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Highly siderophile and chalcophile element behaviour in abyssal-type and supra-subduction zone mantle: new insights from the New Caledonia ophiolite

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Abstract: The New Caledonia Ophiolite hosts one of the largest obducted mantle sections worldwide, offering a unique opportunity to investigate key mantle processes. The ophiolite comprises refractory harzburgites, locally overlain by mafic-ultramafic cumulates, and minor lherzolites. Previous geochemical studies indicated that the lherzolites are akin to abyssal-type peridotites, while the harzburgites underwent multiple melting episodes in MOR and supra-subduction zone environments, followed by late stage metasomatism.

In this work, Os isotopes, highly siderophile (HSE) and chalcophile element data are reported for the New Caledonia peridotites, in order to constrain the behaviour of these elements in abyssal-type and fore-arc mantle.

The variably serpentinised lherzolites (LOI = 6.4 - 10.7 %) yield slightly subchondritic to suprachondritic initial Os isotopic compositions (1870s/1880si = 0.1273-0.1329) and subchondritic to chondritic Re/Os ratios (0.04-0.11). The gently sloping HSE patterns with increasing depletion towards Au show concentrations in the range of other lherzolites from MOR or continental setting. Sulphur contents are high and variable (202-1268 ppm), and were likely increased during serpentinisation. By contrast, Se/Te ratios and concentrations are within the range of primitive mantle (PM) values, meaning that these elements were not significantly mobilised during serpentinisation. Despite displaying homogenous petrographic and geochemical features, the harzburgites are characterised by extremely heterogeneous Re-Os and HSE compositions.

Type-A harzburgites exhibit subchondritic 1870s/1880si (0.1203-0.1266) and low Re/Os ratios (0.01-0.04). The strong IPGE-PPGE fractionations (PdN/IrN = 0.21-0.56) coupled with positive Pt anomalies and S-Se-Te abundances often below the detection limit suggest high melt extraction

rates, resulting in sulphide consumption and Os-Ru metal alloy stabilisation.

Type-B harzburgites possess strongly fractionated, Os-Ir-Pt poor (Os = 0.003-0.072 ng/g, Ir = 0.0015-0.079 ng/g) and Pd-Re enriched patterns, associated with chondritic to suprachondritic measured 1870s/1880s (0.127-0.153). These characters are uncommon for highly depleted mantle residues. Interaction with an oxidised component does not appear as a viable mechanism to account for the IPGE-depleted patterns of type-B harzburgites, as calculated oxygen fugacities are close to the FMQ buffer (Log Δ FMQ= 0.20 to 0.48). The lower fO2 conditions register by type-B harzburgites compared to type-A likely records fO2 lowering due to continued melt extraction at un-buffered fO2 conditions. Type-B patterns may thus derive from slightly higher melting degrees, finally leading to Os-Ir-Pt release into the silicate melt. We propose that HSE geochemistry of the New Caledonia peridotites reveals superimposition of geochemical characters related to the recent Eocene evolution on a mantle source bearing a long term (> 1 Ga) evolution.

Abstract

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The variably serpentinised lherzolites (LOI = 6.4 - 10.7 %) yield slightly subchondritic to suprachondritic initial Os isotopic compositions ($^{187}Os/^{188}Os_i = 0.1273-0.1329$) and subchondritic to chondritic Re/Os ratios (0.04-0.11). The gently sloping HSE patterns with increasing depletion towards Au show concentrations in the range of other lherzolites from MOR or continental setting. Sulphur contents are high and variable (202-1268 ppm), and were likely increased during serpentinisation. By contrast, Se/Te ratios and concentrations are within the range of primitive mantle (PM) values, meaning that these elements were not significantly mobilised during serpentinisation. Despite displaying homogenous petrographic and geochemical features, the

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*Highlights (for review)

A comprehensive Re-Os, highly siderophile and chalcophile element investigation of the New Caledonia peridotites is presented.

HSE patterns of the lherzolites reflect low partial melting degrees of a mantle source that previously experienced melt percolation and radiogenic Os ingrowth.

Based on HSE and Os isotopic signature, two big groups of harzburgites (type-A and B) can be identified.

HSE and Re-Os systematics of type-A harzburgites are consistent with high melt extraction degrees, resulting in sulphide exhaustion and Os-Ru metal alloys stabilisation.

The HSE and Re-Os features of type-B harzburgites may record higher partial melting degrees and Os-Ir-Pt release into silicate melt after continued melt extraction.

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1. Introduction

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Highly siderophile elements (HSE: PGE+ Au-Re) are powerful geochemical tracers that 4 can provide useful information for a variety of mantle processes, such as mantle 5 melting, metasomatism and melt-fluid/mantle interaction (e.g. Luguet et al., 2001, 6 2003, 2007; Lorand et al., 2008; Ackerman et al., 2009). However, our knowledge 7 concerning the behaviour of HSE in mantle source rocks of primitive arc magmas and 8 9 the role of the subduction zone environment on HSE partitioning (i.e. hydrous melting, melt/fluid-mantle interaction) still remains quite fragmentary. Furthermore, although 10 abundant HSE data are now available for different types of mantle peridotites, HSE 11 data on fore-arc peridotites are remarkably scarce (e.g. Becker and Dale, 2016). 12 The New Caledonia ophiolite (Peridotite Nappe) hosts one of the largest and best 13 preserved mantle sections worldwide, providing an excellent opportunity to 14 investigate upper mantle processes. The rock exposures are dominated by harzburgite 15 tectonites bearing a supra-subduction zone affinity (Marchesi et al., 2009; Ulrich et 16 al., 2010; Pirard et al., 2013; Secchiari et al., 2019a). The main geochemical and 17 isotopic features of these rock-types reflect a complex polyphase evolution, including 18 several melting episodes in different geodynamic settings and subduction zone 19 metasomatism (Marchesi et al., 2009; Ulrich et al., 2010; Secchiari et al., 2019a). 20 Minor abyssal-type spinel and plagioclase lherzolites, with compositions similar to 21 abyssal peridotites, occur as discrete bodies in the north-western part of the island. 22 23 The Iherzolites record a different history compared to the extremely refractory harzburgites, as highlighted by their different geochemical signature (Ulrich et al., 24 2010; Secchiari et al., 2016). 25

In this work, a set of fully characterised peridotites (i.e. whole rock and in situ major and trace element contents, Sr-Nd-Pb isotopes) from New Caledonia (Secchiari et al., 2016, 2019a) has been used to investigate Re-Os, HSE and chalcophile element (S-Se-Te) systematics. The main aims of this work are: 1) to examine the behaviour of these elements in the Iherzolites (i.e. presumed abyssal peridotites) and in the ultradepleted harzburgites, which may represent rocks from a former supra-subduction zone mantle wedge; 2) to constrain the behaviour of HSE and chalcophile elements during subduction zone processes.

2. Geological setting and petrological background

New Caledonia is a NW–SE elongated island located in the SW Pacific region, between the eastern margin of Australia and the Vanuatu archipelago (Fig. 1a). The island represents the emerged portion of the submarine Norfolk Ridge and it is composed by a mosaic of volcanic, sedimentary and metamorphic terranes, ranging in age from Permian to Miocene (Aitchison et al., 1995; Cluzel et al., 2001, 2012; Lagabrielle et al., 2013). These terranes were amalgamated during two major tectonic events: 1) an Early Cretaceous tectonic convergence phase (Paris, 1981) and 2) a Paleocene to Late Eocene subduction culminated in the obduction of the ophiolite. Both events were characterized by high-pressure low-temperature (HP-LT) metamorphism in connection with plate convergence. New Caledonia can be sub-divided into four main geological domains (Cluzel et al., 2001; see Fig. 1): (i) the Basement units (pre-Late Cretaceous basement and Late Coniacian-to-Late Eocene sedimentary cover), (ii) the Cenozoic HP-LT metamorphic belt, (iii) the basaltic Poya Terrane and (iv) a large slab of peridotites, i.e. the Peridotite Nappe.

The Peridotite Nappe represents an allochtonous sheet of oceanic lithosphere 51 belonging to the former South Loyalty basin thrust on the continental basement of the 52 Norfolk Ridge at the end of the Eocene subduction. The emplacement of the ophiolitic 53 nappe resulted from the failed subduction of the Norfolk Ridge tip in a NE-dipping 54 subduction zone, which culminated in the obduction of the Loyalty subarc lithosphere 55 ~ 34 Ma ago (Cluzel et al. 2012). 56 The Peridotite Nappe has an extension of about 8000 km² and is mostly exposed in 57 the Massif du Sud, where a thick harzburgite-dunite unit, locally overlain by 58 kilometre-scale lenses of mafic and ultramafic intrusives, crops out. The sequence is 59 believed to represent a crust-mantle boundary that records the onset of Eocene 60 61 subduction in a nascent arc setting (Marchesi et al., 2009; Pirard et al., 2013; Secchiari et al., 2018). Recent geochemical studies have shown that the ultramafic 62 intrusives (i.e. dunites and wehrlites) crystallised from variably depleted melts with 63 island arc basalt affinity, after massive interactions with the underlying harzburgite 64 (Marchesi et al., 2009; Pirard et al., 2013). In contrast, the mafic rocks (i.e. 65 gabbronorites) rather have a cumulate origin (Marchesi et al., 2009; Pirard et al., 66 2013; Secchiari et al., 2018) and derive from crystallization of primitive, non-67 aggregated, ultra-depleted melts showing involvement of a subduction-related 68 component in their source (Secchiari et al., 2018). 69 The harzburgites are also exposed in the northern Tiébaghi massif (Ulrich et al., 2010) 70 or as sparse tectonic klippen in the central part of the island (e.g. Kopeto, Poya, 71 72 Koniambo), where exceptionally fresh peridotites display primary mineral assemblages similar to the more serpentinised rocks of the Massif du Sud. 73 74 The New Caledonia harzburgites bear an overall ultra-depleted composition, inherited from a complex multistage evolution linked to the development of the Eocene 75 subduction system (Marchesi et al., 2009; Ulrich et al., 2010; Secchiari et al., 2019a). 76

Geochemical studies have proposed that the harzburgites formed by high degrees of

fluid-assisted melting (up to 20-25 % in a supra-subduction zone environment, see Marchesi et al., 2009; Ulrich et al., 2010). More recently, the work of Secchiari et al. (2019a) provided further constraints on the evolution of the harzburgites, tracking their history from melting to late stage metasomatism. Accordingly, the harzburgites underwent two partial melting episodes in the spinel stability field: a first melting phase in a MOR setting (15% melting degrees), followed by hydrous melting in a supra-subduction zone setting (up to 18% fluid-assisted melting). Post-melting cooling and re-equilibration at lithospheric conditions was accompanied by interaction with slab-derived hydrous melts bearing an ultra-depleted composition (Secchiari et al., 2019a,b). These metasomatic processes in the harzburgites are indicated by the widespread occurrence of secondary metasomatic phases (i.e. thin films of Al₂O₃-, CaO- poor orthopyroxene, and low Al₂O₃ and Na₂O clinopyroxene), L-MREE and Zr-Hf bulk rock enrichments, as well as by the unradiogenic Nd isotopic ratios shown by some samples (Secchiari et al., 2019a). Compared to the harzburgites from the central and the southern massifs, Tiébaghi samples display a more fertile nature, as indicated by higher trace element concentrations as well as by the occurrence of a small fraction (up to 4 vol.%) of clinopyroxene (see Ulrich et al., 2010; Secchiari, PhD thesis). The main geochemical and petrological features of the spinel and plagioclase lherzolites are thought to reflect moderate melting degrees (8-9%) in a MOR environment, followed by refertilisation by depleted MORB-type melts, yielding plagioclase Iherzolites. The main petrological and geochemical features of these

lithotypes have been described in detail by Secchiari et al. (2016).

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2.1 Sample description

element geochemistry (i.e. major, trace element and Sr-Nd-Pb isotope compositions) 105 were analysed for mass fractions of all PGE, Re, Au, S, Se and Te and ¹⁸⁷Os/¹⁸⁸Os. 106 107 Detailed descriptions of the Iherzolites and harzburgites, including trace element chemistry and Sr-Nd-Pb isotopes, are provided in Secchiari et al. (2016) and Secchiari 108 et al. (2019a, b), respectively. 109 Lherzolite samples are from the Poum and Babouillat areas, while the harzburgites 110 were collected from several outcrops and mine zones along the island: Yaté, Kopeto, 111 Poya, Poro and Tiébaghi (Fig. 1b and Table 1). The Iherzolites include serpentinised 112 (LOI= 6.9 - 10.7 %) spinel and plagioclase lherzolites, while the harzburgites are 113 typically not or only little serpentinised (LOI = 0 - 3%), except for samples YA1, TI1 114 and TI2 (LOI = 6.0 - 9.0 %). 115 Both Iherzolites and harzburgites are low strain mantle tectonites, showing dominant 116 porphyroclastic textures (Fig. S1a-b) and local protomylonite development. Spinel 117 118 Iherzolites have 7-8 vol.-% clinopyroxene and display a typical abyssal-type REE signature. The plagioclase lherzolites show melt impregnation microstructures (Fig. 119 120 S1b) and are slightly enriched in incompatible trace element enrichments (REE, Ti, Y, and Zr) with respect to the spinel lherzolites. Harzburgites are extremely depleted 121 rocks, as highlighted by the general absence of clinopyroxene (with the exception of 122 sample TI2, where clinopyroxene is ~ 4 vol. %, Fig. S1c) and the very low 123 incompatible trace element contents (Secchiari et al., 2019a). The primary mantle 124 paragenesis is composed of olivine, orthopyroxene and spinel. The occurrence of thin 125 films of metasomatic ortho- and clinopyroxene (Fig. S1d) was interpreted as the 126 127 result of percolation by small fractions of subduction-related magmas (Secchiari et al., 2019a-b). 128 Rare sulphide grains with variable size, shape and position have been recognised in 129

the Iherzolites and were analysed for their major element chemistry in samples BA1

In this contribution, seventeen samples of peridotites fully characterised for lithophile

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and POU2 (see Table S1). Small (40 to 100 μ m x 30 to 80 μ m) sulphide inclusions (Fig. S2a-e), bearing polyhedral or spherical shape, have been observed within olivine and pyroxene porphyroclasts. Interstitial sulphide grains occur as polyhedral blebs (Fig. S2f), up to 300 x 100/150 μ m in maximum dimensions, and are generally located at olivine-pyroxene grain boundaries. Sulphide composition is relatively homogeneous (Table S1 and Fig. S2), with Ni-poor (Fe/Ni=1.4-2.5) monosulphide solid solution (Guo et al., 1999) being the most abundant phase. Ni-rich (Fe/Ni =0.7-0.8) pentlandite has also been identified in the sample POU2. The sulphides frequently show lamellae and rims made by Fe-oxides/hydroxides due to desulphidation reactions.

3. Analytical methods

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143 3.1 HSE and chalcophile elements

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- Seven lherzolites and ten harzburgites (including four duplicates) have been analysed in the geochemistry laboratory at Freie Universität for HSE, S, Se, Te mass fractions in whole rocks and ¹⁸⁷Os/¹⁸⁸Os.
- Detailed procedure descriptions have been given in previous work from this laboratory (e.g. Fischer-Gödde et al., 2011; Wang et al., 2013; Wang and Becker, 2013). The
- About 2.5 g of sample powder was weighed into 90 ml quartz glass digestion vessels

methods will only be briefly summarized below.

- and spiked with mixed 191 Ir- 99 Ru- 194 Pt- 105 Pd, 77 Se- 125 Te, 185 Re- 190 Os and 34 S solutions.
- 153 Then, 5 ml 14 mol/L, N_2 -bubbled HNO $_3$ and 2.5 ml 9 mol/L HCl were added. The
- vessels were immediately sealed with Teflon tape and samples were digested for 16 h
- at 320°C and 100 bar. After digestion, osmium was extracted from the reverse aqua

regia into chloroform, back extracted into HBr (Cohen and Waters, 1996), and further 156 purified by micro distillation from a H₂SO₄-dichromate solution into 15 μl of HBr (Birck 157 et al., 1997) 158 Osmium isotopes were determined as OsO₃ in negative mode using the Thermo 159 Finnigan Triton TIMS, using a secondary electron multiplier. Signal intensities of the 160 spike isotope 190 Os of samples were $\sim 150,000-500,000$ cps. Standard runs with 161 different amounts of Os on the filament (10 pg and 100 pg) were also run in between 162 the studied samples, yielding an average value of 0.1139 ± 0.0002 (2 s. d., n = 24) 163 for 100 pg loads. Two hundreds scans were collected in each measurement for high-164 Os samples, while at least 120-140 scans were obtained for the low-Os samples. Raw 165 166 data were corrected for isobaric OsO₃⁻ interferences, mass fractionation using the ¹⁹²Os/¹⁸⁸Os ratio of 3.08271, contributions from the Os spike solution and blank 167 contributions. $^{187}\text{Os}/^{188}\text{Os}$ were finally adjusted relative to the mean of the Os 168 standard. The oxygen isotope compositions used for the oxide correction of Os oxide 169 molecules were ${}^{18}\text{O}/{}^{16}\text{O}$ of 0.00204 and ${}^{17}\text{O}/{}^{16}\text{O}$ of 0.00037 (Nier, 1950). 170 About 50% of the digestion solution was used for separation of the HSE fraction and 171 172 about 30% for S-Se-Te separation. Chemical separation of the HSE fraction from the matrix was performed on columns filled with 10 ml of pre-cleaned Eichrom 50W-X8 173 (100-200 mesh) cation exchange resin (Fischer-Gödde et al., 2011). During 174 separation, the HSE fraction was collected in 14 ml 0.5 mol/L HCl-40 vol.% acetone 175 mixture. After the volume of the solution has been reduced to about 2 ml it was 176 analysed for Au, Re, Ir and Pt. In order to remove interfering Cd, the remaining 177 solution was further purified in 0.2 mol/L HCl on 3 ml Eichrom 50W-X8 (100-200 178 179 mesh) resin. The collecte dsolution was evaporated to near dryness and the residue was taken up in 0.28 M HNO₃ for ICP-MS analysis. The analyses were carried out 180 using a single collector Element XR instrument. We used either a Scott-type spray 181

chamber (Re, Ir, Pt, Au) or an Aridus-I desolvator (Ir, Ru, Pt, Rh, Pd) at an oxide 182 formation rate of $CeO^+/Ce^+ < 0.004$. 183 A two-step ion exchange chromatography method was used for separation of S, Se 184 and Te (see Wang et al., 2013). Sulphur measurements were performed on the S-Se 185 fraction at medium mass resolution mode on the Element XR. Selenium and Te were 186 measured using a double pass Scott type glass spray chamber at low mass resolution 187 mode on the Element XR, combined with a hydride generation sample introduction 188 system by reacting the sample solution with 1 g/100 g NaBH₄ in 0.05 mol/L NaOH 189 (see Wang et al., 2013 for details). 190 For each batch of analysis, one procedural blank has been used. Procedural blanks 191 192 yielded the following mean values (\pm 1 s.d., n = 4-5): Re = 2.5 \pm 2.0 pg; Os = 0.5 \pm 0.3 pg with 187 Os/ 188 Os ratios of 0.14 ± 0.03; Ir = 15 ± 5 pg; Ru = 45 ± 14 pg; Rh = 193 $24 \pm 22 \text{ pg;}$ Pt = $23 \pm 29 \text{ pg;}$ Pd = $640 \pm 330 \text{ pg;}$ Au = $4 \pm 2 \text{ pg;}$ Te = $1.1 \pm 0.8 \text{ ng;}$ 194 Se = 2.3 ± 0.8 ng; S = 2.8 ± 0.7 μ g. Samples were corrected for total procedural 195 196 blanks using the mean values. Blank corrections for Re are negligible for most of the analysed samples ($\leq 0.3 - 0.8$ %), but more significant for the harzburgites ($\sim 4 - 8$ 197 198 %). Blank corrections for Pt and Pd are again negligible for the lherzolites ($\sim 0.2 - 0.3$ %) a few percents for the harzburgites (~ 0.4 - 4 %, with the exception of KPT2, 199 KPT5 and PO3 for which the correction for Pt is $\sim 11 - 36$ %). Blanks of Os, Ir, Ru and 200 Rh are insignificant for most of the samples (≤ 0.4 %) but higher for the most 201 depleted harzburgites, i.e. KPT2, KPT5 (~ 2 - 7 % for Os, Ir, Rh) and PO3 (~ 9 % for 202 Os and Ir, 13 % for Rh,). Blank corrections for S, Se and Te in Iherzolites range 203 between 1 - 1.7 % (S - Se) and 3-6% (Te), while for harzburgites blank corrections 204 205 for these elements strongly affected the obtained results (corrections $\sim 10 - 26$ % for 206 S and up to 40 - 80% for Se and Te), given the very low measured abundances.

3.2 Oxygen fugacity

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Iron oxidation state in studied spinel was measured using flank approach designed for the JEOL JXA-8200 electron microprobe analyser at Freie Universität on a collection of natural and synthetic spinel standards characterized for their Fe³⁺/ΣFe at IPGG RAS (St. Petersburg) using Mössbauer spectroscopy (Goncharov, 2018). The analytical procedure was similar to experiments performed over the last decades to study Fe³⁺/ Σ Fe in mantle garnets after procedure developed by (Höfer and Brey, 2007). The flank positions were calculated from the difference spectrum of almandine-andradite for the spectrometer in wavelength range related to FeLa and FeLβ lines. FeLa and FeLß intensities were collected at a wavelength of flank lines for 300 seconds each with 3 repetitions in the core and rim parts of 4-5 spinel grains within one thin section. Measurement conditions were 15 kV and 60 nA using TAP crystal for intensities at flank positions and with remaining 4 spectrometers were measured chemical composition at the same spot simultaneously. Averaged FeLa and FeLB intensities for one sample were used to calculate iron oxidation state of spinel from the equation obtained after standardization, where FeL\(\beta\)/FeLa ration and FeO total content correlate with $Fe^{3+}/\Sigma Fe$ measured by Mössbauer spectroscopy. Owing to the lack of an appropriate geobarometer for spinel peridotites, the assumed equilibrium pressure for oxygen fugacity calculations has been set at P = 1.5 GPa. Equilibrium temperatures have been calculated using coexisting minerals and the Cain-orthopyroxene geothermometer of Brey and Köhler (1990) and olivine-spinel geothermometer of Li et al. (1995).

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4. Results

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HSE and chalcophile element compositions of the New Caledonia Iherzolites are reported in Table 1 and displayed in Fig. 2, 3, 4. Spinel and plagioclase lherzolites are relatively homogeneous in terms of HSE, Se, Te concentrations, abundance patterns and Os isotopic compositions, with plagioclase-bearing samples showing indistinguishable patterns from those of spinel lherzolites. The concentrations of the highly siderophile and chalcophile elements are in the range of those observed for modern abyssal and ophiolitic peridotites, displaying good correlation for Ir group PGE (IPGE, e.g., Os vs. Ir and Ir vs. Ru) and more dispersed variations for the Pt group PGE (PPGE, Fig. S3). In primitive mantle (PM) normalised concentration diagrams (Fig. 3), the Iherzolites exhibit flat or gently sloping patterns with similar PMnormalized PGE concentrations and depletions in Au (except for sample BAB2B) and Re compared to the PGE (with the exception BAB2B for Au and POU1A and POU3 for Re). Overall, absolute contents of the PGE are similar or slightly lower than primitive mantle (PM) values (Becker et al., 2006; Fischer-Gödde et al., 2011), overlapping the field of the abyssal peridotites and peridotite tectonites from continental settings (e.g., Fig. 2 and Becker and Dale, 2016). Ru/Ir and Pd/Ir ratios are suprachondritic, as observed for other mantle lherzolites (e.g. Lorand et al., 1999; Rehkämper et al., 1999; Luguet et al., 2003; Becker et al., 2006). Initial ¹⁸⁷Os/¹⁸⁸Os ratios calculated at 53 Ma (i.e. the inferred age of initial magmatism in the subduction system, e.g. Cluzel et al., 2006) vary from chondritic to slightly suprachondritic (0.1273-0.1329, Fig. 4a), corresponding to $\gamma Os_{(53Ma)}$ of 0.5 to 4.9. These values overlap with data of abyssal peridotites and orogenic peridotites, but tend to be somewhat higher than for other mantle lherzolites bearing comparable

- depletion degrees (Fig. 4a and 4b). ¹⁸⁷Re/¹⁸⁸Os ranges from subchondritic to slightly
- suprachondritic values (0.186-0.525, see Fig. 4c).
- Se and Te are positively correlated in the Iherzolites (Fig. 5a) and range between 54 -
- 262 91.3 ng/g and 7.3 13.8 ng/g, respectively. Se/Te ratios (5.9-7.1) are slightly lower
- 263 than the PM value and similar to the data previously obtained on depleted lherzolites
- 264 (Wang and Becker, 2013). Se and Te do not display any correlation with PGE
- abundances, with the exception of Te, which shows a weak correlation with Pd (Fig.
- 266 5b). S contents are high and variable (202 1268 μ g/g) compared to unserpentinised
- peridotites, leading to high S/Se ratios (2703-16289, Fig 5c).

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4.2 HSE and chalcophile elements in harzburgites

- On the basis of HSE behaviour and Os isotopic compositions (Fig. 2, 3 and 6, Table 1),
- 272 the studied harzburgites can be grouped into two different sub-types: type-A and
- 273 type-B.
- 274 Type-A harzburgites (TI1, TI2, PO4 and YA1) are characterised by notably lower mass
- 275 fractions of all PGE and most other chalcophile elements (Figs. 2, 4) and more
- 276 fractionated HSE and chalcophile element patterns (Fig. 6a) compared to the
- 277 Iherzolites. Mass fractions of the PGE in these harzburgites are 1.07-2.07 ng/g for Os,
- 278 0.50-1.14 ng/g for Ir, 1.53-2.52 ng/g for Ru, 0.19-0.27 ng/g for Rh, 2.42-2.70 ng/g
- for Pt and 0.49-0.57 ng/g for Pd. Among the sub-group A, sample YA1 displays
- distinct PGE abundances, showing much lower Os, Ir, Pt, Pd contents (0.55 ppb for
- Os, 0.30 ppb for Ir, 0.27 ppb for Pt and 0.26 ppb for Pd), with exceptions for Ru, Rh,
- 282 Au and S.
- 283 HSE and chalcophile element diagrams of type-A harzburgites display fractionated
- 284 patterns, with concentrations decreasing towards Re. Os and Ru are enriched

compared to Ir, leading to correlated suprachondritic Os/Ir and Ru/Ir ratios (Os/Ir=2.9-7.9; Ru/Ir=1.8-2.1, Fig. 4). Pt and Au generally show positive spikes, more pronounced for Au, with the exception of YA1, which displays a negative Pt anomaly. Pd contents are low (< 0.1 PM values) and constant for PO4, TI1, TI2, with Pd/Ir showing subchondritic ratios for all studied samples. Positive correlations are observed between IPGE (Fig. 2a-b-c) and Pt-Ir (not shown), and, somewhat surprisingly, between IPGE and the fertility indicators (i.e. Al_2O_3 and CaO, not shown). Mass fractions of Te, Se and S are low, often close to or below the detection limit, again with the exception of the harzburgite YA1. For all type-A harzburgites, Re concentrations are very low (about 0.02 ng/g), leading to subchondritic ¹⁸⁷Re/¹⁸⁸Os ratios (0.045 to 0.196, the latter value also reflecting the low Os concentrations in YA1). Os isotopic compositions are subchondritic to chondritic (0.1203-0.1266, corresponding -5 $\leq \gamma_{Osi(53 \text{ Ma})} \leq -0.1$) and do not define any correlation with ¹⁸⁷Re/¹⁸⁸Os or incompatible element depletion indices (i.e. Al₂O₃, see

Fig. 4).

Type-B harzburgites comprise very fresh samples from Kopeto (KPT2, KPT3, KPT5), Poro (PO3) and Poya (PY1) massifs. Compared to type-A harzburgites, these samples have much lower HSE abundances and display variable and strong fractionations among PGE and more incompatible chalcophile elements (Fig. 2 and 6b). In detail, Os, Ir and Pt show positive correlations (Fig. 2) and are strongly depleted compared to Ru, Rh and Pd (Os= 0.003-0.071 ppb, Ir= 0.015-0.079 ppb, with Os/Ru=0.01-0.26 and Ru/Ir= 2.5-20). For sample PY1, Pt is enriched relative to IPGE, Rh and Pd (Pt/Rh=7.7; Pt/Pd=5.5). Pd, Re and S-Se-Te have similarly low normalized abundances, with chalcophile elements often close to or below the detection limit. Au

exhibits positive spikes for PY1, KPT3 and PO3 samples and tends to be more enriched

than similar incompatible chalcophile elements (i.e. Pd and Re).

Measured ¹⁸⁷Os/¹⁸⁸Os ratios vary from chondritic to suprachondritic (0.1273-0.1534)

and are coupled with high and variable 187 Re/ 188 Os (1.62-32). Initial Os isotopic

compositions calculated at 53 Ma range from depleted to slightly suprachondritic

315 values (0.1181-0.1365, $-7 \le \gamma Os_{(53Ma)} \le 3$).

Replicate analyses of samples PY1, KPT2, KPT5, PO3 yield quite similar results for Ru

(< 5 % relative deviation, except for sample KPT2) and Au (~ 6% for PY1 and KPT5)

and acceptable results for Pt for PY1-KPT5 (11.5 - 14.0 % relative deviation). Values

appear much more scattered for Os and Re (RSD> 30%) and less dispersed for Rh

and Pd (7≤RSD%≤32). The relative deviation of chalcophile elements is more limited,

321 mostly < 15%.

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322 The large variations of mass fractions of HSE and chalcophile elements in duplicate

samples reflects the very low mass fractions of these elements combined with the

inhomogeneity in the distribution of HSE carrier phases in gram-size quantities of rock

powder, an issue that has already been recognised in peridotitic rocks (e.g. Becker et

326 al., 2006; Luguet et al., 2007).

328 4.3 Oxybarometry

330 Geothermometric estimates and calculated oxygen fugacity values are reported in

Table 2. Equilibrium temperatures calculated for porphyroclastic assemblies with the

Ca-in-orthopyroxene thermometer (Brey and Köhler, 1990) range between 930-

1130°C, with samples YA1 and TI2 displaying the lowest temperatures (930°C-980°C,

see Table 2). Olivine-spinel geothermometry (Li et al., 1995) yields considerably lower

equilibration temperatures (815-940°C).

Oxygen fugacity estimated with the Wood (1990) method yielded values close to the FMQ (fayalite-magnetite-quartz) buffer for all the harzburgites ($0.16 \le \Delta logFMQ \le 0.71$). Similar estimates ($0.18 \le \Delta logFMQ \le 0.76$) are obtained using temperatures calculated with the olivine-spinel geothermometer. No significant difference can be observed between sub-group A and B harzburgites. Remarkably, the highest oxygen fugacity values are recorded by two harzburgites belonging to sub-group A (YA1 and TI2), which are characterised by lower degrees of depletion, as inferred from trace element modelling (see Secchiari et al., 2019a).

By contrast, lherzolite BA1 lherzolite indicates much more reducing oxygen fugacity conditions ($\Delta logFMQ \leq -3.73$).

5 Discussion

5.1 HSE and Re-Os systematics of the Iherzolites

Major element composition and lithophile trace element chemistry of spinel lherzolites indicate moderately depleted compositions, inherited from low partial melting degrees (8-9%) of a DMM source, whereas plagioclase lherzolites originated through reactive melt percolation of spinel lherzolites by highly depleted, incremental melt fractions of a DMM source in the shallow lithosphere (Secchiari et al., 2016). In the following sections the processes that may have affected HSE and Os isotopic signature of the New Caledonia lherzolites will be discussed: low temperature alteration, in particular serpentinisation, partial melting and the role of melt infiltration and chemical disequilibrium of the HSE in mantle rocks.

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Serpentinisation is a widespread process of hydrothermal alteration in ultramafic lithologies. However, its influence on HSE behaviour has not been investigated with much detail, despite some authors have proposed it as a possible cause for ¹⁸⁷Os ingrowth and Re variations in the upper mantle (Snow and Reisberg, 1995; Walker et al., 1996; Standish et al., 2002). Recent experimental studies have shown that during serpentinisation the formation of secondary sulphides, Fe-Ni alloys and native metals (Au-Cu) is promoted by reducing f_{02} conditions (Klein and Bach, 2009; Foustoukos et al., 2015) and thus, with the exception of Au, HSE may be retained in the host rock. Comparison of partially serpentinised and unserpentinised peridotites displaying similar major element features supports the notion that at least PGE ratios are little changed by moderate to strong serpentinisation (Becker and Dale, 2016). The New Caledonia Iherzolites underwent intermediate serpentinisation (LOI = 6.4 to 10.7 %, see paragraph 2.1 and Table 1), which had limited effects on the budget of fluid immobile moderately incompatible lithophile trace elements in these rocks (Secchiari et al., 2016). Notably, PGE contents and ratios in the lherzolites are similar to other unaltered and serpentinised lherzolites from the modern oceans and ophiolitic complexes (see Fig. 2 and 3 e.g.; Snow et al., 2000; Luguet et al., 2001, 2004; Pearson et al., 2004; Alard et al., 2005; Becker et al., 2006; Fischer-Gödde et al., 2011; Becker and Dale, 2016). This observation supports the hypothesis that PGE abundances are comparable in fresh and variably serpentinised ultramafic rocks (Becker et al., 2006; Liu et al., 2009; Fischer-Gödde et al., 2011; Marchesi et al., 2013; Becker and Dale, 2016), implying that serpentinisation results in minor changes in PGE ratios.

By contrast, the possible influence of serpentinisation on Au and Re is more difficult to evaluate, as no study has systematically investigated its effect on the behaviour of the aforementioned elements. In the lherzolites from New Caledonia, Au displays similar normalized concentrations as Re and, with the exception of a few samples, both elements are depleted relative to Pd, Te and Se. Au abundances tend to be somewhat lower than abundances in other lherzolites with similar major element composition (Fig. 3). Although the compositions can be entirely explained by magmatic fractionation processes (see subsequent chapters), minor losses of Au due to hydrothermal alteration cannot be ruled out (e.g. Lorand et al., 1999). The lack of correlation between Au and Al₂O₃ (not shown) could be a hint that Au abundances may have been affected by a combination of magmatic processes and serpentinisation (i.e. Fischer-Gödde et al., 2011). Rhenium is slightly depleted compared to the PGE for most of the studied lherzolites, but displays higher concentrations than other mantle Iherzolites (Fig. 3). In addition, Re contents do not correlate with LOI and Re/Os ratios cover the range generally reported for moderately depleted mantle rocks. The samples with the lowest Re contents display the highest LOI values, suggesting that no significant quantities of Re were added during the interaction with seawater during serpentinisation. Likewise, the chondritic to slightly suprachondritic ¹⁸⁷Os/¹⁸⁸Os cannot be ascribed to serpentinisation, as unrealistically high water-rock ratios (~ 10^3 - 10^4) would be required in order to perturb the whole rock 187 Os/ 188 Os at the % level or higher (e.g. Becker and Dale, 2016). The Iherzolite data also shows mass fractions of Se and Te and Se/Te that are similar to values in unserpentinised lherzolites (e.g., Wang and Becker, 2013). In contrast sulfur in most lherzolites from New Caledonia shows much higher concentrations than typical for peridotites, which is readily explained by contamination with seawater-derived sulfur during serpentinisation.

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We thus conclude that the HSE (perhaps with the exception of Au), Se, Te and Re-Os signature of the lherzolites offer no conclusive evidence that serpentinisation and associated reactions affected these elements in a noticeable way.

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5.1.2 Partial melting and chemical disequilibrium of the HSE in the mantle

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Spinel and plagioclase Iherzolites exhibit comparable HSE contents and patterns, similar to other Iherzolites from oceanic or continental settings that underwent low to moderate degrees of melt extraction (Fig. 2 and 3). Partial melting has often been invoked as a possible cause for HSE and ¹⁸⁷Os/¹⁸⁸Os variations in mantle rocks (e.g. Reisberg and Lorand, 1995; Meisel et al., 2001). Studies of the behaviour of the HSE during mantle melting and their abundances in mantle rocks have supported the hypothesis that HSE concentrations in residual peridotites result from sulphide-silicate partitioning during magmatic processes (i.e. Becker and Dale, 2016; Brenan et al., 2016 and references therein) and that at temperatures relevant for mantle processes homogeneous sulphide liquid and, in special cases, sulfide solid solutions, coexist in equilibrium with silicate melt, olivine, pyroxenes and an Al-rich phase (e.g. Rehkämper et al., 1999; Mungall and Brenan, 2014; Brenan et al., 2016). Experimentally determined sulphide melt-silicate melt partition coefficients (D^{sulph/sil}) for PGE have been shown to be high and constant (10⁵ to 10⁶, e.g. Mungall and Brenan 2014; Brenan et al. 2016), while Au shows slightly lower D^{sulph/sil} (~10⁴). Therefore, up to moderately high degrees of melting, PGE behave as compatible elements and their inter-elemental ratios remain similar as long as sulphide is present in the mantle residue. By contrast, Re is much less chalcophile (D^{sulph/sil}~300-800, e.g. Fonseca et al., 2007; Brenan, 2008) and is expected to become more quickly depleted in the residual mantle.

Major element compositions of sulphides in the spinel lherzolites (Table S1) are 438 consistent with a residual origin after incongruent melting processes (e.g. Bockrath et 439 al., 2004; Ballhaus et al., 2006). The occurrence of homogeneous monosulphides also 440 441 suggests relatively high cooling rate after the melting event. Overall, the studied lherzolites are characterised by flat to gently sloping PGE 442 patterns, with similar PM-normalized abundances, no PPGE fractionation and nearly 443 constant ratios for IPGE (i.e. Os/Ir, Ru/Ir). By contrast, Au and Re display the 444 strongest depletion. These features imply that HSE, with the exception of Au and Re, 445 exhibit a similar compatible behaviour during mantle melting, as expected for low to 446 moderate melting degrees in presence of residual sulphide melt. This observation is 447 448 consistent with the previous estimates obtained through geochemical modelling (Secchiari et al., 2016) and with the occurrence of a residual subsolidus sulphide 449 assemblage in spinel lherzolites. 450 Although the PGE patterns are nearly flat, with only slight depletion of Pd in a few 451 samples, the depletion of Au and Re, the range of chondritic to slightly suprachondritic 452 γ^{187} Os_i and the higher mass fractions of Se and Te compared to Re and the other HSE 453 suggest a multi-stage history of the lherzolites. Notably, γ^{187} Os_i do not correlate with 454 mass fractions of incompatible HSE such as Re, Re/Os nor with fertility indicators (Fig. 455 4), as was observed in some other suites of lherzolites (Becker and Dale, 2016). 456 The Os isotopic signature may be a pre-existing feature of the mantle source, i.e. 457 already present before the recent melt extraction event (Secchiari et al., 2016). This 458 is supported by the dispersed Os isotopes-fertility indicators trends, as well as by 459 some old model ages recorded by our lherzolite samples ($T_{MA}(PM) = 0.4-0.8$ Ga, see 460 461 Table 1). In addition, the remarkable absence of magmatic Cu-Fe-rich sulphides (e.g. see Lorand et al., 2013) argue against a recent, post-melting sulphide addition. We 462 thus speculate that the bulk HSE, Se, Te and Os isotope compositions of the 463

lherzolites are the result of partial melting event which affected a mantle source previously characterised by an heterogeneous sulphide population including both residual and magmatic sulfides precipitated along grain boundaries by infiltrating melts (Burton et al., 1999; Lorand et al., 1999; Alard et al., 2000, 2002).

Sulfur mass fractions are variable in the Iherzolites from New Caledonia and typically

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5.2 Sulphur, Se and Te behaviour in the New Caledonia Iherzolites

much higher compared to estimates of the depleted MORB mantle source (DMM ~ 150-200 ppm, Mathez, 1976; Salters and Stracke, 2004) and mass fractions of S in unaltered lherzolites (e.g., Wang and Becker, 2013). In addition, sulphur does not correlate with fertility indicators (i.e. Al₂O₃), as commonly observed in unserpentinised mantle tectonites (e.g., Lorand and Alard, 2010; Wang and Becker, 2013). The high S concentrations and the lack of correlation with melting indicators suggest that S was added late in the evolution of the rocks. The sulphur budget of mantle peridotites can be strongly influenced by seawater-rock interaction, because of the high sulphate content of seawater, leading to hydrothermal sulphides and sulphate precipitation (Alt and Shanks, 1998). We note that the major element chemistry indicates a residual origin for the sulphide phases of the lherzolites (see paragraph 4.1 and 5.1.2). Hydrothermal sulphides or sulphates could not be identified. Recent geochemical works have demonstrated the role of serpentine as a sink of S under various oxidation states (S^{2-} , S^{-} , S^{0} and S^{6+} , Debret et al., 2017). These studies have shown that S concentrations can be anomalously high in serpentinised peridotites (up to 1 wt.%, see Alt et al., 2003), as S can be accommodated in serpentine minerals, accounting from 60 to 100% of the sulphur budget of the peridotite (Debret et al., 2017). In situ analysis of serpentine minerals have revealed that S can be hosted in nano-phases associated with serpentine or trapped either via

Si substitutions in the tetrahedra, or as a sulphate ion in the network of the 490 tetrahedral sheet of serpentine minerals (Debret et al., 2017). 491

The addition of S during serpentinization is also reflected in the high suprachondritic S/Se ratios (up to 16500) and the excellent correlation observed between S concentrations and S/Se ratios (see Fig. 5c). Despite the strong S enrichments, Se and Te display 'normal' concentrations and Se/Te ratios are in the range of other Iherzolites (see Wang and Becker, 2013). These data confirm that Se-Te contents and ratios were not significantly impacted by serpentinisation, as previously observed for other peridotites that experienced low to moderate serpentinisation degrees (e.g. Wang and Becker, 2013; Marchesi et al., 2013).

Moreover, Se-Te show a good correlation between each other (Fig. 5a), implying that they are controlled by the same mineral phases. In mantle peridotites, Se can replace S as a chalcogen anion within the crystalline structure of sulphides (e.g. Bulanova et al., 1996; Hattori et al., 2002; Helmy et al., 2010) or can form Se-rich micro phases, while Tellurium, owing to its semi metal nature, tends to partition between sulphides and late exsolved micrometric tellurides (Pt, Pd, Te, As, Bi phases). The latter are thought to crystallise at low temperatures during cooling, once sulphide melt becomes saturated with respect to Te (Luguet et al., 2004; Lorand et al., 2008; Lorand and Alard, 2010). In the Iherzolites, Se and Te do not correlate with melting indicators, but Te displays a rough positive correlation with Pd (Fig. 5b), which suggests that the sulphide melt-bulk silicate partition coefficient of Te should be between Se and Pd (e.g., Figs. 3, 4, 5).

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5.3 Type-A harzburgites: highly siderophile element systematics of a residual sub-arc mantle section

The distinct HSE patterns and Os isotopic signature recorded by the New Caledonia 516 harzburgites hint that they experienced a different history compared to the northern 517 Iherzolites. 518 519 Three of the four samples belonging to the sub-group A (TI1, TI2, PO4) show similar HSE patterns and chalcophile elements depletion, indicating that the same processes 520 contributed to the HSE and chalcophile element budget of these rocks. The low 521 chalcophile element concentrations, close to or below the detection limit, coupled with 522 low Pd/Ir ratios and subchondritic ¹⁸⁷Os/¹⁸⁸Os_i, point out that type-A harzburgites are 523 residues of high degree of melt extraction, where sulphides melts must have been 524 nearly completely dissolved in coexisting silicate melt. 525 Experimental studies have in fact predicted a compatible behaviour for all the PGE 526 during mantle melting as long as sulphide is retained in the peridotite (Mungall and 527 Brenan, 2014). Depending on the initial S content of the mantle rocks, ~12-20% of 528 melting is required for sulphide exhaustion (Lorand et al., 1999; Luguet et al., 2007; 529 530 Brenan et al., 2016). As melting proceeds, sulphides are progressively dissolved into the melt and the PGE become concentrated in the residual sulphide melt (Mungall and 531 532 Brenan, 2014). Provided that chemical equilibrium is attained, increasing degrees of partial melting should slightly increase whole rock PGE contents, but element ratios 533 should remain almost constant. At the point when sulphide is completely removed, 534 IPGE and Pt are accommodated in metallic alloys, while Re, Au and Pd mass fraction 535 should become extremely low, as these elements are not hosted in any residual 536 mantle phase (Mungall and Brenan, 2014). Hence for high melting degrees, the 537 abundances of the HSE in the residue should reflect mineral-melt partitioning and the 538 539 P-T and f_{02} -dependent solubility of Pt and IPGE alloys in silicate melt (Fonseca et al., 540 2011, 2012; Mungall and Brenan, 2014; Brenan et al., 2016). The IPGE-PPGE fractionation and the resolvable fractionations between specific PGE 541 displayed by the type A harzburgites bear witness of the high melting degrees, which 542

resulted in the formation of a S-free mantle residue. The fractionated Os-Ir-Ru-Rh segments of HSE patterns and the positive Pt anomalies in type A harzburgites are likely carried by residual sulphides (i.e. laurite) and metallic alloys (Os-Ir and Pt-Ir, see Lorand et al., 1999; Luguet et al., 2001, 2007). These latter are thought to precipitate from sulphide melt shortly before the complete exhaustion of sulphide (Mungall and Brenan, 2014) or immediately after sulphide consumption, due to $f_{\rm S2}$ lowering and diminished metal-sulphide complexation in the silicate melt (Fonseca et al., 2012). The variable, but broadly systematic inter-element fractionation among the IPGE (high Os/Ir and Ru/Ir), high Ru/Rh and the variable Pt anomaly relative to Rh and Pd indicate the different proportions of residual Ir-Pt alloys in different samples and preferred retention of Ru and Os relative to Ir in the residual PGE alloys (e.g. Brenan and Andrews, 2001; Fonseca et al., 2012). The HSE fractionations of type A harzburgites are different from HSE patterns of modern harzburgites from MOR environments (Fig. 6a), as the latter are characterised by flat or weekly fractionated Os-Ir-Ru triplet and they rarely display positive Pt spikes (Snow and Schmidt, 1998; Luguet et al., 2001, 2003). By contrast, HSE elemental fractionations of type-A harzburgites are similar to patterns of some arc xenoliths (Saha et al., 2005; Liu et al., 2015; Scott et al., 2019) or ophiolitic peridotites bearing a supra-subduction zone signature (see Büchl et al., 2002, 2004; O'Driscoll et al., 2012). Similar fractionations have been reported for some mantle xenoliths from the Chatam Islands (New Zealand) and from some other areas (Pearson et al., 2004). On the other hand, geochemical modelling based on lithophile incompatible elements has shown that the extreme depletion in trace element contents displayed by the New Caledonia harzburgites was achieved through a polyphase evolution, including a first melting event in a mid-ocean ridge setting, followed by fluid-assisted melting reaching clinopyroxene exhaustion after involvement in a subduction system (see Secchiari et al., 2019a). Such high melting degrees are permissible in supra-

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subduction zone environments, where hydrous conditions at relatively low pressures
can produce melt fractions substantially exceeding 20% without invoking extremely
high temperature (e.g. see Ulmer, 2001). Likewise, the observed LREE and FME (Sr,
Ba, Pb) enrichments coupled with variable Pb isotope compositions of the Type A
harzburgites may be explained by syn- and post-melting interactions with different
subduction-related components, possibly aqueous fluids and melts originated in the
forearc setting (Secchiari et al., 2019 a,b).
We thus conclude that the HSE and chalcophile element signature displayed by TI1,

We thus conclude that the HSE and chalcophile element signature displayed by TI1, TI2 and PO4 predominantly reflect high degrees of melt extraction, presumably in a supra-subduction zone environment. The positive Pt spikes suggest that Pt-rich alloys were stable in the mantle residue and were only in part dissolved in the melt during melt extraction.

The enrichments of Au are modest (0.2-1.3 ng/g Au) and must be related to fluid overprint, either from slab-derived fluids (McInnes et al., 1999; Kepezhinskas et al., 2002) or from low-T alteration, e.g., during obduction (e.g. Snow et al., 2000).

The harzburgite YA1 shows higher S and Se concentrations (Fig. 5a), which, considering the significant LOI value of 6.83%, could be related to serpentinisation and precipitation of secondary sulphides. The strongly fractionated HSE pattern and the low concentrations of the incompatible HSE (i.e. Pd, Re) indicate that the HSE budget of YA1 is also governed by melting, as for TI1, TI2 and PO4 harzburgites. The low Os, Ir and Pt concentrations, are much closer to the values reported for type-B harzburgites (see Table 1 and Fig. 5b). Sample YA1 can thus be seen as transitional between type-A and type-B sub-group.

5.4 Origin of type-B harzburgites – strong depletion followed by subduction zone metasomatism?

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Type-B harzburgites mostly occur in the central massifs, however one sample (PO3) 598 has also been identified in the eastern zone, close to the area where one type-A 599 600 harzburgite (PO4) was sampled. Type-B harzburgites display remarkably different HSE and Re/Os behaviour compared to type-A sub-group (Fig. 2 and 6), indicating that 601 different processes need to be invoked in their genesis. 602 Type-B harzburgites show low abundances of incompatible chalcophile elements, i.e. 603 Pd, S, Se and Re, with values in the range of type-A harzburgites. In principle such 604 low concentrations could be reconciled with high melting degrees and sulphide 605 exhaustion in the mantle source. The strong depletion of Os, Ir and Pt, (Fig. 6b) 606 coupled with slightly subchondritic to suprachondritic ¹⁸⁷Re/¹⁸⁸Os (0.23 to 37) and 607 ¹⁸⁷Os/¹⁸⁸Os (0.1239 to 0.302) are remarkable and do not occur in residual 608 harzburgites from convecting mantle, as represented by abyssal peridotites. 609 The HSE in type-B harzburgites share similarities with some xenoliths from arc 610 settings such as low Os contents associated with chondritic to suprachondritic 611 ¹⁸⁷Os/¹⁸⁸Os (Brandon et al., 1996, 1999; Saha et al., 2005; Widom, 2011). These 612 613 features have been ascribed to interactions with subduction zone fluids (i.e. fluids 614 from subducted altered oceanic crust and/or its sedimentary cover), which may have induced sulphide breakdown in the residual arc mantle, locally modifying the Os 615 isotopic signature of the mantle (Wisdom et al., 2003). Such qualitative observations 616 have also been supported by experimental works, that highlighted the critical 617 influence of oxygen fugacity (f_{02}) on sulphide and alloy stability (e.g. Andrews and 618 Brenan, 2002; Fonseca et al., 2011, 2012; Mungall and Brenan, 2014). 619 620 Moreover, the strongly fractionated, IPGE-depleted, HSE patterns displayed by type-B harzburgites closely resemble those observed in some refractory harzburgites and 621 replacive dunites that underwent interaction with S-undersatured melts in some 622 ophiolites (Büchl et al., 2002; Lorand et al., 2004). 623

As metasomatism by subduction fluids and hydrous melts has been proposed for the New Caledonia harzburgites based on isotopic and incompatible element studies, we have determined oxygen fugacities on a set of five harzburgites and one lherzolite, in order to test if the different HSE signature of type-B harzburgites could reflect higher f_{02} conditions. As a whole, the harzburgites record more oxidised conditions compared to the Iherzolite (see Table 2). This is consistent with the hypothesis that the Iherzolites represent slightly depleted mantle rocks not directly involved in the Eocene subduction system (Secchiari et al., 2016). However, the harzburgites exhibit f_{02} values close to the FMQ buffer and in the range of the suboceanic mantle (e.g. Bryndzia and Wood, 1990). Also, type A harzburgites tend to be slightly more oxidised than type B harzburgites. The relatively low oxygen fugacity values may be related to the high depletion experienced by the harzburgites. A recent study of Bénard and coauthors (2018) has in fact shown that increasing melt depletion at un-buffered $f_{\rm O2}$ conditions can induce oxygen fugacity variation in the sub-arc peridotites, as $\mathrm{Fe^{3+}}$ is more extensively extracted than Fe^{2+} as melting proceeds, producing residues with lower $Fe^{3+}/\Sigma Fe$ than in the original mantle source. We note that the most depleted harzburgites (i.e. type-B PY1, KPT5 and PO3) in our dataset yield the lowest computed oxygen fugacities. Following this interpretation, we suggest that the harzburgites recorded higher f_{02} conditions compared to the Iherzolites due to fluid-assisted melting and interaction with an oxidising component, but increasing degrees of melt extraction from type A to type B harzburgites possibly led to progressive lowering of f_{O2} . Alternatively, calculated oxygen fugacity values may partly reflect f_{O2} conditions related to the melting conditions of the harzburgites in the suboceanic mantle (Lee et al., 2005). From this perspective, one may speculate that the oxidising capacity of slab-derived fluids or melts was not high enough to erase the memory of the original f_{O2} conditions. For the New Caledonia archipelago, Eocene subduction started close to

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or in correspondence of an active oceanic spreading center, where hot and young (~ 6-9 My old, Cluzel et al., 2016) lithosphere was forced to subduct. In such a context, fluid fluxes from the downgoing slab must have been limited, due to the young age of the subducted material and the intra-oceanic nature of the subduction (Cluzel et al., 2016). In addition, post-melting metasomatism is thought to have occurred through interaction with small fractions (0.5-1%) of boninitic magmas, which may have not been able to maintain their oxidising capacity during percolation through the sub-arc mantle.

In conclusion, the effect of f_{O2} on the different HSE patterns of type A and B harzburgites is unclear, and perhaps insignificant, as their difference in $\Delta log f_{O2}$ is minor. Thus, most likely the stronger fractionation and depletion of some IPGE in type B harzburgites reflects the higher degrees of melting.

5.5 Type-B harzburgites: a broader perspective

Despite being similar in terms of chemistry or mineralogy, type-A and B harzburgites possess distinct HSE signatures. In addition, the HSE signatures of type-B harzbugites have not yet been identified in other mantle tectonites, either from modern oceanic lithosphere or ophiolitic complex. However, similar compositions have been recently reported for some moderately depleted to highly refractory peridotites and mantle xenoliths from New Zealand (Scott et al., 2019). The New Zealand mantle is composed of isotopically heterogeneous mantle fragments with evolutionary histories extending over 2.75 Ga (Os model ages= 0.1-2.75 Ga, with a broad peak at 1.2 Ga), and PGE systematics decoupled from major element compositions (see Scott et al., 2019; Liu et al., 2015). These features have been explained by accretion of Zealandia lithospheric mantle

from amalgamation of genetically unrelated convecting mantle fragments which were 677 swept together beneath the Gondwana subduction margin, variably re-melted and 678 laterally accreted (Scott et al., 2019). 679 680 Among the New Zealand peridotite suites, mantle xenoliths from Lake Moana and Chatam Island show HSE patterns that are similar to our dataset (Fig. 8). Lake Moana 681 Cretaceous xenoliths include cpx-free harzburgites that experienced up to 30% 682 melting, while Eocene-aged Chatam Island harzburgites exhibit a less refractory 683 nature, as attested by the presence of primary clinopyroxene (up to 1.8% modal, 684 Scott et al., 2016). Such depletion degrees were achieved either by plume melting or 685 hydrous melting in an arc setting for the Lake Moana xenoliths, whereas Chatam 686 Island samples are thought to represent fragments of fore-arc lithophere (Scott et al., 687 2016, 2019). 688 Overall, HSE diagrams highlight that New Zealand mantle xenoliths reproduce with 689 good approximation both patterns observed in our harzburgites, namely type-A and 690 691 type-B (Fig. 7). IPGE patterns are broadly sub-parallel, with the New Caledonia harzburgites falling within or at the lower range of values displayed by the New 692 693 Zealand samples, whereas Pt, Pd and Re exhibit much more variability. 694 For most of the New Zealand xenoliths, Ru concentrations are higher compared to the contents in the New Caledonia harzburgites. Higher Ru contents cannot be ascribed to 695 different degrees of melting, as fertility indicators (i.e. Mg#(OI), Cr#(SpI)) indicate 696 similar degrees of melt extraction for both peridotite suites (see Scott et al., 2016, 697 2019; Secchiari et al., 2019a). The increased Ru retention in the New Zealand 698 samples may be reconciled to the higher Cr-spinel content of these lithologies (up to 699 700 2.8%, see Scott et al., 2016, vs. up to 0.8% for our harzburgites). Numerous studies have in fact demonstrated that spinel can be a significant host for Ru (D^{spinel/melt} ~ 20, 701 Capobianco and Drake, 1990). In addition, increase in oxygen fugacity markedly 702 enhances Ru compatibility in Cr-rich spinel (D^{spinel/melt} up to 500 for f_{O2} of -0.5 FMQ, 703

Park et al., 2012), which can accomodate Ru within its crystal lattice (Pagé and 704 Barnes, 2016) or as laurite and/or Ru-rich alloy inclusions (Brenan and Andrews, 705 2001). Recent geochemical works have also illustrated the importance of Ru retention 706 707 in the sub-arc mantle for the HSE signature of arc lavas (Dale et al., 2012; Park et al., 2013). These studies explain the low Ru concentrations and the high Pt/Ru ratios of 708 the volcanic products as related to Ru retention in the mantle source due to the 709 presence of Cr-rich spinel or PGM (see Dale et al., 2012; Park et al., 2013). Likewise, 710 positive Ru anomalies in our type-B harzburgites may reflect the presence of small 711 laurite or Ru-rich phase inclusions, which could have escaped the high melting 712 713 degrees due to the shield effect of spinel. Other HSE (e.g. Pd and Re) in the New Caledonia harzburgites display a wider range 714 of values compared to the New Zealand samples, possibly related to variable 715 proportions of traces of sulfide precipitated from silicate melt. 716

Enrichments of Pd have also been recognised in other sub-arc mantle sections, where 717 they have been attributed to slab-derived fluids metasomatism, due to the high 718 solubility of Pd in aqueous fluids (McInnes et al., 1999; Park et al., 2013). Likewise, 719 720 Re addition in mantle wedge peridotites may be due to Re release in slab derived 721 fluids during dehydration of the mafic portion of the subducting slab (see Dale et al., 2009). Hence, we conclude that Pd-Re (as well as Te-Se-S) re-enrichments in type-B 722 harzburgites may have been facilitated by minor sulfide precipitation from slab 723 derived fluids or melts. 724

By contrast, the widespread negative Pt anomalies may reflect destabilisation of a pre-existing Pt-alloy phase, possibly related to continued melting after alloy saturation (see Mungall and Brenan, 2016), and Pt release into the melt.

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5.6 Inferences from Re-Os systematics of the New Caledonia harzbugites

All type-A harzburgites, including YA1, show unradiogenic ¹⁸⁷Os/¹⁸⁸Os_i and very low 731 Re/Os relative to the range of chondritic values (Walker et al., 2002) or primitive 732 mantle estimates (Meisel et al., 1996). The γ_{Osi} overlap with data from depleted 733 abyssal peridotites and mantle sections from some ophiolites (e.g., Becker and Dale, 734 2016). 735 In order to obtain an estimate of the time of melt depletion, we calculated Re-Os 736 model ages (T_{RD}) and Re-depletion model ages (T_{RD}) . The Re-depletion model ages 737 are generally used for samples with low Re contents and Re/Os ratios. However it is 738 important to note that this method provides minimum depletion ages, as melting is 739 not expected to remove all of the Re on a whole-rock scale. 740 The Re-depletion ages for type-A harzburgites are quite homogeneous for three out 741 four samples (TI1, TI2, YA1) ranging between 0.4 and 0.7 Ga for (see Table 1), while 742 sample PO4 yields an older Re-depletion age of 1.3 Ga. The younger model ages can 743 be linked to the previous evolution of the New Caledonia mantle in relationship to the 744 745 eastern Australian margin, from which the New Caledonia archipelago was separated via marginal rifting about 90 Ma ago (Cluzel et al., 2001, 2012; Whattam, 2009). The 746 747 ancient Re depletion age of the harzburgite PO4, on the other hand, reflects a mantle domain that experienced long-term low Re/Os ratio. This age is also mirrored by Nd 748 isotopic signature, which shows a highly radiogenic value (ε_{Ndi} =+13.32, Secchiari et 749 750 al., 2019a), indicative of a mantle reservoir that underwent long-term depletion of Nd. In addition, similar depletion ages (i.e. 1.2 Ga) occur in the New Zealand mantle rocks 751 (see Scott et al., 2019). 752 These results are consistent with recent Re-Os studies on abyssal peridotites and 753 mantle tectonites from ophiolitic sequences showing that the convecting mantle 754 contains harzburgite domains that underwent depletion events much older than the 755 age of peridotite processing under the ridge (e.g., Harvey et al., 2006). 756

In summary, the New Caledonia harzburgites show Re-Os systematics typical of depleted upper mantle showing Early Paleozoic Os isotopic equilibration and evidence for ancient depletion events (> 1.0 Ga). This multi-stage history led to the depleted nature of the harzburgites and possibly resulted in their complex HSE geochemical signatures.

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Summary and conclusions

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A Re-Os isotopes, highly siderophile and chalcophile element investigation of the New 765 Caledonia peridotites was carried out to unravel the behaviour of the aforementioned 766 767 elements in Iherzolites and harzburgites from the New Caledonia ophiolite. The Iherzolites exhibit subchondritic to slightly suprachondritic ¹⁸⁷Os/¹⁸⁸Os_i (0.1273-768 0.1329). PM-normalised HSE abundance diagrams are characterised by gently sloping 769 patterns showing increasing depletion towards Re-Au, similar to lherzolites that

experienced low to moderate melt extraction. However, the lack of correlation 771 between HSE and fertility indicators, as well as the slightly suprachondritic Os isotopic 772 773 ratios, argue against a simple partial melting history. Rather, the aforementioned

features and the presence of included and interstitial residual monosulphides possibly

indicate that melting occurred on a mantle domain that has previously experienced a

melt percolation event. The high S concentrations of the lherzolites (202-1268 ppm)

most likely result from late-stage seawater-rock reactions.

By contrast, the New Caledonia harzburgites record higher degrees of melt extraction, as attested by the strikingly low, often below the detection limit, concentrations of incompatible chalcophile elements. Despite their homogeneity in terms mineralogical and major element compositions, HSE patterns and Os isotopic compositions indicate the occurrence of two distinct harzburgite sub-groups.

Type-A harzburgites are characterised by steeply plunging HSE patterns, showing IPGE-PPGE and Os-Ir-Ru fractionation, coupled with low Re/Os ratios and subchondritic ¹⁸⁷Os/¹⁸⁸Os_i. The strongly fractionated HSE patterns and the positive Pt anomalies, coupled with the high modelled melting degrees, indicate that melting occurred under hydrous conditions in sub-arc mantle.

Type-B harzburgites display notably different HSE patterns, showing depleted Os-Ir compared to Ru, positive anomalies and Pd-Re re-enrichments (relative to IPGE), coupled with chondritic to strongly suprachondritic measured Os isotopic ratios $(^{187}Os)^{188}Os = 0.127-0.153$). These features have not been yet identified in mantle tectonites and might indicate Os-Ir-Pt release into silicate melt for higher partial melting degrees.

The HSE signature carried by the studied peridotites, as well as the puzzling similarity observed between the New Caledonia harzburgites and the New Zealand mantle xenoliths, might attest the presence of a mantle source bearing a long lasting evolution (> 1 Ga), possibly linked to the Zealandia formation.

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- 1080 Figure captions

- 1081 Fig. 1. a) Present-day structures of the Southwest Pacific region modified after Cluzel
- et al. (2012). Dark orange, land; light orange, continental plateau; white, oceanic
- basins (LHR: Lord Howe Rise, NR: Norfolk ridge, LR: Loyalty ridge, HP: Hikurangi

Plateau); b) simplified geological map of New Caledonia showing distribution of the Peridotite massifs (modified after Cluzel et al., 2012).

1086

Fig. 2. Variations of Os, Ru, Rh, Pt, Pd, Au and Re vs. Ir for the New Caledonia peridotites. Abyssal peridotites (Kane fracture zone: Snow and Schmidt, 1998; Brandon et al., 2000; Luguet et al., 2001, 2003; Marchesi et al., 2013; MAR: Harvey et al., 2006; Lena trough: Lassiter et al., 2014) and ophiolitic peridotites (IL-EL: Internal and External Ligurides, Snow et al., 2000; Luguet et al., 2004; Fischer-Godde et al., 2011; Lanzo: Becker et al., 2006; Pyrenees: Becker et al., 2006; Luguet et al., 2007) are shown for comparison.

1094

Fig. 3. Primitive mantle normalised HSE and chalcogen patterns for the New Caledonia spinel and plagioclase lherzolites. Grey shaded area includes oceanic lherzolites from Mid-Atlantic and South West Indian ridges (Snow and Schmidt, 1998; Luguet et al., 2001; Luguet et al., 2003) and ophiolitic lherzolites from the Ligurian Units (Snow et al., 2000; Luguet et al., 2004; Fischer-Gödde et al. 2011). Normalising values after Becker et al. (2006), Fischer-Gödde et al. (2011) and Wang and Becker (2013).

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Fig. 4. a) Al₂O₃ (wt.%)-¹⁸⁷Os/¹⁸⁸Os_i and b) Os-¹⁸⁷Os/¹⁸⁸Os_i and c) ¹⁸⁷Re/¹⁸⁸Os-187Os/188Os diagrams showing data from New Caledonia Iherzolites and type-A harzburgites in comparison to PM compositions. PM data from Meisel et al. (1996). Data for abyssal peridotites are from Harvey et al., (2006) for Atlantic peridotites, Lassiter et al., (2014) for Lena through and Liu et al. (2015) for Gakkel ridge. See Fig. 2 for ophiolitic peridotites references. Fig. 5. a) Te vs. Se, b) Te vs. Pd and c) S vs. S/Se correlation diagrams for New Caledonia spinel and plagioclase lherzolites. Also showing for comparison data for orogenic lherzolites from Wang and Becker (2013).

Fig. 6. a) Primitive mantle normalised HSE and chalcogen abundances in type-A harzburgites. Light grey shaded field encompasses the area of modern MOR harzburgites (Snow and Schmidt, 1998; Luguet et al., 2001, 2003; Harvey et al., 2006; Marchesi et al., 2013); b) Primitive mantle normalised diagram showing HSE and chalcogen patterns of type-B harzburgites. Normalising values are after Becker et al. (2006), Fischer-Gödde et al. (2011) and Wang and Becker (2013).

Fig. 7. Primitive mantle normalised HSE abundances of a) type-A and b) type-B harzburgites compared to the HSE composition displayed by New Zealand mantle xenoliths. See text for further detail.

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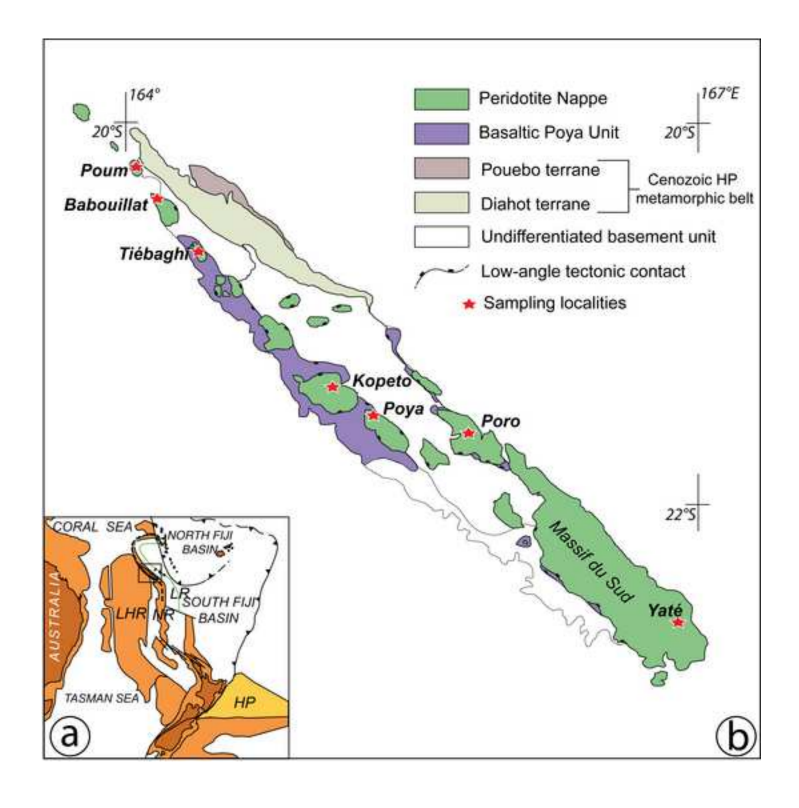


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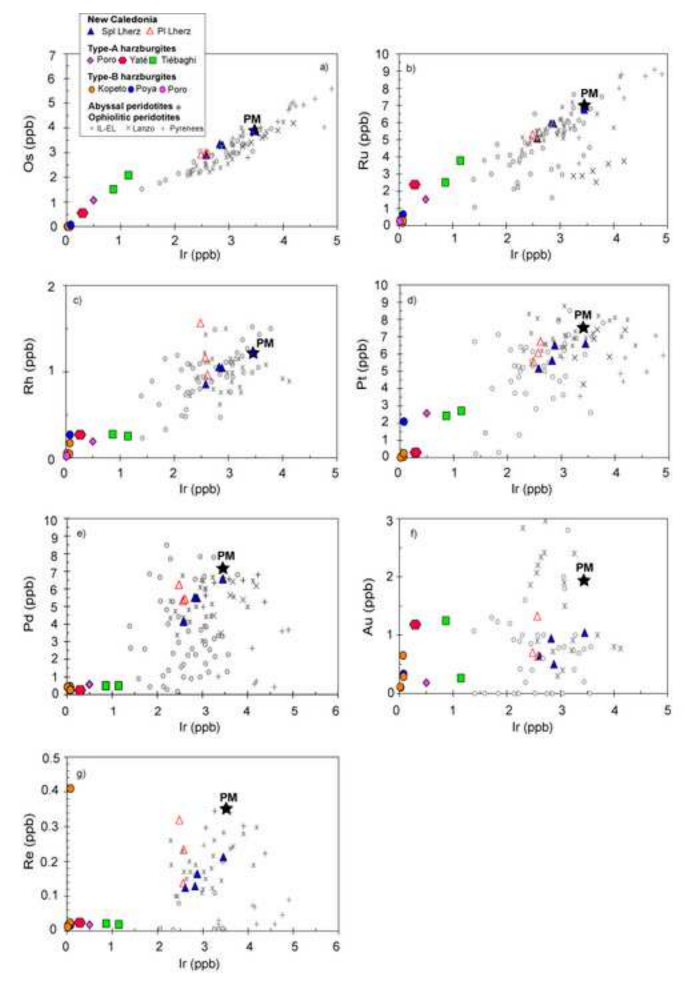


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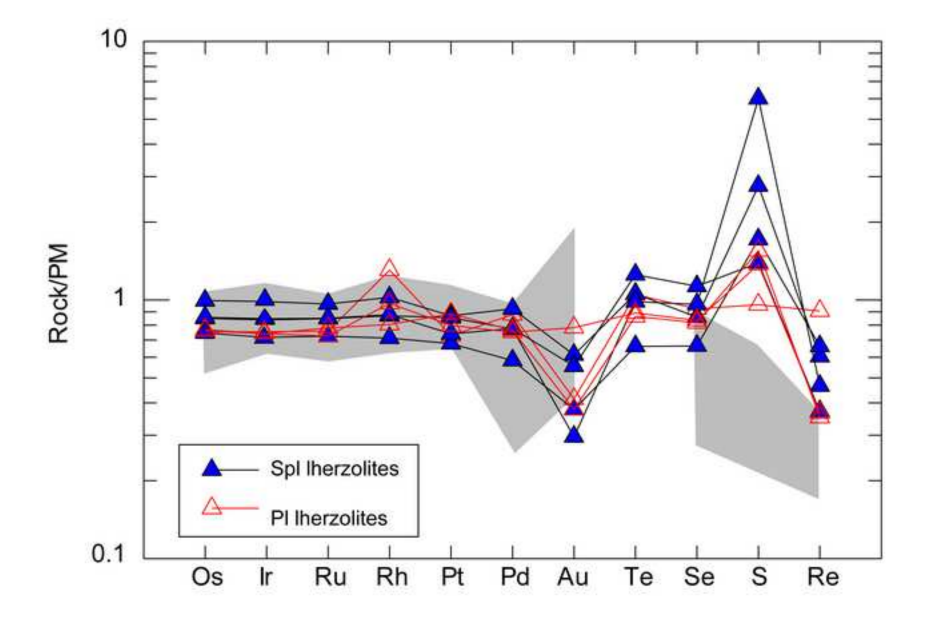


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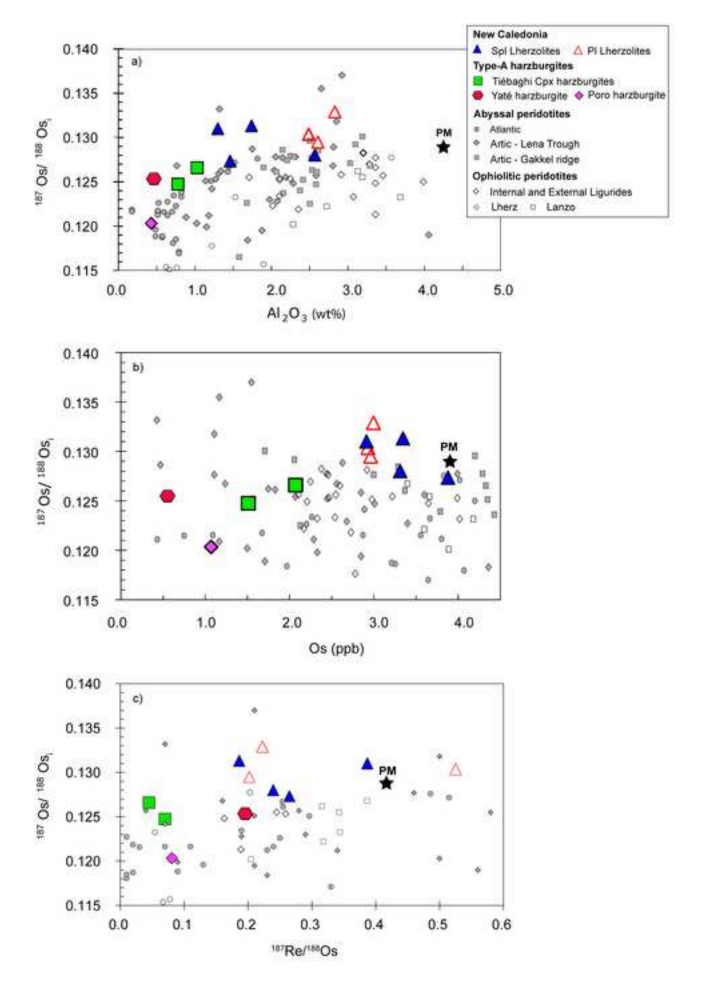


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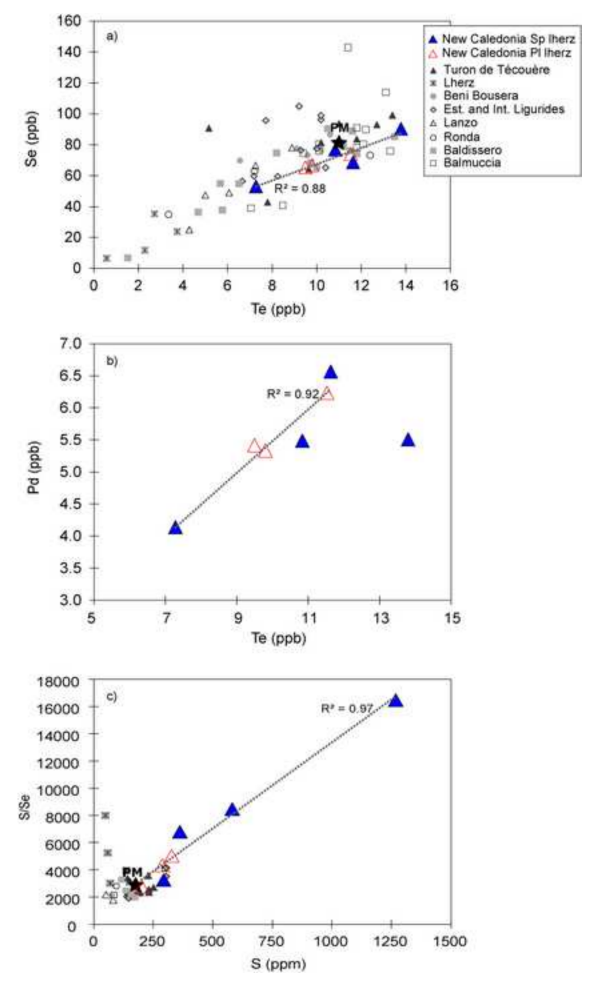


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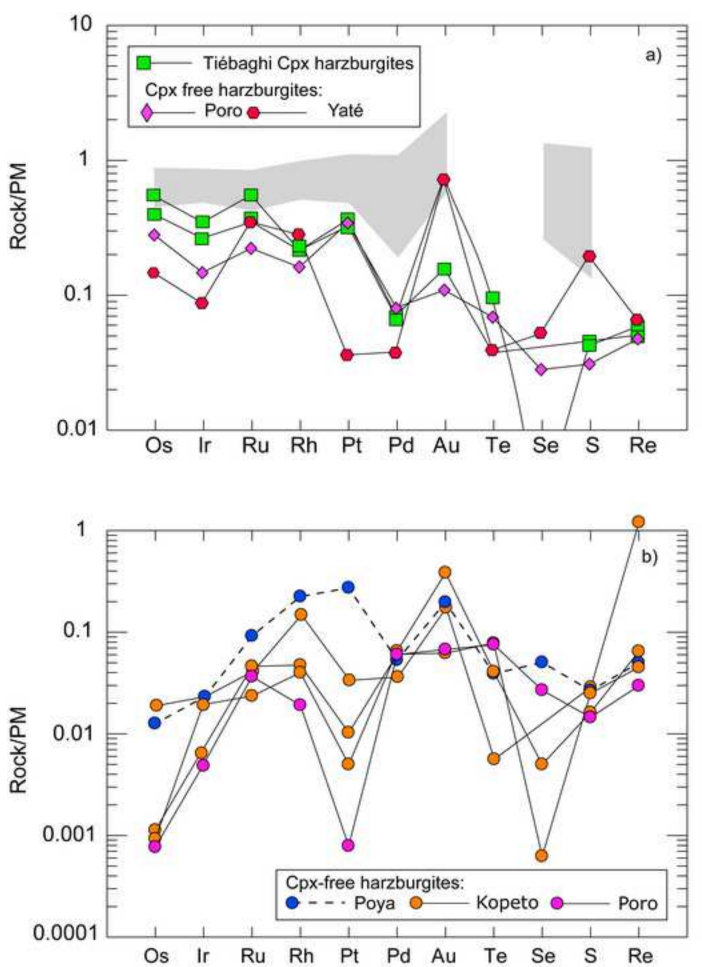
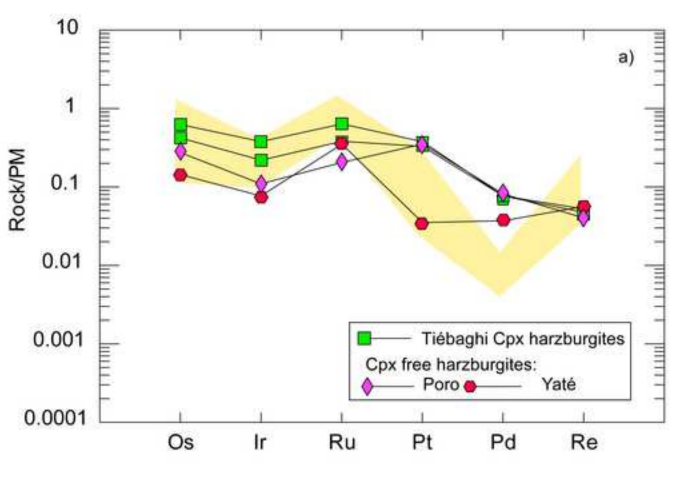
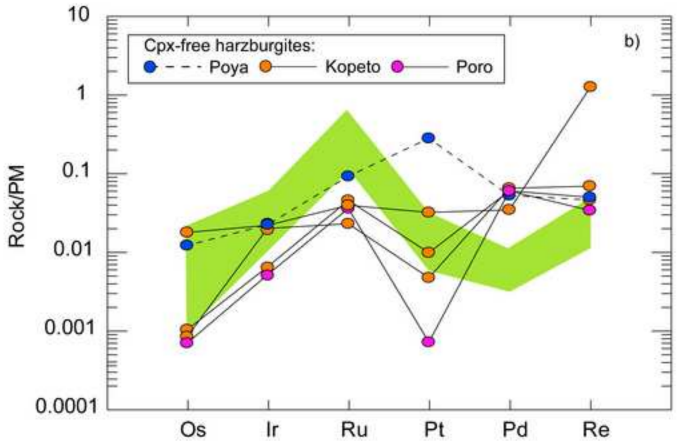
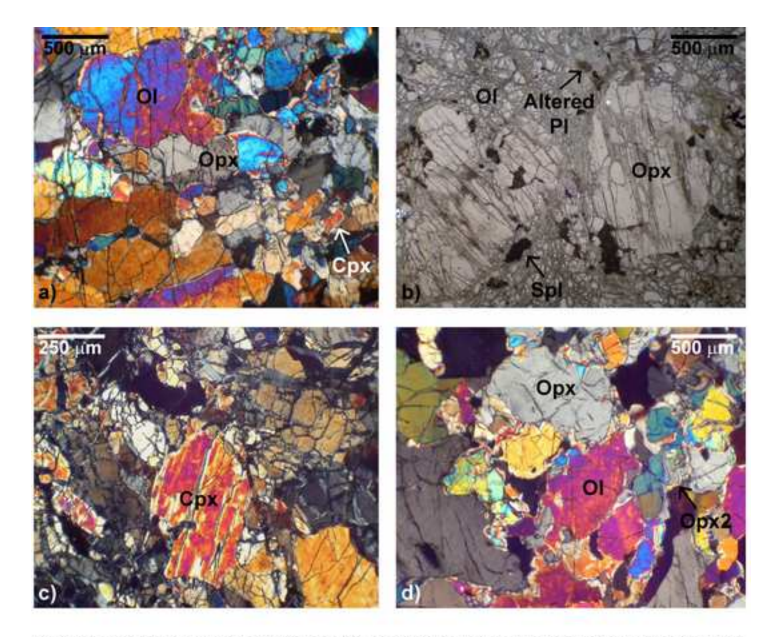


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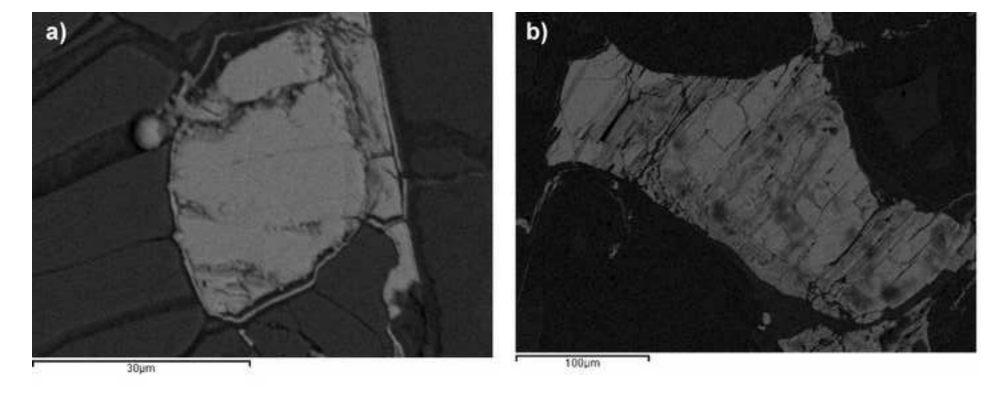






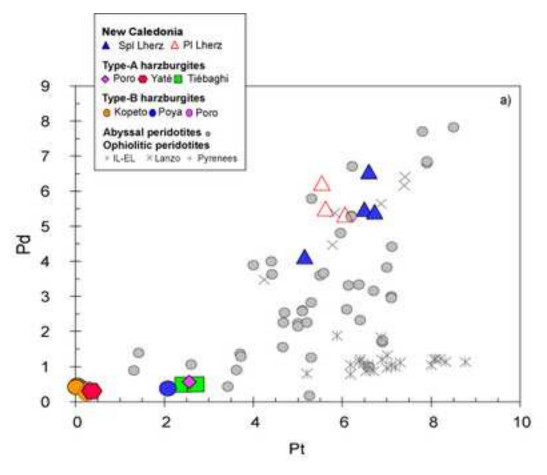
Microphotographs of the New Caledonia peridotites: a) porphyroclastic texture marked by deformed and stretched olivine and orthopyroxene crystals in spl-lherzolite POU2; b) plagioclase-lherzolite thin section (BAB2A, parallel polars); c) strongly exsolved, primary clinopyroxene occurrence in TI2 harzburgite; d) interstitial orthopyroxene (opx2) formed at the expense of primary olivine (KPT5 harzburgite).

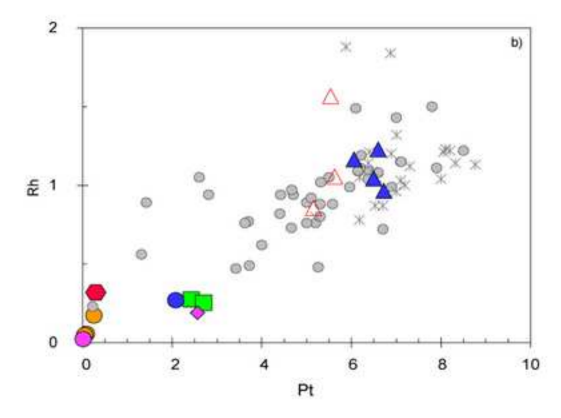
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BSE images of sulphides in the New Caledonia Iherzolites: a) small included sulphide (BA1); b) larger intergranular sulphide in POU2 Iherzolite.

Figure S3
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Pt vs. a) Pd and b) Rh variation diagrams for the New Caledonia peridotites. Abundances are in ppb.

Table 1 Click here to download Table: Table 1.xlsx

 Table 1

 Concentrations of the HSE, S, Se and Te, Os isotopes, selected major elements and ratios for the New Caledonia peridotites

Sample	Rock	Al ₂ O ₃ wt.%	LOI %	Os (ng/g)	Ir (ng/g)	Ru (ng/g)	Rh (ng/g)	Pt (ng/g)	Pd (ng/g)	Au (ng/g)	Re (ng/g)	Os _N /Ir _N	Os _N /Ru _N	Pd _N /Ir _N	Ru _N /Ir _N	Pt _N /Ir _N	Pt _N /Ru _N	¹⁸⁷ Re/ ¹⁸⁸ Os (2SE)	¹⁸⁷ Os/ ¹⁸⁸ Os measured	2SE	¹⁸⁷ Os/ ¹⁸⁸ Os _i	$\gamma Os_{(53Ma)}$	T _{MA} (PM) Ga	T _{RD} (PM) Ga	T _{RD2} (PM) Ga		Se (ng/g)	Te (ng/g)	S/Se	Se/Te
POU1A	PIL	2.49	7.56	2.93	2.48	5.31	1.57	5.54	6.23	0.704	0.319	1.06	0.99	1.24	1.07	1.03	0.96	0.525(1)	0.130822	9.1E-06	0.130358	2.9	0.8	f	f	202	74.9	11.5	2703	6.5
POU2	Sp L	1.46	8.63	3.88	3.45	6.76	1.23	6.60	6.56	1.04	0.213	1.01	0.18	0.94	5.71	0.88	0.15	0.265(1)	0.127574	6.8E-06	0.127340	0.5	0.7	f	0.3	582	69.6	11.6	8364	6.0
POU2B	Sp L	2.57	10.69	3.31	2.88	5.93	1.04	6.49	5.49	0.503	0.165	1.03	1.09	0.94	0.95	1.04	1.09	0.240(1)	0.128227	8.0E-06	0.128016	1.1	0.4	f	0.2	1268	77.8	10.8	16289	7.2
POU3	Sp L	1.30	9.60	2.91	2.58	5.07	0.86	5.15	4.14	0.642	0.234	1.01	1.05	0.79	0.97	0.92	0.95	0.387(1)	0.131356	9.1E-06	0.131015	3.4	f	f	f	362	54.0	7.3	6691	7.4
BA1	Sp L	1.74	6.39	3.34	2.85	5.95	1.05	5.62	5.51	0.943	0.129	1.06	0.27	0.96	3.88	0.92	0.24	0.186(1)	0.131485	7.9E-06	0.131320	3.7	f	f	f	294	91.3	13.8	3218	6.6
BAB1B	PI L	2.61	6.98	2.96	2.61	5.46	0.965	6.73	5.42	0.641	0.125	1.02	0.24	1.02	4.30	1.19	0.28	0.202(1)	0.129664	8.9E-06	0.129485	2.2	f	f	0.0	327	66.2	9.5	4943	7.0
BAB2B	PIL	2.83	8.43	3.00	2.57	5.08	1.16	6.06	5.33	1.33	0.138	1.05	1.67	1.02	0.63	1.09	1.73	0.223(1)	0.133084	8.9E-06	0.132887	4.9	f	f	f	289	67.6	9.8	4278	6.9
TI1	Н	0.78	9.01	1.51	0.864	2.52	0.275	2.42	0.488	1.25	0.022	1.57	1.08	0.28	1.46	1.29	0.89	0.070(25)	0.12479	1.7E-05	0.12473	-1.5	0.8	0.7	0.7	9	1.1	1.0	8188	1.0
TI2	Н	1.03	6.04	2.07	1.14	3.78	0.254	2.70	0.499	0.264	0.019	1.63	0.98	0.21	1.65	1.09	0.66	0.045(18)	0.12662	1.3E-05	0.12658	-0.1	0.5	0.4	0.4	10	0.7	0.4	14053	1.7
PO4	Н	0.43	0.18	1.07	0.499	1.53	0.191	2.56	0.570	0.160	0.018	1.93	1.26	0.56	1.53	2.36	1.54	0.081(35)	0.12040	2.4E-05	0.12033	-5.0	1.5	1.3	1.3	6	3.1	0.8	2084	4.1
YA1	Н	0.46	6.83	0.554	0.297	2.35	0.326	0.268	0.263	1.17	0.022	1.67	0.42	0.44	3.95	0.42	0.11	0.196(68)	0.12551	4.7E-05	0.12534	-1.0	1.0	0.6	0.6	40	4.9	0.4	8117	11.4
PY1	н	0.78	0.00	0.049	0.077	0.643	0.269	2.08	0.378	0.336	0.016	0.58	0.14	2.43	4.19	2.97	2.97	1.62(8)	0.1299	5.3E-04	0.1284	1.4	0.0	0.0	0.2	6	0.9	0.9	6103	1.1
Duplicate				0.023	0.133	0.693	0.171	2.44	0.588	0.311	0.062							13.0(1)	0.127	1.0E-03	0.115	-8.9				53	0.3	1.0	196197	0.3
KPT2	н	0.70	3.03	0.004	0.022	0.323	0.057	0.078	0.431	0.107	0.018	0.18	0.02	9.50	7.22	0.22	0.22	19(1)	0.148	5.6E-03	0.131	3.6	0.1	f	f	5	1.3	0.4	4164	2.8
Duplicate				0.036	0.017	0.211	0.040	0.210	0.620	0.144	0.010							1.4(1)	0.302	1.3E-03	0.301	137.8				7	1.3	8.2	4947	0.2
KPT5	н	0.74	0.12	0.004	0.066	0.165	0.048	0.038	0.464	0.654	0.024	0.05	0.04	3.46	1.25	0.21	0.21	32(2)	0.147	6.2E-03	0.118	-6.6	0.0	f	f	3	3.0	0.7	1133	4.1
Duplicate				0.007	0.062	0.165	0.054	0.046	0.512	0.603	0.056							37(1)	0.160	3.2E-03	0.127	0.5				3	0.7	1.3	4453	0.5
PO3	Н	0.41	0.00	0.003	0.015	0.256	0.023	0.006	0.428	0.084	0.012	0.18	0.02	14.12	8.57	0.02	0.02	19(2)	0.153	8.4E-03	0.136	8.0	0.1	f	f	3	1.4	0.6	2129	2.2
Duplicate				0.055	0.018	0.243	0.016	0.009	0.573	0.293	0.0078							0.69(7)	0.1239	4.7E-04	0.1233					bdl	bdl	0.8	-	-
KPT3	Н	0.67	0.67	0.072	0.079	0.280	0.172	0.248	0.250	0.288	0.410	0.81	1.87	0.17	0.19	6.92	6.92	28(1)	0.1273	3.5E-04	0.1204	-5.1	0.0	0.3	1.3	6	bdl	bdl	-	-

Duplicate: replicate digestion of the same sample powder

¹l L= plagioclase lherzolite, Sp L= spinel lherzolite, H= harzburgite

Values of PM 187Os/188Os = 0.1296 and 187Re/188Os = 0.434 used for calculation of T_{MA} and T_{RD} ages (Meisel et al., 2001); f= future model ages. T_{RD2} (PM) indicates depletion ages calculated taking into account Re addition that may have occurred during peridotite evolution.

Table 2Equilibration Temperature, Pressure, and Oxygen Fugacity Calculated for selected peridotite samples

Sample	Туре	T BK (°C)	P (GPa)	Wood (199	90)	T OI-Spl (°C)	Wood (1990))
				log f(O2)	ΔFMQ		log f(O2)	ΔFMQ
TI2	Type-A harz	930	1.5	-10.40	0.71	870	-11.37	0.76
YA1	Type-A harz	980	1.5	-9.83	0.50	815	-12.66	0.50
KPT5	Type-B harz	1130	1.5	-7.91	0.39	840	-12.20	0.48
PO3A	Type-B harz	1000	1.5	-9.86	0.17	940	-10.76	0.18
PY1B	Type-B harz	1050	1.5	-9.17	0.16	930	-10.90	0.20
BA1	Spl lherz	1060	1.5	-13.1503	-3.96	880	-15.66	-3.73

T BK= Equilibration temperature calculated on Opx porphyroclasts using Brey and Köhler (1990)

T OI-SpI = Equilibration temperature calculated using Li et al. (1995) formulation Oxygen fugacity estimates are from Wood et al. (1990) and are reported as Δ log fO2 from the quartz-fayalite-magnetite (FMQ) buffer using the Fe³⁺/ Σ Fe values of the analysed spinels. See text for further details.

Table S1 Click here to download Table: Table S1.xlsx

Table S1Major element composition for selected sulphides from the New Caledonia Iherzolites

Sample Mineral Occurrence	POU2 mss Intergr	POU2 mss Intergr	POU2 mss Incl	POU2 pn Incl	POU2 pn Incl	POU2 pn Incl	POU2 pn Incl	BA1 mss Incl	BA1 mss Incl	BA1 mss Incl	BA1 mss Incl	BA1 mss Incl
wt%	р	р	р	р	р	р	р	р	р	р	р	р
S	34.32	34.39	32.74	33.24	41.19	32.68	40.82	33.22	33.08	32.47	32.53	33.13
Fe	41.23	43.80	36.65	28.09	23.90	29.14	24.68	36.85	42.99	43.25	42.19	43.02
Co	0.64	0.59	0.78	0.46	1.93	1.91	2.26	0.46	0.50	0.55	0.80	0.58
Ni	23.28	20.44	29.19	37.55	32.99	35.74	32.23	25.77	23.44	24.72	24.54	23.27
Cu	0.52	0.57	0.64	0.66	bdl	0.53	bdl	3.70	bdl	bdl	bdl	bdl
Total	99.99	99.79	100.00	100.00	100.01	100.00	99.99	100.00	100.01	100.99	100.06	100.00
Fe/Ni	1.8	2.1	1.3	0.7	0.7	8.0	8.0	1.4	1.8	1.7	1.7	1.8

Pn= pentlandite; mss= monosulphide solid solution