tween 5 and 10 mg of crushed material was fully digested by sequential mixtures of HF–HNO₃ and HNO₃–HCl. Iron in the dissolved samples was then purified using anion-exchange chromatography (see Table 2 for the detailed procedure; Strelow, 1980; Dauphas et al., 2004, 2009b). The purification was repeated twice, thus ensuring the removal of all the matrix elements. Most separation chemistries were carried out in the clean laboratory at Washington University in St. Louis (WUSTL) and supplemented by chemistries for selected samples at the University of Chicago (UofC) to confirm inter-laboratory concordance and reproducibility. Most iron isotope analyses were performed with a Thermo Scientific Neptune MC-ICP-MS at the UofC and some were done with a Thermo Scientific Neptune Plus MC-ICP-MS at WUSTL, both following identical protocol described in Dauphas et al. (2009b).

In order to further confirm the validity of the iron separation technique used in this study (marked as “routine” chemistry), two other Fe separation techniques were applied to selected samples (marked as “long” and “UTEVA” chemistry in Table 2). The “long” column method uses long Teflon columns (10.5 cm) filled with 3 ml AG1-X8 anion-exchange resin (Dauphas et al., 2009b; Craddock and Dauphas, 2011). The “UTEVA” column method uses pre-packaged 1 mL Eichrom UTEVA resin cartridges (Horwitz et al., 1992; Tissot and Dauphas, 2011). All samples measured following purification by the three chemistry protocols yield identical iron isotope compositions within analytical uncertainty (Fig. 1 and Table 1).

All data are reported in $\delta^{56}\text{Fe}$ and $\delta^{57}\text{Fe}$ notations relative to the isotopic reference IRMM-014, defined as

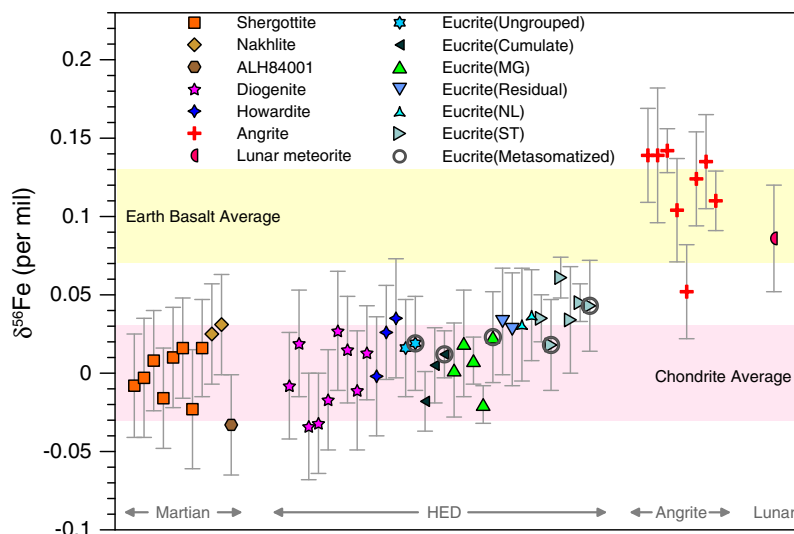


Fig. 3. Iron isotope compositions ($\delta^{56}\text{Fe}$) of martian meteorites, HEDs, and angrites. Colored shadow areas schematically show “Earth Basalt Average” and “Chondrite Average”, established by a compilation of literature data (Dauphas and Rouxel, 2006; Dauphas et al., 2009a). Eucrite (MG) = Eucrite Main Group, Eucrite (NL) = Eucrite Nuevo Laredo Trend, Eucrite (ST) = Eucrite Stannern Trend. Each point represents one meteorite. For replicate measurements in Table 1, only the average is plotted. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

$$\delta^x\text{Fe} = \left[\frac{(^x\text{Fe}/^{54}\text{Fe})_{\text{sample}}}{(^x\text{Fe}/^{54}\text{Fe})_{\text{IRMM-014}}} - 1 \right] \times 1000 \quad (1)$$

where $x = 56$ or 57 . In a three-isotope plot ($\delta^{56}\text{Fe}$ vs. $\delta^{57}\text{Fe}$), all samples define a straight line of slope 1.48 (Fig. 2), consistent with mass-dependent isotope fractionation. Note that during measurements, IRMM-524a was used as bracketing standard because it has identical isotope composition to IRMM-014 and it is more readily available. The analytical uncertainties reported here for each sample are 95% confidence intervals (see Dauphas et al. (2009b) for a detailed account on how error bars are calculated). Weighted averages and uncertainties are calculated for replicate analyses of meteorites using the following two equations:

$$\bar{x} = \frac{\sum_{i=1}^n (x_i/e_i^2)}{\sum_{i=1}^n (1/e_i^2)} \quad (2)$$

$$\sigma^2 = \frac{1}{\sum_{i=1}^n (1/e_i^2)} \quad (3)$$

where e_i is the standard deviation for each independent analysis. The uncertainties of previously published $\delta^{56}\text{Fe}$ data cited in this paper are typically 2 standard deviation (2SD) or 2 standard error (2SE). Caution should be taken when comparing these weighted average uncertainties to the 2SD or 2SE in previous studies because in most instances, accuracy of the measurements has not been tested below $\pm 0.03\text{‰}$.

3. RESULTS

Our iron isotope data are reported in Table 1, and illustrated in Figs. 2 and 3. The iron isotope compositions are compared to those of chondrites and terrestrial basalts,

for which 10 carbonaceous chondrites define an average $\delta^{56}\text{Fe}$ value of -0.010‰ and Earth basalts define an average value of $\sim +0.1\text{‰}$ relative to IRMM-014 (Craddock and Dauphas, 2011).

3.1. Earth and Moon

The terrestrial geostandards measured here (BIR-1, BCR-2, BHVO-2, AGV-1, AGV-2, GSP-1 and AC-E) give iron isotope compositions that are consistent with previous reports (see Craddock and Dauphas (2011) for a data compilation). The composition of BCR-2 measured here is slightly lighter than that reported by Craddock and Dauphas (2011). Three modern island arc basalts (IABs) treated with different techniques (routine, long and UTEVA chemistries) also agree with the data in a previous study (Dauphas et al., 2009a).

The iron isotope composition of the lunar highland meteorite MAC88105 is $\delta^{56}\text{Fe} = +0.09 \pm 0.03\text{‰}$, which is consistent with previously reported Apollo lunar highland rocks (Poitrasson et al., 2004; Weyer et al., 2005). Other lunar lithologies have variable iron isotope compositions that encompass this value, from $\delta^{56}\text{Fe} = +0.03\text{‰}$ for low-Ti mare basalts (Weyer et al., 2005) to $+0.27\text{‰}$ for high-Ti mare basalts (Moynier et al., 2006). The formation of these highly fractionated iron isotope compositions in lunar samples has been discussed in several recent papers (Poitrasson et al., 2004; Weyer et al., 2005, 2007; Poitrasson, 2007; Liu et al., 2010), and will not be further addressed here.

3.2. Martian meteorites

The eight shergottites studied have uniform $\delta^{56}\text{Fe}$ equal to $0.00 \pm 0.01\text{‰}$. These new and more precise data are in

agreement with previous reports (average $+0.01\%$; [Poitrasson et al., 2004](#); [Weyer et al., 2005](#); [Anand et al., 2006](#)), and confirm that shergottites have an iron isotope composition identical to chondrites. The $\delta^{56}\text{Fe}$ of two nakhlites ($+0.03 \pm 0.02\%$) and ALH84001 ($-0.03 \pm 0.03\%$) are also very similar to shergottites (within analytical precision). A close look at the data hints at the presence of possibly different iron isotope compositions among different groups of martian meteorites. However, these differences are at the limits of current analytical precision.

3.3. HED meteorites

Our new data confirm the findings of previous reports that most eucrites (cumulate eucrites, Main Group, Nuevo Laredo Trend and residual eucrites) have a chondritic iron isotope composition (average $\delta^{56}\text{Fe} = 0.00 \pm 0.01\%$). However, it is significant that Stannern Trend (ST) eucrites (average $\delta^{56}\text{Fe} = +0.05 \pm 0.01\%$) are isotopically heavier than other eucrites. Our results for six individual ST eucrites are more precise than the four previous measurements, which were limited to two different ST eucrites samples: Stannern ($\delta^{56}\text{Fe} = 0.00 \pm 0.05\%$, [Schoenberg and von Blanckenburg, 2006](#); $+0.01 \pm 0.02\%$, [Weyer et al., 2005](#); $-0.06 \pm 0.06\%$, [Zhu et al., 2001](#)) and Bouvante ($\delta^{56}\text{Fe} = +0.03 \pm 0.04\%$, [Weyer et al., 2005](#); $+0.03 \pm 0.05$, [Poitrasson et al., 2004](#)). To confirm the heavier iron isotope composition of ST eucrites measured, we independently processed and analyzed the same splits at UofC using three independent chemistry methods ([Table 2](#)). All these replicated measurements of ST eucrites show consistent results ([Fig. 1](#) and [Table 1](#)). ST eucrites are enriched in heavy iron isotopes compared to other classes of eucrites. Ungrouped eucrites, Pasamonte ($+0.02 \pm 0.03\%$) and NWA1240 ($+0.02 \pm 0.03\%$), have iron isotope compositions identical to other eucrites. Three howardites (average $\delta^{56}\text{Fe} = +0.02 \pm 0.02\%$) and nine diogenites ($0.00 \pm 0.01\%$) are isotopically identical to all classes of eucrites, excluding ST. Hence, iron isotopes, unlike oxygen isotopes, cannot distinguish ungrouped eucrites from other HEDs.

3.4. Angrites

The eight samples of angrites (average $\delta^{56}\text{Fe} = +0.12 \pm 0.01\%$) are isotopically heavier than chondrites, but similar to terrestrial basalts. To ensure that these data were correct, we again processed samples independently at WUSTL and UofC using three chemistry methods. All replicate measurements of the same samples processed by different methods gave the same iron isotope compositions within analytical uncertainty ([Table 1](#) and [Fig. 1](#)) and confirm that angrites have heavy fractionated iron isotope compositions.

4. DISCUSSION

In the following section, we discuss the implications of the enrichment in the heavy isotopes of Fe observed in angrites and Stannern Trend (ST) eucrites in terms of the redox state of the parent bodies and magmatic differentiation.

4.1. Redox-controlled iron isotope fractionation on the angrite parent-body?

Several processes can be considered in order to explain the fractionated iron isotope composition of basaltic angrites compared to other basaltic meteorites, including fractionation during low temperature terrestrial alteration, metal–silicate segregation, mineral fractionation, volatilization during impact, and partial silicate mantle melting. All angrites studied here are finds from the Sahara desert, Antarctica or Argentina. Leaching of isotopically light iron during terrestrial weathering can raise the $\delta^{56}\text{Fe}$ value of the residual rock up to $+0.1\%$ ([Saunier et al., 2010](#)). This process, however, is an unlikely candidate to explain the enrichment of heavy isotopes in angrites because all the angrites have a very narrow range of $\delta^{56}\text{Fe}$ ($+0.12 \pm 0.01\%$). These samples were found across a wide range of environments (from icy to hot desert) and have been subject to different exposure histories, which would not fractionate iron isotopes during alteration to an almost identical extent. In addition, for all other achondrite groups measured in this study, there is no difference in iron isotope composition between meteorite falls and finds indicating that terrestrial exposure has not affected isotope behavior for the samples in this study.

Equilibrium iron isotope fractionation between metal core and mantle is also not a viable explanation for the fractionated iron isotope compositions of angrites. High temperature ($2000\text{ }^\circ\text{C}$) and high pressure (7.7 GPa) equilibrium melting experiments of chondritic materials show no iron isotope fractionation between metal alloy and silicate melt ([Poitrasson et al., 2009](#)). Low-temperature, low-pressure metal silicate segregation would enriched the metal in heavy isotopes and the silicates in lighter isotopes (as seen in pallasites by [Poitrasson et al. \(2005\)](#)). [Campbell and Humayun \(2005\)](#) proposed that IVB iron meteorites, enriched in heavy iron isotopes ([Williams et al., 2006](#)), could be the core of the angrite parent-body. If this is the case, angrite is expected to be enriched in light iron isotopes, which is the inverse of what is observed in angrites. Only at extremely high pressures ($>100\text{ GPa}$) encountered at the core–mantle boundary of the Earth, metallic core could be enriched in light iron isotopes while silicate mantle could be enriched heavy iron isotopes according to theoretical calculations ([Polyakov, 2009](#)). Although the parent-body of angrites is still unidentified; there is little doubt that these meteorites come from a small-scale asteroid ([Burbine et al., 2006](#); [Rivkin et al., 2007](#); [Trilling et al., 2007](#)).

Preferential evaporation of lighter Fe during impact events (and enrichment in the heavier in the residue) is a mechanism that has been proposed to explain the difference between terrestrial and lunar basalts ([Poitrasson et al., 2004](#)). All angrites (except Angra dos Reis and NWA2999) are unshocked and do not show particular impact features ([Scott and Bottke, 2011](#)). In addition, angrites are depleted in moderately volatile elements, but they are not notably more depleted in moderately volatile elements (*e.g.*, Zn, Cd) than eucrites ([Weisberg et al., 2006](#)). If volatilization due to impacts was the origin of the fractionation observed in angrites, similar effects should be found in eucrites, which

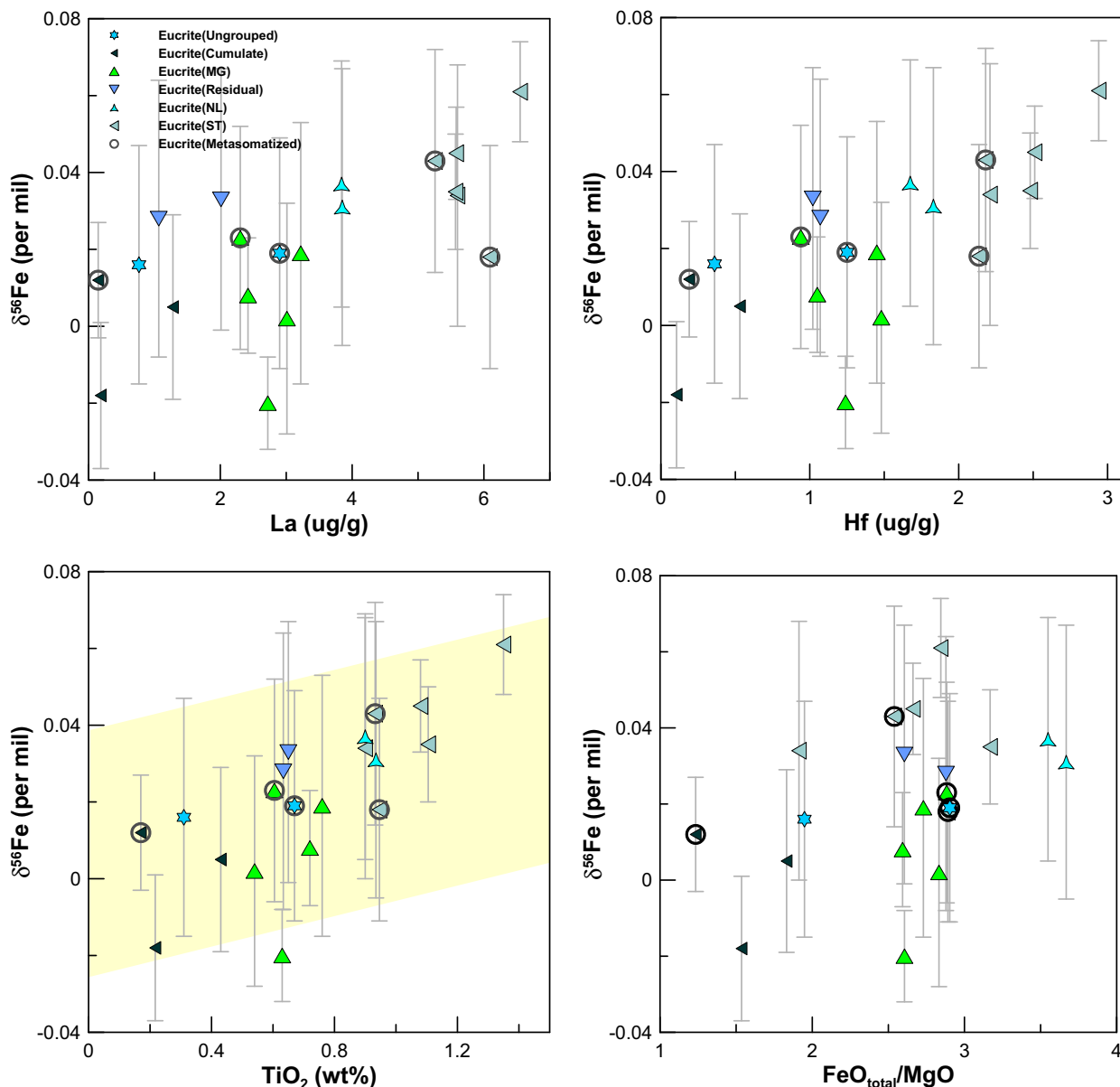


Fig. 4. Iron isotope compositions ($\delta^{56}\text{Fe}$) vs. La, Hf, TiO_2 and $\text{FeO}_{\text{total}}/\text{MgO}$. The yellow shaded area represents ilmenite-controlled iron isotope fractionation (see text for details). Eucrite (MG) = Eucrite Main Group, Eucrite (NL) = Eucrite Nuevo Laredo Trend, Eucrite (ST) = Eucrite Stannern Trend. Each point represents one meteorite. For replicate measurements in Table 1, only the average is plotted. Data sources: The concentration data of Pasamonte, Moore County, Cachari, Jonzac, Serra de Magé, and Camel Donga come from Barrat et al. (2000); Nuevo Laredo, Juvinas, Stannern, NWA4523, and Bouvante from Barrat et al. (2007b); NWA049 and NWA1240 from Barrat et al. (2003); Sahara 02501 from Barrat et al. (2007a); NWA2061 from Barrat et al. (2011); Yamato75011 from Fukuoka and Ikeda (1983); Agoult and Dag945 from Yamaguchi et al. (2009); EET87548 comes from Warren et al. (2009); Pomozdino from Kitts and Lodders (1998). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

is not the case. Therefore the enrichment in the heavier iron isotopes of angrites is unlikely caused by volatilization during impact processing, although this cannot be definitely ruled out.

Redox-controlled iron isotope fractionation through partial mantle melting provides one possible explanation for the fractionated iron isotope compositions of angrites. Redox proxies indicate that angrites probably formed in a

relatively oxidized environment (IW+1 to +2; Jurewicz et al., 1991, 1993; McKay et al., 1994), even though the exact redox conditions are difficult to quantify. Dauphas et al. (2009a) devised a quantitative model of iron isotope fractionation between source and melt as a function of $\text{Fe(III)}/\text{Fe(II)}$ ratio. According to this model, given the low $\text{Fe(III)}/\text{Fe(II)}$ ratio of angrites, little Fe isotope fractionation would be expected during partial melting. How-

ever, the exact redox condition of the mantle of angrite's parent body remains poorly known and it is also uncertain whether redox conditions can affect iron equilibrium isotope fractionation factors by modifying the structure of the melt even in systems with little Fe(III). Further work remains to be done to understand iron isotopic fractionation in angrites.

4.2. Iron isotope fractionation on the HED parent-body

Petrological arguments indicate that HED meteorites were formed in a low oxygen fugacity environment, close to the IW buffer, with Fe(III) absent (Stolper, 1977; Hewins and Ulmer, 1984). The chondritic $\delta^{56}\text{Fe}$ values of diogenites, howardites, cumulate eucrites, MG-NL eucrites, and residual eucrites are consistent with a chondritic iron isotope composition of the HED parent-body mantle that has not been modified by core segregation or partial melting, as is also seen on Mars (Poitrasson et al., 2004).

The $\delta^{56}\text{Fe}$ values of ST eucrites are higher than those of other eucrites. ST eucrites do not show any particular depletion in volatile elements or impact features when compared to other eucrites (Barrat et al., 2007b). Therefore, volatilization of light isotopes of Fe due to impact is a very unlikely origin for iron isotope difference between ST eucrites and other eucrites. Iron isotope fractionation is also observed in lunar basalts where redox state is extremely low (IW–2 to –1; Liu et al., 2010). It has been proposed that ilmenite fractionation controls the 0.1‰ difference in $\delta^{56}\text{Fe}$ between low-Ti and high-Ti lunar mare basalts (Craddock et al., 2010). Ilmenite is enriched in the heavy isotopes of iron between 0.16‰ and 0.42‰ in $\delta^{56}\text{Fe}$ (Craddock et al., 2010). ST eucrites are richer in Ti than the other eucrites and petrographic studies have also shown there is relatively more ilmenite in ST eucrites than in other eucrites (0.93–1.20 vol % vs. trace–1.00 vol %, respectively; Delaney et al., 1984; Warren et al., 1990). By mixing ilmenite with $\delta^{56}\text{Fe} = +0.42\text{‰}$ (the maximum fractionation) to main-group or cumulate eucrites ($\delta^{56}\text{Fe} \approx 0$), we obtained a $\delta^{56}\text{Fe} = 0.02\text{‰}$ (see Appendix for the mixing calculation; the calculation results are plotted as the shaded area in Fig. 4). This could be enough to explain the observed ST eucrites values within analytical errors ($\pm 0.03\text{‰}$). If the ilmenites have a lower $\delta^{56}\text{Fe}$, only a fraction of the isotopic effect observed in ST eucrites can be explained by ilmenites and other source of isotope fractionation cannot be excluded. It is possible that other Fe-bearing minerals also controlled the iron isotope composition of eucrites, and further work is needed to characterize Fe isotope fractionation between minerals in eucrite meteorites.

The origin of ST eucrites is debated (see review by McSween et al. (2010)). ST eucrites are petrographically and geochemically similar to MG eucrites, but are enriched in incompatible elements (Fig. 4). Their origin was first explained as smaller degrees of partial melting of the same source as other eucrites (Consolmagno and Drake, 1977; Stolper, 1977; Hsu and Crozaz, 1996). However, this model is not consistent with the growing database for trace lithophile and siderophile elements (Mittlefehldt and Lindstrom, 2003; Barrat et al., 2007b). It has been proposed recently

that ST eucrites are derived from the same parent magma to MG eucrites and have been modified by the assimilation of the asteroidal crust (Barrat et al., 2007b). This model is supported by our iron isotope data. It is observed in this study that $\delta^{56}\text{Fe}$ is correlated with Ti and other incompatible elements (Fig. 4). Non-modal re-melting of previously formed asteroidal crust would preferentially consume ilmenite (Barrat et al., 2007b). Partial melts formed from the eucritic crust will be rich not only in Ti, but also in the heavy isotopes of Fe because ilmenite in the case of the Moon has been demonstrated to be consistently heavy by up to 0.42‰ in $\delta^{56}\text{Fe}$ (Craddock et al., 2010). Contamination of ordinary basaltic eucrites by such components could explain the origin of ST eucrites with fractionated iron isotope compositions.

4.3. Iron isotopes and fluid–rock interactions on asteroidal bodies

A few eucrites display pre-terrestrial secondary minerals, which may have recorded interactions with fluids. Serra de Magé contains quartz veinlets, which have been ascribed by Treiman et al. (2004) to the circulation of water. A few other eucrites display Fe-enrichment along the cracks that crosscut the pyroxenes, and sometimes deposits of Fe-rich olivine and anorthitic plagioclase inside the fractures (Barrat et al., 2011). The origin of these phases is a matter of debate. Barrat et al. (2011) have proposed that this Fe-enrichment is the result of a metasomatic event. The nature of the metasomatic agent is not clear, but could have been an aqueous fluid. Subsequently, Roszjar et al. (2011) proposed that Fe-olivine and anorthite were formed by incongruent in situ melting of pyroxene at a temperature slightly above the temperature of formation of the primary pyroxenes, at about 1150 °C.

Fluid–rock interaction and exsolution of aqueous fluids could preferentially remove light iron isotopes from bulk rock and is an important mechanism for iron chemical and isotope fractionation in both low- and high-temperature processes on Earth (Rouxel et al., 2003; Poitrasson and Freyrier, 2005; Chapman et al., 2009). Thus, the fingerprint of aqueous fluid interaction in eucritic rocks could be detected using Fe isotopes. Among our samples, five eucrites have been suspected to interact with aqueous fluids: Serra de Magé, Pasamonte, NWA049, NWA2061 and Y-75011 (circled points in Figs. 3 and 4). None of these samples show iron isotope compositions fractionated relative to other similarly grouped eucrites. These results do not rule out definitively the involvement of fluids during eucrite evolution, but imply that fluid interactions on eucritic parent-bodies had negligible impact on iron isotope compositions. These results exclude the possibility that secondary processes on HED parent-body are the reason for the fractionated iron isotope compositions of ST eucrites.

5. CONCLUSION

We have found that Fe is enriched in the heavier isotopes in angrites (by $+0.12 \pm 0.01\text{‰}$; similar to terrestrial

Table A1

Parameters of the mass-balance used in Eq. (A1) to calculate the ilmenite-controlled iron isotope fractionation.

Parameter	Meaning	Value	Reference
$A_{\text{cumulate_MG}}$	Ilmenite abundance in eucrite (cumulate) and eucrite (MG) (vol %)	Trace-1.00	Delaney et al. (1984)
A_{ST}	Ilmenite abundance in eucrite (ST) (vol %)	0.93–1.20	Delaney et al. (1984) and Warren et al. (1990)
ρ_{ilmenite}	Ilmenite density (g/cm^3)	4.79	Lodders and Fegley (1998)
ρ_{eucrite}	Eucrite average density (g/cm^3)	3.20	Kitts and Lodders (1998)
C_{ilmenite}	Iron concentration in ilmenite (%)	36.81	Ilmenite (FeTiO_3) stoichiometric value
C_{ST}	Iron concentration in eucrite (ST) (%)	13.72–15.27	Kitts and Lodders (1998)
$\delta^{56}\text{Fe}_{\text{ilmenite}}$	Iron isotope composition of separated ilmenite	0.16–0.42	Craddock et al. (2010)
$\delta^{56}\text{Fe}_{\text{cumulate_MG}}$	Average iron isotope composition of eucrite (cumulate) and eucrite (MG)	~ 0.00	This study and previous references (Poitrasson et al., 2004; Weyer et al., 2005; Schoenberg and von Blanckenburg, 2006)

Notes: Eucrite (MG) = Eucrite Main Group, Eucrite (ST) = Eucrite Stannern Trend.

basalts) and in Stannern Trend eucrites (by $+0.05 \pm 0.01\%$). Martian and HEDs (except Stannern Trend) meteorites have iron isotope compositions identical to chondrites. Isotope fractionations during magmatic processes (partial melting, mineral fractionation and fluid exsolution) under different redox conditions are the best explanations to explain iron isotope variations in planetary crusts. However, further work remains to be done to document experimentally equilibrium iron isotope fractionation between melts and minerals to understand iron isotope variations in magmatic rocks. Our results show that not only terrestrial and lunar basalts are isotopically fractionated in iron but also crustal rocks from at least two asteroids with different sizes and volatilization histories.

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APPENDIX A

Ilmenite is enriched in the heavy isotopes of iron between 0.16‰ and 0.42‰ in $\delta^{56}\text{Fe}$ (Craddock et al., 2010). By using the $\delta^{56}\text{Fe}$ value of separated ilmenite and the modal abundance of ilmenite in ST eucrites, we can construct a simplified mixing model of ilmenite-controlled iron isotope fractionation (Eq. (A1)).

$$\delta^{56}\text{Fe}_{\text{ST}} = \delta^{56}\text{Fe}_{\text{cumulate_MG}} + \Delta_{\text{ilmenite}} \cdot \delta^{56}\text{Fe}_{\text{ilmenite}} \quad (\text{A1})$$

$\delta^{56}\text{Fe}_{\text{ST}}$ is the iron isotope composition of ST eucrites; $\delta^{56}\text{Fe}_{\text{cumulate_MG}}$ is the iron isotope composition of cumulate or MG eucrites; $\delta^{56}\text{Fe}_{\text{ilmenite}}$ is the iron isotope composition of mineral ilmenite. Δ_{ilmenite} is the enrichment of ilmenite in ST eucrites compared to cumulate or MG eucrites; Δ_{ilmenite} is a function of the abundance of ilmenite in ST and in cumulate or MG eucrites. Considering the conversion of volume percentage reported in references to weight percentage, and iron concentration in mineral ilmenite and in bulk eucrites, we could write:

$$\Delta_{\text{ilmenite}} = (A_{\text{ST}} - A_{\text{cumulate_MG}}) \cdot \frac{\rho_{\text{ilmenite}}}{\rho_{\text{eucrite}}} \cdot \frac{C_{\text{ilmenite}}}{C_{\text{ST}}} \quad (\text{A2})$$

The meanings and values of the parameters used in Eq. (A2) are listed in Table A1.

Hence, Eq. (A1) can be developed into Eq. (A3):

$$\delta^{56}\text{Fe}_{\text{ST}} = \delta^{56}\text{Fe}_{\text{cumulate_MG}} + (A_{\text{ST}} - A_{\text{cumulate_MG}}) \cdot \frac{\rho_{\text{ilmenite}}}{\rho_{\text{eucrite}}} \cdot \frac{C_{\text{ilmenite}}}{C_{\text{ST}}} \cdot \delta^{56}\text{Fe}_{\text{ilmenite}} \quad (\text{A3})$$

By using the maximum fractionation value ($\delta^{56}\text{Fe} = 0.42\%$) measured in ilmenite (Craddock et al., 2010), we obtained a $\delta^{56}\text{Fe} = 0.02$ for ST eucrites, which could largely explain the difference between ST eucrites and cumulate or MG eucrites within analytical errors ($\pm 0.03\%$). However, if we use the minimum fractionation value ($\delta^{56}\text{Fe} = 0.16\%$) measured in ilmenite (Craddock et al., 2010), we obtained a $\delta^{56}\text{Fe} = 0.01$ for ST eucrites. In this case, other minerals

might also contribute to the enrichment of iron isotope in ST eucrites. Due to the lack of iron isotope data of relevant minerals, there is no further constraint on this problem.

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