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# Tracing metal–silicate segregation and late veneer in the Earth and the ureilite parent body with palladium stable isotopes

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# Abstract

Stable isotope studies of highly siderophile elements (HSE) have the potential to yield valuable insights into a range of geological processes. In particular, the strong partitioning of these elements into metal over silicates may lead to stable isotope fractionation during metal–silicate segregation, making them sensitive tracers of planetary differentiation processes. We present the first techniques for the precise determination of palladium stable isotopes by MC-ICPMS using a  $106Pd-110Pd$  doublespike to correct for instrumental mass fractionation. Results are expressed as the per mil (‰) difference in the <sup>106</sup>Pd/<sup>105</sup>Pd ratio ( $\delta^{106}Pd$ ) relative to an in-house solution standard (Pd\_IPGP) in the absence of a certified Pd isotopic standard. Repeated analyses of the Pd isotopic composition of the chondrite Allende demonstrate the external reproducibility of the technique of  $\pm 0.032\%$  on  $\delta^{106}$ Pd. Using these techniques, we have analysed Pd stable isotopes from a range of terrestrial and extraterrestrial samples. We find that chondrites define a mean  $\delta^{106}Pd_{chondrite} = -0.19 \pm 0.05\%$ . Ureilites reveal a weak trend towards heavier  $\delta^{106}$ Pd with decreasing Pd content, similar to recent findings based on Pt stable isotopes (Creech et al., 2017), although fractionation of Pd isotopes is significantly less than for Pt, possibly related to its weaker metal–silicate partitioning behaviour and the limited field shift effect. Terrestrial mantle samples have a mean  $\delta^{106}Pd_{\text{mantle}} = -0.182 \pm 0.130\%$ , which is consistent with a late-veneer of chondritic material after core formation.

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Keywords: Palladium; Stable isotopes; Terrestrial planet accretion; Late-veneer; Meteorites

# 1. INTRODUCTION

The platinum group elements (PGE; Rh, Ru, Pd, Os, Ir and Pt) all exhibit highly siderophile behaviour, with the bulk of Earth's highly siderophile element (HSE; i.e.,  $PGE + Re$ , Au) budget concentrated into the core. As such, the HSE have played a key role in constraining models of Earth's accretion and differentiation. Combined with their extreme depletion in the bulk silicate Earth, changes in

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oxidation state and bonding environment between mantle silicates and the Fe–Ni metallic core may lead to significant fractionation in HSE stable isotopes in the silicate mantles of terrestrial planets. This has led to considerable interest in stable isotope measurements of HSE, and stable isotope data for various sample types have recently been documented for Pt [\(Creech et al., 2014, 2017](#page-12-0)), Ru ([Hopp](#page-12-0) [et al., 2016\)](#page-12-0) and Os [\(Nanne et al., 2017\)](#page-13-0). Palladium represents a natural addition to this suite of stable isotope tracers, with potential to complement and contrast with these other HSE stable isotope systems.

Palladium is a transition metal with six naturally occurring stable isotopes— $^{102}Pd$ ,  $^{104}Pd$ ,  $^{105}Pd$ ,  $^{106}Pd$ ,  $^{108}Pd$ ,

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<span id="page-1-0"></span> $110Pd$ —with relative abundances of 1.02%, 11.14%, 22.33%, 27.33%, 26.46% and 11.72%, respectively [\(Meija et al.,](#page-12-0) [2016](#page-12-0); Fig. 1). Palladium can occur in a number of oxidation states, although principally occurs in nature as Pd(0) and Pd(II) ([Dennen, 1954; Borisov et al., 1994\)](#page-12-0). Palladium shares the highly siderophile nature of the PGEs, as evidenced by its relative depletion in the mantle as compared with chondritic meteorites (chondrites:  $\sim$ 1 µg g<sup>-1</sup> Pd; bulk silicate Earth:  $4 \text{ ng g}^{-1}$  Pd; [McDonough, 2014\)](#page-12-0). However, reported metal–silicate partition coefficients for Pd at conditions relevant to core formation are around an order of magnitude lower than those for the other PGE, including Pt  $(D_{Pd}^{met/sil} \sim 10^{3} - 10^{4}; D_{Pt}^{met/sil} \sim 10^{4} - 10^{5};$  [Holzheid et al.,](#page-12-0) [2000; Mann et al., 2012\)](#page-12-0). The PGE also exhibit chalcophile behaviour and in the conditions found in the mantle and crust they are most commonly found in sulphide components of ultramafic complexes. In addition to its weaker metal–silicate partitioning, Pd has the lowest density  $(11,995 \text{ kg m}^{-3})$ ; [Kaye and Laby, 1995\)](#page-12-0), melting point (1555 °C; [Kaye and Laby, 1995\)](#page-12-0), and 50% condensation temperature (1324 K; [Lodders, 2003\)](#page-12-0) of the PGE. Previous isotopic studies of Pd have been limited to searches for cosmogenic and nucleosynthetic effects in iron meteorites [\(Mayer et al., 2015](#page-12-0)), and as part of an experimental and theoretical study of nuclear volume effects [Fujii et al.](#page-12-0) [\(2011\)](#page-12-0). However, stable isotope variations of Pd in nature remain unexplored. The properties of Pd, in particular the relatively low condensation temperature, combined with subtle differences in the geochemistry of Pd as compared with Pt, its closest associate amongst the PGE, could lead to differences in stable isotopic compositions in nature.

Stable isotopic fractionations at temperatures relevant to metal–silicate differentiation have now been reported in several stable isotope systems, (e.g., Si (e.g., [Young et al.,](#page-13-0)



from [Meija et al. \(2016\).](#page-12-0)

[2015](#page-13-0)), Mo ([Hin et al., 2013; Burkhardt et al., 2014](#page-12-0)), Zn [\(Mahan et al., 2017\)](#page-12-0) and Fe ([Elardo and Shahar, 2017](#page-12-0))). The metal–silicate partitioning of Pd is significantly greater than for these elements, and combined with the differences in oxidation state and bonding environment between mantle silicates and the Fe–Ni metallic core, as well as evidence for Pt stable isotope fractionation related to metal–silicate partitioning ([Creech et al., 2017](#page-12-0)), significant Pd stable isotopic fractionation could occur during planetary core formation. Stable isotopic studies of Pd, especially where contrasted with similar data for Pt, may provide useful insights into a variety of planetary and geological processes.

In this paper, we present the first high-precision method to analyse the stable isotopic composition of Pd. We apply this technique to estimate the Pd isotopic composition of a wide range of chondrites and some modern terrestrial samples in order to provide a reference for the Pd isotopic composition of Solar System materials. We compare this solar system Pd isotopic composition to terrestrial Archean samples from Isua and South Africa to investigate the potential preservation of Pd isotopic signatures from prior to the late arrival of chondritic material, similar to those recently observed for Pt ([Creech et al., 2017\)](#page-12-0). We further analysed the Pd isotopic composition of a series of ureilite meteorites. Ureilites have a complex petrogenesis, but are considered to be mantle restites formed on a parent body that never totally melted (e.g., [Scott et al., 1993; Barrat](#page-13-0) [et al., 2015; Greenwood et al., 2016\)](#page-13-0). The ureilites have HSE contents ranging from approximately chondritic to depletions of several orders of magnitude relative to chondrites ([Warren et al., 2006; Rankenburg et al., 2008\)](#page-13-0), which have been interpreted to reflect the early stages of core formation [\(Warren et al., 2006\)](#page-13-0). These depletions have also been recently shown to have correlated stable isotope fractionations for the element Pt ([Creech et al., 2017](#page-12-0)). These factors, combined with ureilites being among the most common achondrites in our meteorite collections  $(\sim 400$  ureilites are currently reported in the Meteoritical Bulletin Database) make these meteorites some of the best available samples to test the effects of metal–silicate or metal–sulphide segregation on the Pd isotopic composition.

# 2. DESIGN AND PREPARATION OF THE PALLADIUM DOUBLE-SPIKE

The double-spike method is a well established approach for precise determination of isotope ratios, and has been applied to a broad range of isotope systems using both TIMS and MC-ICPMS analysis (e.g., Pb, Cr, Zn, Fe, Ni, Ti, Mo, Ca, Pt, W; [Compston and Oversby, 1969;](#page-12-0) [Eugster et al., 1969; Hamelin et al., 1985; Beard et al.,](#page-12-0) [1999; Beard and Johnson, 1999; Anbar et al., 2001;](#page-12-0) [Siebert et al., 2001; Baker et al., 2004; Dideriksen et al.,](#page-12-0) [2006; Gopalan et al., 2006; Schoenberg et al., 2008;](#page-12-0) [Arnold et al., 2010; Gall et al., 2012; Creech et al., 2013;](#page-12-0) [Hin et al., 2013; Burkhardt et al., 2014; Abraham et al.,](#page-12-0) [2015; Bonnand et al., 2016; Millet et al., 2016; Bezard](#page-12-0) [et al., 2016; Krabbe et al., 2017](#page-12-0)). These include several recent applications to studying mass-dependent fractiona-Fig. 1. Relative abundances of isotopes in the mass range of Pd<br>tion of PGEs [\(Creech et al., 2014, 2017; Hopp et al.,](#page-12-0)

<span id="page-2-0"></span>[2016; Nanne et al., 2017](#page-12-0)). An advantage of the double-spike approach is that, where mass-dependent isotopic fractionation is being studied, the natural isotopic fractionation can be determined from a single isotope measurement, and the double-spike can correct for any mass-dependent isotope fractionation induced during chemical purification and mass spectrometry. Therefore, the double-spike method is less susceptible to potential analytical artefacts induced by trace amounts of sample matrix that may remain and incomplete yields of the analyte following chemical purification. In addition, the double-spike also permits for precise and accurate determination of Pd concentrations from the same analysis as the isotope data.

## 2.1. Palladium double-spike design

With six stable isotopes, there are many different possible combinations available for the Pd double-spike and the double-spike inversion. The choice of Pd double-spike was made on the basis of minimum predicted errors as calculated using the *Double Spike Toolbox* of [Rudge et al. \(2009\)](#page-13-0), with the additional constraint that the isotopes used in the double-spike and the double-spike inversion not contain  $102$ Pd or  $104$ Pd due to the potential for interference on those isotopes from small amounts of  $102$ Ru and  $104$ Ru [\(Fig. 1\)](#page-1-0). We therefore chose to use a  $^{106}Pd-^{110}Pd$  double-spike using a 105Pd–106Pd–108Pd–110Pd inversion, as this combination has a small predicted error (ca. 29 ppm/amu; [Rudge et al.,](#page-13-0) [2009\)](#page-13-0) and a broad plateau in predicted error across a range of sample–spike mixtures (Fig. 2A), indicating that this combination is not highly sensitive to the sample–spike ratio.

# 2.2. Preparation of Pd double-spike

Two Pd isotope spikes ( $^{106}Pd$ ,  $^{110}Pd$ ) were obtained from Isoflex [\(www.isoflex.com\)](http://www.isoflex.com) in the form of metal powders, with certified isotopic compositions as given in [Table 1.](#page-3-0) The spikes were carefully weighed before transferring to Savillex Teflon beakers for dissolution with aqua regia. Once the spike material was fully digested, the spike solutions were repeatedly evaporated to dryness in concentrated HCl, and finally redissolved in 6 M HCl.

The  $^{106}Pd-^{110}Pd$  double-spike was prepared by mixing the two spikes in the optimal proportions of (47.17% <sup>106</sup>Pd, 52.83% <sup>110</sup>Pd as calculated using the *Double Spike* Toolbox of [Rudge et al. \(2009\)](#page-13-0). The double-spike mixture was then evaporated to dryness, taken up in a small volume of concentrated HCl, evaporated to dryness again, and finally taken up in 6 M HCl. Precise isotopic determination by the double-spike technique requires accurate calibration of the double-spike composition. The method for calibration of the double-spike prepared here and the accuracy of the calibrations are described below.

#### 3. MATERIALS AND METHODS

# 3.1. Chemicals and standard solutions

All laboratory work was performed in a class-100 clean room environment using class-10 laminar flow hoods at the



Fig. 2. (A) for the  $^{106}Pd^{-110}Pd$  double-spike with  $105Pd-106Pd-108Pd-110Pd$  inversion, as a function of doublespike–sample proportions, calculated using the Double Spike Toolbox of [Rudge et al. \(2009\)](#page-13-0). Note that this combination results in a relatively broad region around the optimum where low errors are found. (B) Measured  $\delta^{106}$ Pd as a function of the proportion of double-spike in the double-spike–sample mixture from 0.2 to 0.7, equivalent to a standard–spike ratio from 4.0 to 0.4, or a factor of 0.38–1.3 times the theoretical optimal spike–sample proportion of 0.525. Double-spike corrected results yield delta values ( $\delta^{106}$ Pd) that are within the typical reproducibility of measurements of the standard over this range of sample-spike mixtures. Error bars represent the 2 s.e. on individual measurements.

Institut de Physique du Globe de Paris, France. Pre-cleaned Savillex Teflon PFA beakers were used for all samples and solutions processed in this study. BASF Selectipur<sup>®</sup> AR grade acids  $(69\% \text{ HNO}_3, 37\% \text{ HCl})$  were further purified by sub-boiling distillation using Savillex DST-1000. All dilutions were carried out using ultra-pure (18.2 M $\Omega$  cm) Milli-Q water.

Due to the unavailability of any certified isotope reference materials for Pd, we employed a Pd ICP solution standard for this purpose (Fischer Chemical 1000 mg  $L^{-1}$ Pd in 5% HCl, lot #C64130). This solution standard is hereafter referred to as Pd\_IPGP; aliquots of Pd\_IPGP standard can be made available to other laboratories on request.

<span id="page-3-0"></span>Table 1

Certified isotopic compositions of single isotope spikes provided by Isoflex, and calibrated composition of 106Pd–110Pd double-spike.

Spike	Form	Certificate No.	$102$ Pd (%)	104Pd(%)	105Pd(%)	106Pd(%)	$108$ Pd $(%)$	110 <sub>pd</sub>
Pd-106	metal powder	6106	$<$ 0.03 $\,$	0.06	0.68	98.4	0.80	0.6
$Pd-110$	metal powder	6107	$0.01$	0.05	0.09	0.13	0.33	99.4
$106Pd - 110Pd$ double-spike			0.002	0.052	0.396	48.110	0.557	50.882

## 3.2. Terrestrial and meteorite samples

Details of the samples and amounts used in this study are summarised in [Table 4.](#page-7-0)

Eight terrestrial samples were studied. OKUM is an ultramafic komatiite (previously distributed by Geosciences Laboratories, now distributed by IAG) sampled from Serpentine Mountain, Ontario, Canada. This area is located at the western end of the Abitibi Greenstone Belt, and was emplaced into the crust ca. 2.7 Ga (Houlé [et al.,](#page-12-0) [2009](#page-12-0)). PGE in these rocks are found as Ni–Cu–PGE minerals (Houlé [et al., 2009](#page-12-0)). B25 is a komatiite from the ca. 3.5 Ga Barberton greenstone belt, South Africa [\(Caro](#page-12-0) [et al., 2006](#page-12-0)). BC7 is a peridotite xenolith from Batchelor's Crater, a Plio–Pleistocene alkali basalt vent in the Chudleigh volcanic province, North Queensland, Australia. LK-NIP is a dolerite from the Nipigon Diabase, Ontario, Canada distributed by Geosciences Laboratories. The Nipigon diabase sills were intruded into the crust ca. 1100 Ma [\(Hart and MacDonald, 2007; Heaman et al., 2007](#page-12-0)). Four metabasalts from the Isua Supracrustal Belt were analysed from Pd cuts that were kept from previous Pt chemistry by [Creech et al. \(2017\)](#page-12-0). In the southwest part of the belt, pillow lava structures are recognised within low deformation zone windows within a sequence of amphibolite rocks. Compositionally, the pillows are high-MgO  $(12-15 \text{ wt.}\%)$  tholeiitic basalts and are associated with ultramafic schists of komatiitic composition. A felsic unit which cross-cuts this pillow lava-bearing sequence gives U-Pb SHRIMP ages clustering around  $\sim$ 3.85 Ga [\(Michard-Vitrac et al., 1977; Baadsgaard](#page-12-0) [et al., 1984; Nutman et al., 1997](#page-12-0)) suggesting that these amphibolites are amongst the oldest of the Isua supracrustals.

The Pd isotopic composition of 11 chondritic meteorites were analysed in this study. These comprise six carbonaceous chondrites, four ordinary chondrites and one enstatite chondrite. For one carbonaceous chondrite (Allende), a large specimen of ca. 15 g was crushed and homogenised with an agate mortar and pestle, and from this homogenised powder seven replicate digestions of  $\sim 0.7-1.0$  g each were prepared. Ten Antarctic ureilite meteorites were analysed from a suite provided by NASA-JSC. These specimens span a large compositional range, with Pd contents ranging from ca. 21–165 ng/g [\(Rankenburg et al., 2008;](#page-13-0) [Table 4](#page-7-0)), and have also been analysed for Pt stable isotopes by [Creech et al. \(2017\)](#page-12-0).

## 3.3. Sample preparation and dissolution

Digestions of meteorite and terrestrial samples were carried out using a NiS fire assay method as described by

[Creech et al. \(2014, 2017\).](#page-12-0) Terrestrial samples were coarsely crushed and then powdered using an agate ball mill. Powders were weighed directly into porcelain crucibles for NiS digestion. Meteorite fragments were taken from sliced specimens, weighed and then cleaned by rinsing in ultrapure water, methanol and again in ultra-pure water. The meteorite fragments, typically weighing 0.5–1.0 g, were then placed in a Savillex Teflon beaker in ca. 4 mL of 2 M HCl. The beakers were heated to 120  $\degree$ C for ca. 10 min and then ultrasonicated for ca. 3 min. After this gentle acid washing, the samples were rinsed twice with water and then dried in the beaker on a hotplate. The samples were then crushed to a fine powder in an agate mortar and weighed before being transferred to porcelain crucibles for NiS fire assay digestion. NiS fire assays were carried out as described by [Creech et al. \(2014\)](#page-12-0), with 10 g of sodium carbonate, 20 g of sodium tetraborate, 1 g of nickel powder and 0.75 g of sulphur being added per 15 g of sample. Note that where less than 15 g of sample was processed, the amount of sample was made up to 15 g with pure silica.

Samples were double-spiked prior to NiS digestion by adding the spike directly to the mixture in the crucible and allowing to dry before being fused in a furnace at ca. 1030 °C for 90 min. Samples were optimally spiked in approximate proportions of 52.5% Pd from the doublespike and 47.5% from the sample. Where published Pd concentrations were unavailable they were estimated based on published results for similar samples. NiS beads were extracted from fused charges and dissolved in concentrated HCl as described by [Creech et al. \(2014\).](#page-12-0) Once fully digested, the samples were brought into solution in 10 mL of 0.5 M HCl for anion-exchange chemistry.

#### 3.4. Chemical separation of Pd

A protocol for the chemical separation of Pt from geological materials was presented by [Creech et al. \(2014\),](#page-12-0) utilising AG1-X8 (100–200 mesh) anion exchange resin [\(Table 2\)](#page-4-0). In that method, the major cationic species in geological samples are readily removed from the column using dilute acids, and the PGE are eluted using sequentially stronger HCl and  $HNO<sub>3</sub>$ . In order that in the future Pt and Pd isotopic composition may be analysed from the same samples, we employed this same method for Pd.

In this method, 1 mL of pre-cleaned AG1-X8 (100–200 mesh, chloride form) was loaded as a slurry onto Eichrom polypropylene columns with a 2 mL resin capacity and a 20 mL reservoir, and allowed to gravity settle. The resin bed was cleaned with  $20 \text{ mL of } 0.8 \text{ M HNO}_3$ ,  $10 \text{ mL of } 10^{-1}$ 11 M HCl,  $25 \text{ mL of } 13.5 \text{ M HNO}_3$  and  $40 \text{ mL of } 6 \text{ M}$ HCl, and then conditioned with 40 mL of 0.5 M HCl prior

<span id="page-4-0"></span>Table 2 Elution scheme for Pd based on [Creech et al. \(2014\)](#page-12-0).

Step	Eluent	Volume (mL)	
Load 1 ml AG1-X8 100-200 $\#$			
Clean column	0.5 M HNO <sub>3</sub>	20	
Clean column	11 M HCl	10	
Clean column	13.5 M $HNO3$	25	
Clean column	6 M HCl	40	
Conditioning	$0.5 M$ HCl	40	
Sample load	$0.5 M$ HCl	10	
Rinse (elute matrix)	$0.5 M$ HCl	40	
Rinse	1 M HCl	20	
Discard transition metals	$0.8$ M HNO <sub>3</sub>	20	
Collect Pd $(+Re, Ru)$	11 M HCl	60	
Collect Pt	13.5 M HNO <sub>3</sub>	40	

to sample loading. Samples were loaded onto the columns in 10 mL of 0.5 M HCl, which was followed by a rinse with 40 mL of 0.5 M HCl, during which virtually all of the major cationic constituents of the sample and some of the weakly retained PGE (Rh, Ru and Ir) were eluted. This was followed by elution of 20 mL of 1 M HCl to elute any remaining weakly retained PGE, and  $20 \text{ mL of } 0.8 \text{ M HNO}_3$  to elute any remaining transition metals such as Zn. The remaining PGE are recovered from the column in 11 M HCl (Ru, Pd, Re) and  $13.5 M HNO<sub>3</sub>$  (Pt).

Palladium is eluted from the column in 11 M HCl with recovery of ca. 96%. However, Ru and Re are also eluted in the same cut, with yields of ca. 37% and 94%, respectively (Fig. 3). Given that (1) there are no isobaric or likely polyatomic interferences arising from the presence of small amounts of Re, (2) that while Ru has isotopes at masses 102 and 104, these isobars of  $^{102}$ Pd and  $^{104}$ Pd can be avoided in the double-spike inversion, and (3) the doublespike approach has been shown to be less sensitive to the presence of minor amounts of remaining matrix elements



Fig. 3. Elution profile for a synthetic matrix with a chondritic composition. Major cationic components are not retained on the column and are eluted with the load and 0.5 M HCl rinse. The PGE are eluted in a sequence of HCl and HNO<sub>3</sub> acids. Pd is eluted in 11 M HCl with recovery of ca. 96%, although the Pd cut also contains some Ru and Re.

than sample–standard bracketing methods (e.g., [Dideriksen et al., 2006; Creech et al., 2014](#page-12-0)), we trialled making measurements on Pd cuts without further purification. A range of doping tests were conducted to test the suitability of this approach for analysis of natural samples, and these are presented in Section [4.2](#page-6-0) below.

## 3.5. Mass spectrometry and data reduction

Palladium stable isotope measurements were carried out using a Thermo-Scientific Neptune Plus multi-collector inductively-coupled-plasma mass-spectrometer (MC-ICPMS) at the Institut de Physique du Globe de Paris, France. The operating and measurement conditions of this instrument are summarised in [Table 3](#page-5-0). For the isotope measurements, the samples were dissolved in 0.5 M HCl and introduced into the mass spectrometer using an ESI Apex-HF desolvator with a  $100 \mu L \text{ min}^{-1}$  PFA nebuliser. The Neptune was run in low resolution mode, with a mass resolution of  $\sim$ 1700 (M/ $\Delta$ M as defined by the peak edge width from 5% to 95% full peak height; [Table 3\)](#page-5-0). The Pd isotopic composition of the sample and standard solutions were typically analysed at concentrations of 100 ppb Pd (i.e., samples were prepared with nominal Pd concentrations of 100 ng mL $^{-1}$ , although in some cases samples were run at concentrations of ca. 50 ng mL<sup>-1</sup>, against intensity matched standards). During Pd isotope measurements, ion beams at <sup>102</sup>Pd, <sup>103</sup>Rh,<sup>104</sup>Pd, <sup>105</sup>Pd, <sup>106</sup>Pd, <sup>107</sup>Ag, <sup>108</sup>Pd, <sup>109</sup>Ag, and <sup>110</sup>Pd were simultaneously collected in a single cycle using Faraday cups connected to amplifiers with  $10^{11}\Omega$  resistors for all Pd channels. This configuration resulted in a combined ion beam intensity for all Pd isotopes of  $\sim$ 90 V at an uptake rate of  $\sim$ 100 µL min<sup>-1</sup>, corresponding to a sensitivity of  $\sim 900 \text{ V ppm}^{-1}$ . During some analytical tests, a second cycle was added in which  $^{102}Pd$  was placed in the centre cup, permitting  $^{98}Ru$ ,  $^{99}Ru$ ,  $100$ Ru and  $101$ Ru to be collected in the low-mass side. Faraday collector gains were calibrated prior to the beginning of each analytical session.

Isotope results are reported in  $\delta^{106}$ Pd notation, which reports the per mil (‰) difference in the  $106Pd/105Pd$  ratio relative to our in-house Pd isotope standard (Pd\_IPGP). All data reduction was conducted offline using the freely available double-spike data reduction tool IsoSpike[\(www.](http://www.isospike.org) [isospike.org;](http://www.isospike.org) [Creech and Paul, 2015\)](#page-12-0).

# 4. RESULTS

#### 4.1. Double-spike calibration

For precise determination of isotope ratios by doublespike, it is critical that the isotopic composition of the double-spike be accurately calibrated to the composition of the isotope standard. To correct for instrumental mass bias during the calibration we used external normalisation to a Ag ICP-standard solution, assuming natural abundances for the Ag isotopes (51.84% for  $107$ Ag and 48.16% for 109Ag) from [Meija et al. \(2016\)](#page-12-0). Measurements of both the reference standard and the double-spike solution had matched Ag:Pd concentrations (1:2) and instrumental mass

Instrumental operating conditions and measurement parameter for the Neptune Plus MC-ICPMS.

<b>Instrument operating conditions</b>				
RF power		1300 W		
Plasma cool gas flow rate		16 L min <sup>-1</sup>		
Interface cones	-sampler	Jet cone		
	-skimmer	Ni skimmer, H-type		
Source slit width		$0.25$ mm		
Acceleration voltage		10 kV		
Instrument resolution		$\sim$ 1700		
Mass analyser pressure		ca. $8 \times 10^{-9}$ mbar		
Detector		9 Faraday detectors; all		
		Pd channels using $10^{11} \Omega$		
		preamplifiers		
Sample introduction system		ESI Apex HF		
Sample uptake rate		50 or 100 $\mu$ L min <sup>-1</sup>		
<b>Measurement parameters</b>				
Solution concentration		$100 \text{ ng}$ ;mL <sup>-1</sup>		
Typical sensitivity		$\sim$ 900 V;ppm <sup>-1</sup> total Pd		
Sample measurement time		$100 \times 8.4$ s integrations		
Washout time		$\sim$ 20 min		
Background measurement time		$100 \times 8.4$ s integrations, on-peak		



Fig. 4. Results of doping the Pd\_IPGP standard at a range concentrations relative to Pd with various elements that are abundant in rocks (Na, Mg, Ca, Ni, Fe, Zn) or with potential for isobaric interferences in the mass range of Pd (Ru, Cd).

fractionation was corrected assuming an exponential fractionation law. In this approach with both the standard and double-spike measured in the same session, the calibration of the double-spike is relative to the calibration of the standard, and thus this method gives accurate results even if the absolute isotopic composition of Pd\_IPGP is not precisely known. Ten analyses of both solutions were measured in the same session and the averages of these analyses

assumed as the isotope ratios of each solution relative to each other.

The double-spike calibration was tested by analysing a series of double-spike–standard mixtures in different mixing proportions, with the proportion of double-spike varying from 0.2 to 0.7 (i.e., standard–spike ratio 4.0–0.4, or a factor of 0.38–1.3 times the theoretical optimal spike–sample proportion of 0.525). The results of these tests show that

<span id="page-5-0"></span>Table 3

<span id="page-6-0"></span>

Fig. 5. (A) Pooled measurements of the Pd\_IPGP standard over a period of ca. six months, with different coloured symbols corresponding to separate measurement sessions. Small offsets occur between measurement sessions, resulting in an external reproducibility of  $\delta^{106}Pd = 0.006 \pm 0.017\%$  (2sd). (B) The same measurements of Pd\_IPGP as in panel A, where data for each session has been renormalised such the session mean for Pd\_IPGP has  $\delta^{106}Pd = 0$ , giving an external reproducibility of  $\delta^{106}Pd = 0.000$  $\pm$  0.007‰ (2sd). (C) Results of analyses of Pd\_IPGP standard at lower solution concentrations (as indicated) and thus lower signal intensities. Dashed boxes indicate the 2 sd uncertainty of the analyses they enclose. In all panels, error bars reflect the 2 se error of each analysis; where no error bars are visible, internal errors are smaller than the size of the data point.



Fig. 6. Assessment of external reproducibility for natural samples based on seven replicate digestions of the carbonaceous chondrite Allende, which yield  $\delta^{106}Pd = -0.167 \pm 0.032\%$ .

the double-spike corrected values yield delta values ( $\delta^{106}Pd$ ) that are within the typical reproducibility of measurements of the standard over this range of sample-spike mixtures ([Fig. 2B](#page-2-0)). These results demonstrate that our double-spike approach should yield accurate results even when sample Pd concentrations are not well known at the time of double-spiking.

## 4.2. Isotopic fractionation associated with matrix effects

Non-spectral matrix effects may arise due to the presence of other elements besides the analyte in the sample solution, which may depress the ionisation and throughput of the analyte in the plasma. In order to test for such effects, solutions of the Pd\_IPGP standard were doped with different elements (Na, Mg, Ca, Ni, Fe, Zn) in varying concentrations relative to Pd  $([X]/[Pd] = 0.01-1.00;$  [Fig. 4](#page-5-0)). These tests show that, using the double-spike approach, the presence of common rock-forming elements at proportions up to  $[X]/[Pd] = 1$  do not impact our double-spike corrected results at levels comparable to our analytical reproducibility.

Due to the high relative abundances of 102Ru and 104Ru (31.6% and 18.7%, respectively), Pd isotope analyses at these masses may be strongly affected by the presence of Ru in samples. As described in Section [2.1](#page-2-0) above, the choice of the  $106Pd-110Pd$  was partly motivated by the avoidance of these interferences, and does not use  $102$ Pd or  $104$ Pd in the double-spike inversion. To confirm that the presence of Ru in our samples does not impact the double-spike corrected Pd isotope analyses, measurements of the Pd\_IPGP standard were made with varying amounts of Ru ([Fig. 4\)](#page-5-0). At [Ru]/[Pd] up to  $\sim$ 1, double-spike corrected data remain within error of zero. Given that the relative yields of Pd and Ru from the chemical separation procedure are 96% and 37%, respectively, and that Pd and Ru are usually present in similar abundance in natural rocks, this demonstrates that our choice of double-spike inversion renders this approach insensitive to the presence of Ru at the levels that it is expected to be present in our samples.

<span id="page-7-0"></span>



 $35\,$ 

(continued on next page)



certificate value for that reference material. Pt data are from Creech et al.  $(2017)$ .

certificate value for that reference material. Pt data are from Creech et al. (2017).

Due to the possibility of isobaric interferences from Cd at masses 106, 108 and 110, we also made measurements of the Pd\_IPGP standard doped with varying amounts of Cd [\(Fig. 4](#page-5-0)). Unsurprisingly, these tests showed that with our choice of double-spike inversion isotopes, double-spike corrected isotope ratios are significantly affected by the presence of small amounts of Cd. In our tests, even at the lowest [Cd]/[Pd] ratio (0.01), the Pd standard falls well outside the reproducibility of the method, and at higher amounts of Cd, isotope ratios tend towards extremely inaccurate values. These results indicate that double-spike corrected ratios will be affected at levels greater than our analytical reproducibility above  $\text{[Cd]}/\text{[Pd]} = 0.003$ . However, Cd is throughly separated from Pd during our sample digestion and chemistry, and in our natural samples, [Cd]/[Pd] ratios in Pd cuts were always 1–2 orders of magnitude lower than this threshold. Thus, Cd in natural samples is not considered to have had any effect on the accuracy of our doublespike corrected data.

# 4.3. Accuracy and reproducibility of double spike MC-ICP-MS measurements

# 4.3.1. Solution standards

Double-spike corrected analyses of the Pd\_IPGP standard, under the analytical conditions given in [Table 3,](#page-5-0) yield an internal precision on  $\delta^{106}Pd$  of ca. 0.003‰ (2 se,  $n = 100$ ). Within an analytical session, the 2 sd uncertainty of repeat measurements was typically on the order of  $\sim 0.007\%$ . However, offsets are observed between the Pd IPGP averages from analytical sessions, which has a negative impact on the long-term apparent reproducibility, giving a mean of  $\delta^{106}Pd = 0.006 \pm 0.017\%$  (2 sd,  $n = 84$ ; [Fig. 5A](#page-6-0)) over a period of ca. six months. Such offsets are often noted in double-spike studies (e.g., [Schoenberg et al., 2008; Moeller et al., 2012; Creech](#page-13-0) [et al., 2014](#page-13-0)), and may be related to small changes in the abundance sensitivity of the MC-ICPMS, drifts in the relative gains of the Faraday collectors, or a variable component of non-exponential mass bias that is not readily corrected for by the DS method. Therefore, the Pd compositions of unknowns are corrected for this offset by normalising to the session mean of Pd\_IPGP measurements, which gives the long-term reproducibility on Pd\_IPGP measurements of  $\delta^{106}Pd = 0.000 \pm 0.007\%$  (2 sd,  $n = 84$ ; [Fig. 5](#page-6-0)B).

Analyses of Pd\_IPGP were also made at lower signal intensities, with concentrations of 50 ng mL<sup>-1</sup>,  $10 \text{ ng } \text{mL}^{-1}$  and  $1 \text{ ng } \text{mL}^{-1}$ , corresponding to ca. 40 ng, 8 ng and 0.8 ng of Pd consumed per analysis, respectively, as compared with ca. 80 ng for typical 100 ng/mL run solution. The results of this test are given in [Fig. 5](#page-6-0)(C). Solutions with 50 ng  $mL^{-1}$  of Pd yield an average  $\delta^{106}Pd = 0.00 \pm 0.01\%$ , which is only marginally worse than those at 100 ng mL<sup>-1</sup>. At 10 ng mL<sup>-1</sup>, the average  $\delta^{106}Pd = 0.00 \pm 0.02\%$ , and at 1 ng mL<sup>-1</sup> the data become inaccurate, giving  $\delta^{106}Pd = -0.07 \pm 0.03\%$ . Note that samples were typically run at a concentration of  $100 \text{ ng } \text{mL}^{-1}$ .

#### <span id="page-9-0"></span>4.3.2. Natural samples

The reproducibility of our technique for samples with a geological matrix was tested by analysis of replicate digestions,  $\sim 0.7$ –1.0 g each, of the carbonaceous chondrite Allende taken from a homogenised powder prepared from a large (15 g) fragment of that meteorite [\(Fig. 6](#page-6-0); [Table 4](#page-7-0)). Of the seven replicate Allende samples, three were processed without double-spiking prior to digestion and four were double-spiked prior to digestion. While there is a small offset (ca. 0.020‰) between the means of samples spiked before and after chemistry, almost all analyses are within the 95% confidence interval of the mean of both groups, suggesting that there is no significant difference in samples spiked before or after digestion. However, a difference is apparent in the Pd concentration derived from these groups, indicating that the Pd recovery from the NiS fire assay method is variable and potentially as low as ca. 60%. The Pd concentrations derived from replicates that were spiked prior to NiS fire assay are consistent with literature data for Allende (average Pd  $\sim$ 705 ng/g; [Tagle and](#page-13-0) [Berlin, 2008](#page-13-0)). These data suggest that there is no significant isotopic fractionation arising from incomplete Pd yields from the NiS fire assay. Together, these replicates yield a mean Allende composition  $\delta^{106}Pd = -0.167 \pm 0.032\%$  (2) sd). This is taken to represent the external reproducibility of the technique for natural samples, and error bars on subsequent plots of  $\delta^{106}Pd$  are either the 2 sd uncertainty for that sample or the reproducibility based on Allende replicates, whichever is larger.

#### 4.4. Natural Pd isotope variations

All data for natural samples are summarised in [Table 4](#page-7-0) and Fig. 7.

Data for 11 chondrites (six carbonaceous chondrites: 2 CM, 1 CV, 1 CO, 1 CR, 1 CK; four ordinary chondrites: 1 L, 1 LL, 2 H; one enstatite: 1 EH3), define a mean  $\delta^{106}Pd = -0.19 \pm 0.05\%$ , which is indistinguishable from the composition defined by replicates of Allende ([Table 4](#page-7-0);

Fig. 7). The terrestrial samples analysed here define a terrestrial composition of  $\delta^{106}Pd = -0.18 \pm 0.13\%$  [\(Table 4](#page-7-0)), although this terrestrial composition is discussed further in Section [5.2](#page-10-0), below. On average, the ureilites have a Pd isotopic composition similar to that of chondrites, with a mean  $\delta^{106}Pd = -0.197 \pm 0.065\%$  (2 sd). One ureilite sample, EET87157, appears to be an outlier, falling more than 2 sd from the mean. Excluding that sample from the mean gives  $\delta^{106}Pd = -0.205 \pm 0.044\%$  (2 sd).

## 5. DISCUSSION

#### 5.1. Palladium isotopic compositions of meteorites

The mean of 11 chondrites from all chondrite groups is indistinguishable from the composition defined by replicates of Allende [\(Table 4;](#page-7-0) Fig. 7), thus defining our chondritic Pd isotope composition,  $\delta^{106}Pd_{\text{chondrite}} = -0.19$  $\pm 0.05%$ .

There is some indication of potential systematic variations between the different chondrite groups. With the exception of Allende and EET 92002 (CK5), all of the carbonaceous chondrites are isotopically lighter than the mean chondrite value. The single enstatite chondrite sample is also slightly isotopically lighter than the mean, similar to the carbonaceous chondrites. Amongst the ordinary chondrites, the H chondrites are very close to the mean value, whereas the L chondrites are the isotopically heaviest chondrite samples. However, these variations are not significant relative to our analytical uncertainties, and further analyses are required to confirm this speculation.

The largest compositional range is observed amongst the carbonaceous chondrites, which could potentially indicate the presence of uncorrected mass-independent effects in those samples arising from the greater proportions of matrix and refractory inclusions (amoeboid olivine aggregates and Ca–Al-rich inclusions) found in that group, similar to observations made for Pt stable isotopes ([Creech](#page-12-0) [et al., 2017](#page-12-0)). Given that  $\delta^{106}$ Pd represents a mass difference



Fig. 7.  $\delta^{106}$ Pd data for chondrite and ureilite meteorites and terrestrial samples. Error bars represent the 2 sd uncertainty for each sample or the reproducibility of the technique based on replicate measurements of Allende (i.e.,  $\pm 0.032\%$ ), whichever is larger. Shaded boxes and horizontal lines represent the mean and 2 sd of the samples they enclose. The terrestrial samples are separated as described in the text, with the present-day mantle composition being defined by the three youngest samples. The terrestrial composition is extended over the Archean samples, which are all within uncertainty of  $\delta^{106}Pd_{\text{mantle}}$ .

<span id="page-10-0"></span>

Fig. 8. (A) Ureilite  $\delta^{106}Pd$  data vs. 1/[Pd], where [Pd] is the Pd concentration in ng g<sup>-1</sup>. Note that the Pd concentrations from [Rankenburg](#page-13-0) [et al. \(2008\)](#page-13-0) were used in lieu of our concentrations as double-spiking was not carried out before digestion for these samples. (B) Ureilite  $\delta^{106}$ Pd data vs.  $\delta^{198}$ Pt. Note that a 1:1 correlation with Pt isotope data would yield a slope of  $\sim$ 0.25, as  $\delta^{198}$ Pt represents a mass difference of 4 amu, compared with 1 amu for  $\delta^{106}Pd$ . Note that in both panels, the regression through the data excludes the sample EET87157, indicated by the half filled symbol, as an outlier, as it falls outside of a 2 sd error envelope.

of only 1 amu, the compositional range observed in chondrites is significant, and further investigations of these components are warranted. However, the mean of seven replicate digestions from a large homogenised sample of Allende is indistinguishable to that of all ordinary and enstatite chondrites, indicating that the bulk value of all chondrite groups is identical within our uncertainties.

The suite of ureilites presented here was also analysed for Pt stable isotopes by [Creech et al. \(2017\)](#page-12-0) and showed range of Pt isotope compositions with a correlation between  $\delta^{198}$ Pt and Pt concentration, which was interpreted to reflect Pt stable isotopic fractionation during metal–silicate partitioning. The range observed in Pd stable isotope compositions of ureilites is significantly smaller than that for Pt ([Table 4;](#page-7-0) [Fig. 7\)](#page-9-0), indicating that Pd isotopes are less strongly fractionated during metal–silicate differentiation. A subtle correlation is observed between Pd stable isotope compositions and Pd concentrations in ureilites, and a regression through  $\delta^{106}$ Pd vs 1/[Pd] gives a slope of 1.2 with an  $R^2 = 0.64$  (Fig. 8A). This slope suggests a Pd isotopic of fractionation 0.3 ppm per ng  $g^{-1}$  of Pd depletion. However, given the small range in Pd isotope compositions relative to our uncertainties, this correlation does not give a strong indication that a measurable Pd isotopic fractionation would be expected to accompany metal–silicate planetary differentiation. The Pd stable isotope data show a moderate correlation with Pt stable isotopes, where a linear regression through  $\delta^{106}$ Pd vs.  $\delta^{198}$ Pt gives a slope of 0.087 (Fig. 8); note that a 1:1 correlation would have a slope of 0.25, as  $\delta^{198}$ Pt represents a mass difference of four amu, compared with one amu for  $\delta^{106}$ Pd. The smaller range in Pd stable isotopes and weaker correlation with Pd concentration as compared with Pt may relate to the weaker partitioning of Pd between metal and silicate  $(D_{\text{Pd}}^{\text{met/sil}} \sim 10^3 - 10^4 \text{ vs.}$  $D_{Pt}^{met/sil} \sim 10^{4}$ –10<sup>5</sup>), or may support the idea that nuclear volume effects play a more significant role in stable isotope fractionation for the heavier element Pt as compared with

Pd, as has been observed for other elements [\(Schauble,](#page-13-0) [2007; Fujii et al., 2009; Moynier et al., 2013](#page-13-0)).

## 5.2. Palladium isotopic composition of Earth's mantle

The terrestrial samples we analysed represent different compositions, geological settings and ages. These samples also span a range of Pd stable isotopic compositions that exceeds that observed in the meteorite samples analysed so far.

In Pt stable isotope data ([Creech et al., 2017\)](#page-12-0), Archean samples from Isua have been recently shown to have heavy, non-chondritic Pt stable isotope compositions, potentially indicating preservation of some fraction of a pre-lateveneer signature of core-formation [\(Creech et al., 2017\)](#page-12-0). As such, we draw a distinction between some Archean samples in this study, as these may pre-date the complete equilibration of late-veneer with the upper mantle. [Maier et al.](#page-12-0) [\(2009\)](#page-12-0) showed, based on komatiite data, that PGE contents of the mantle increased through the Archean (4.5–2.9 Ga), which has been interpreted to reflect the progressive equilibration of late-veneer material with the mantle. Thus, the  $\sim$ 3.8 Ga Isua suite and the  $\sim$ 3.5 Ga komatiite B25 are treated as a separate group. The komatiite sample OKUM, while Archean, has an age of  $\sim$ 2.7 Ga, and it thus considered to represent mantle that had fully equilibrated with the late-veneer. Furthermore, OKUM has been shown to have a Pt stable isotope composition that is identical to the modern, chondritic mantle ([Creech et al., 2014, 2017\)](#page-12-0). The mantle-derived samples BC7 and OKUM may best characterise the upper mantle, suggesting a  $\delta^{106}Pd_{\text{mantle}} =$  $-0.22 \pm 0.03\%$ . However, there is no evidence yet that igneous processes fractionate Pd stable isotopes, and given that the dolerite sample LK-NIP also has a mantle-like Pt stable isotope composition [\(Creech et al., 2014](#page-12-0)), it cannot be readily excluded. Thus, we derive a terrestrial mantle value  $\delta^{106}Pd_{\text{mantle}} = -0.18 \pm 0.13\%$ . Further analyses of

terrestrial samples may refine this value. Based on our data so far, the mantle Pd isotope composition is indistinguishable from that of chondrites. At present is not clear that core formation would have imparted a Pd isotopic signature on Earth's mantle, and thus we cannot say conclusively that a chondritic Pd isotopic composition in the mantle requires a chondritic late-veneer. However, a chondritic isotopic signature remains fully consistent with that hypothesis.

Archean samples from Isua have been recently shown to have heavy, non-chondritic Pt stable isotope compositions, potentially indicating preservation of some fraction of a pre-late-veneer signature of core-formation ([Creech](#page-12-0) [et al., 2017](#page-12-0)). Conversely, Pd stable isotopes in the same samples are within error of our current estimate of the modern mantle composition [\(Fig. 7\)](#page-9-0). The lack of a heavy isotopic signature could indicate either that Pd isotopes were not fractionated to the same extent as Pt during core formation, or that the Pd has been impacted by secondary processes occurring in the crust. The Isua supracrustals have been variably affected by metamorphism, as indicated by disturbances in their Sm-Nd and Lu-Hf systematics (e.g., [Rizo et al., 2011, 2013\)](#page-13-0). In detail, three of the four Isua samples analysed in this study show uncorrelated  $\epsilon^{143}$ Nd(T) (calculated at 3.8 Gyr) and  $\mu^{142}$ Nd variations indicative of disturbance of the Sm-Nd systematics [\(Rizo et al., 2013\)](#page-13-0). Despite this, Pt stable isotope data show virtually no variability beyond the analytical uncertainties for the same samples  $(\delta^{198}Pt = +0.09 \pm 0.07\%)$ ; 2sd; [Creech et al., 2017](#page-12-0)), while the Pd isotopes show significant variability ([Fig. 7\)](#page-9-0). The variability in the Pd stable isotope data could indicate that Pd is less robust than Pt during secondary processes.

# 6. CONCLUSIONS

The new method presented in this paper allows for near quantitative extraction of Pd from geological samples suitable for high-precision isotope ratios measurements by double-spike MC-ICPMS. Using this new technique the first important results are as follows.

- Replicate digestions of the carbonaceous chondrite Allende demonstrate the external reproducibility of the technique of  $\pm 0.032\%$  on  $\delta^{106}$ Pd.
- Chondrites define a mean  $\delta^{106}Pd_{\text{chondrite}} = -0.19$  $± 0.05%$
- Ureilites data suggest that Pd isotopes are significantly less fractionated during metal–silicate segregation than for Pt, likely due to its weaker metal–silicate partitioning behaviour or the lesser impact of nuclear volume effects at the mass of Pd.
- Terrestrial mantle samples analysed so far give a  $\delta^{106}Pd_{\text{mantle}} = -0.18 \pm 0.13\%$ , which is consistent with a late-veneer of chondritic material after core formation.

Our data show that the Pd stable isotope system holds considerable potential for studying a range of geological and planetary processes.

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