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3	X-ray absorption records of Pd ²⁺ on Ni site in pentlandite
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17	Abstract
18	Norilsk sulfide ores are one of the largest known sources of Pd on Earth. Palladium in
19	these ores is presented in platinum-group minerals (PGM) and base metal sulfides (BMS),
20	especially in pentlandite ((Fe,Ni) ₉ S ₈). Although several studies demonstrated high concentrations

along with heterogeneous distribution of Pd in pentlandites from Norilsk, the form of Pd in 21 pentlandite has not been established. Here, we provide the first evidence for Pd incorporation in 22 the lattice of pentlandite from Norilsk ores. These results were obtained by combining X-ray 23 absorption near edge structure (XANES) spectroscopy, synchrotron-based micro-X-ray 24 fluorescence (μXRF), and electron backscatter diffraction (EBSD). We present the first ever 25 26 measured XANES spectra of Pd in pentlandite and atokite ((Pd,Pt)₃Sn) as well as in other common Pd minerals. Divalent Pd in pentlandite was detected by XANES. The Pd spectra in 27 pentlandite show no similarities with Pd spectra in PGM, metallic Pd, PdS, PdCl₂, and PdSO₄ 28 which signifies that Pd incorporates into the lattice of pentlandite. Substitution of Ni by Pd in the 29 lattice of pentlandite is supported by negative correlations shown by uXRF and electron probe 30 microanalysis (EPMA) and complies with the previous studies. The additional EBSD study 31 demonstrates a resemblance in cell parameters of the Pd-rich and Pd-poor parts of the pentlandite 32 grains and reflects that Pd incorporation into the pentlandite structure does not imply any notable 33 structure distortion. The combination of analytical techniques used in the present study 34 demonstrates the great potential of these methods for understanding the mechanisms of noble 35 metal incorporation into ore minerals. 36

Keywords: Palladium, pentlandite, platinum-group minerals, XANES, μXRF, EBSD,
EPMA

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Introduction

Palladium, like other platinum-group elements (PGE), occurs as discrete platinum group
minerals (PGM) or in base metal sulfides (BMS), associated with mafic intrusions (Godel et al.,
2007; Barnes et al., 2008) and komatiite flows (Keays et al., 1981; Lesher and Keays, 2002).
Pentlandite (Pn) among other base metal sulfides appears to be a principal host in most deposits.

The highest concentration of Pd in Pn (up to 11.26 wt%; Kalugin et al., 2021) was reported from 44 the Southern 2 orebody of the Talnakh deposit (Russia). The origin of Pd in Pn is debated. 45 Kalugin et al. (2021) attributed the high Pd concentrations in Talnakh pentlandite to the influence 46 of Pd-rich fluids. The Pd-rich pentlandite from the J-M reef ores, Stillwater deposit (Montana) 47 contains up to 9.8 wt% of Pd, which is believed to be a secondary enrichment during the 48 49 hydrothermal redistribution of Pd from the surrounding PGM (Li and Ripley, 2006). Until recently, most works assumed based on experimental work (Kelly and Vaughan, 1983; 50 Etschmann et al., 2004; Helmy et al., 2021), that pentlandite formed by exsolutions from 51 monosulfide solid solution (MSS) in the non-altered magmatic sulfide ores. However, the 52 partition coefficient of Pd into MSS is <0.2, thus the presence of bulk Pd in pentlandite requires 53 an explanation (Barnes et al., 1997; Liu and Brenan, 2015). Makovicky et al., 1986 showed that 54 Pd solubility in MSS increases with temperature. Upon cooling, MSS rejects Pd and transforms 55 into pyrrhotite, and pentlandite, the latter can incorporate Pd. Some authors suggested that PGE 56 are present in silicate liquid as clusters (Tredoux et al., 1995; Helmy et al., 2020), and these 57 clusters were captured by the sulfide liquid and incorporated into the MSS. Pre-nucleation 58 clusters of Ir, Os, Ru, and Rh are present as metal-metal, metal-S, and metal-AsS complexes, but 59 60 Pt and Pd form semimetal (As, Te, Bi, Sb) complexes (Helmy et al., 2013; Laurenz et al., 2013; Helmy and Bragagni, 2017; Liang et al., 2022). The experimental work of Kitakaze et al. (2016) 61 offers an alternative namely that of pentlandite having formed by peritectic reaction at high 62 temperature between monosulfide solid solution (MSS) and a residual sulfide liquid. An example 63 of this is the granular Pn that occurs at the contact between pyrrhotite (Po) and chalcopyrite 64 (Cpy) and tends to be enriched in Pd at the contact with Cpy compared to the Pn in contact with 65 Po (Dare et al., 2010; Mansur et al., 2019; Brovchenko et al., 2020). In this case, it is argued that 66 Pd preferentially partitions into Pn and thus should be a part of the lattice structure. According to 67

Makovicky et al. (2016) pentlandite dissolves up to 5.4 at.% of Pd and 92 % of Pd occupy the 68 octahedral site in the lattice. Based on correlations between Pd content and Ni/Fe ratio in zonal 69 Pd-rich Pn, Kalugin et al. (2021) concluded the isomorphic substitution of Ni by 0.71 apfu Pd and 70 0.3 apfu Fe. Wirth et al. (2013) and Junge et al. (2015) used focused ion beam (FIB) combined 71 with transmission electron microscopy (TEM) to study the site of Pd in Pn from the Merensky 72 73 and UG-2 reefs of the Bushveld Complex (South Africa). Wirth et al. (2013) found Pd to be present as nanonuggets (of Pd-Sn and Pt-Pd-Sn compounds) in Pn and Junge et al. (2015) found 74 Pd to be present both as part of the Pn lattice and as nanonuggets. The question arises as to 75 whether the nanonuggets represent clusters and are an important part of the budget of Pd. Some 76 authors argue that nanonuggets are essential to the formation of PGE deposits (Tredoux et al., 77 1995; Helmy et al., 2013; Anenburg and Mavrogenes, 2020; Kamenetsky and Zelenski, 2020). 78 On the other hand, those working with laser-ablation inductively coupled plasma mass 79 spectrometry (LA-ICP-MS) argue that for time-resolved analysis signals (TRA) each time slice 80 represents ~ 0.2 to 0.6 microns of material ablated and a constant signal shows that the element 81 uniformly present in the mineral on this scale (Page and Barnes, 2016) implying that the element 82 (in this case Pd) is in solid solution in the mineral (Pn). 83

The Norilsk-Talnakh deposits of Siberia (Russia) are one of the largest known sources of Pd in the world (Naldrett, 2011; Barnes et al., 2020). The distribution of Pd in pentlandite from various Norilsk-Talnakh ore deposits has been studied by a number of analytical methods, including electron probe microanalysis (EPMA), LA-ICP-MS, and particle-induced X-ray emission (PIXE) (Cabri et al., 1984; Czamanske et al., 1992; Distler et al., 1996; Cabri et al., 2003; Barnes et al., 2006; Mansur et al., 2019; Brovchenko et al., 2020; Kalugin et al., 2021). These studies provided valuable information on the concentration and distribution of Pd in

pentlandite. Brovchenko et al. (2020) showed that primary magmatic pentlandite in massive ores 91 92 of Mt. Rudnaya offshoots, Norilsk-Talnakh deposits, contains a high concentration of Pd (up to 93 4.62 wt%). A comprehensive geological review of the Mt. Rudnaya offshoots is presented in Brovchenko et al. (2020). The deepest terminations of the Mt. Rudnaya ores had a specific 94 globular texture composed of Cu-rich intermediate solid solution (ISS) globules up to 4 mm in 95 96 size surrounded by MSS-ISS fine-grained matrix with MSS outward bands between matrix and 97 globules. Both MSS and ISS are relatively high-temperature sulfides that are transformed to BMS 98 at subsolidus temperatures (Craig and Kullerud, 1969; Cabri, 1973). Therefore, the presence of 99 MSS and ISS in natural sulfide ores indicates that these ores formed due to the fast quenching of a primary magmatic sulfide melt (Helmy et al., 2021). The most enriched in Pd and the biggest 100 pentlandite grains (with a size up to $100 \ \mu m$) are predominately confined to the contact between 101 ISS globules and MSS outward bands. Moreover, Pd in these pentlandite grains is 102 103 heterogeneously distributed with an increase of concentration from the contact with MSS toward the contact with ISS. Such type of enrichment of contact pentlandite was considered to display its 104 peritectic reactional origin (Brovchenko et al., 2020). The current work focuses on Pd 105 106 incorporation in Mt. Rudnaya pentlandite.

107 X-ray absorption spectroscopy (XAS), which includes X-ray absorption near-edge 108 structure (XANES) and extended X-ray absorption fine structure (EXAFS), can be used to 109 determine the local structure of elements in minerals. These techniques have been successfully 110 applied to reveal chemical state and atomic position of Au in Fe-sulfides (Tagirov et al., 2016; 111 Trigub et al., 2017; Merkulova et al., 2019; Pokrovski et al., 2019), Hg in pyrite and marcasite 112 (Manceau et al. 2018), Pt in synthetic pyrite (Filimonova et al., 2019), As and Se in pyrite 113 (Manceau et al., 2020) and Pt in pyrrhotite (Filimonova et al., 2021). To the best of our

knowledge, no attempts have been made to use XAS to detect the speciation and occurrence of Pd in pentlandite. The main limitation could be the low concentration of Pd in natural pentlandite, difficulties in the experimental synthesis of homogenous Pd-doped pentlandite, and lack of familiarity of geoscientists with this technique combined with its difficult accessibility. Thus, besides evidence of the form of Pd in pentlandite from the Bushveld complex, the site of Pd in pentlandite from other major PGE ore deposits is still unconfirmed.

In the present study, we report the oxidation state and structural form of Pd in pentlandite 120 121 from the Norilsk ore deposit by using XANES in combination with synchrotron-based micro-Xray fluorescence (µXRF) and electron backscatter diffraction (EBSD). Additionally, we 122 determine the oxidation state of Pd in metallic Pd, PdS, PdCl₂, PdSO₄, and in six different PGM 123 124 to compare XANES spectra of Pd in pentlandite and other common Pd-bearing compounds. This work is the first to show Pd L₃-edge XANES measurements in pentlandite and PGM. The EBSD 125 126 showed the granular structure of the sample and did not demonstrate any deformations inside the 127 crystallites of Pd-rich pentlandite. The outcome of this study provides new insights into the substitutional mechanism of Pd in pentlandite and encourages further use of XAS methods in 128 129 mineralogy and ore geology.

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Materials and Methods

Sulfide mineral samples were collected from the deepest terminations of Mt. Rudnaya subvertical massive sulfide offshoots in the NE of the Norilsk 1 deposit (Brovchenko et al., 2020). Twenty polished blocks were examined using a petrographic light microscope, a scanning electron microscope (SEM), XRF, EPMA, LA-ICP-MS, and X-ray computed micro-tomography (micro-CT) prior to this study. The descriptions of these samples, analytical methods and the results of the analyses are described in detail in Brovchenko et al. (2020) and Sittner et al. (2022).

Apart from BMS, Mt. Rudnaya MSS-ISS ores also contain PGM up to 125 µm in size such as 137 tetraferroplatinum (PtFe), Pt-Fe alloys having the compositions Pt₂Fe and Pt₃Fe₂, cooperite (PtS), 138 Pd- rich rustenburgite (Pt₃Sn), Pt-rich atokite (Pd₃Sn), and Au-Ag alloys, mostly occurring in the 139 ISS globules (Sittner et al., 2022). Three sample blocks, namely RM 10, RM 29, and RM 30, 140 containing pentlandite with the highest concentration of Pd (up to 4.84 wt%), were selected for 141 142 the current study. Pentlandite grains in these selected samples occur at the contact with Cu-rich ISS globules and Cu-poor MSS outward bands, surrounded by MSS-ISS fine-grained matrix as 143 described in Brovchenko et al. (2020). Additionally, six different Pd-bearing PGM were selected 144 from the Mineralogical Collections of the Technische Universität Bergakademie Freiberg 145 (Germany) for the XANES and µXRF measurements. The origin and composition of the 146 investigated minerals are listed in Table 1. 147

The sulfide mineral samples from Norilsk and PGM grains were mounted in epoxy and 148 polished. The epoxy mounts were polished with 3-to-1 µm diamond suspension before every 149 150 analytical procedure, including synchrotron measurements. This was done to remove the oxidized layer that can form on the surface of sulfide minerals and PGM. Synchrotron-based XANES and 151 µXRF measurements were acquired on beamline PHOENIX I at the Swiss Light Source (SLS, 152 153 Switzerland). Measurements were performed in fluorescence-yield mode and with a focused 154 beam for Norilsk samples and PGM grains. Concentrated powders of Pd compounds (PdS, PdCl₂, 155 PdSO₄), used as references for XANES, were measured with an unfocused beam in electron-yield mode. The electron-yield mode was used to exclude self-absorption effects that occur in 156 fluorescence XAS for concentrated samples. 157

The incoming X-ray beam was monochromatized by the 111 reflection of a Si double crystal monochromator. The flux on the sample was approximately 5×10^{10} photons s⁻¹. The size

160 of the unfocused beam was $1 \times 1 \text{ mm}^2$. The beam focused with a set of Kirkpatrick-Baez (K-B) 161 mirrors had a size of $4 \times 4 \mu \text{m}^2$. The fluorescence intensity was measured with a Si drift detector 162 (4 element Vortex, Hitachi, USA). The samples were positioned at 45° with the respect to the 163 incident X-ray beam and the detector. All measurements took place under vacuum at 1×10^{-6} 164 mbar.

Elemental μ XRF maps were obtained by scanning the sample with an incident focused beam with an energy of 3174 eV, and a dwell time of 1 s. Single-point μ XRF spectra were collected with a beam energy of 3174 eV and a collection time of 300s. All XRF data were analyzed using PyMCA software (Sole et al., 2007).

The spots with the high Pd signal on the μ XRF maps also called "hot spots" were selected 169 170 for the XANES analysis. The incident energy for the Pd L₃-edge XANES spectra was scanned from 3100 to 3327 eV for all samples and references. The XANES spectra could not be evaluated 171 over a larger energy range due to the presence of the Pd L_2 -edge at 3330 eV. The spectra were 172 recorded with a step of 5 eV for the pre-edge region (3100-3165 eV), 0.3 eV for the edge and 173 XANES region (3166-3190 eV), and with a step increasing from 1 to 2.4 eV for the post-edge 174 region (3191-3327 eV). One XANES spectrum per measurement was taken for Pd references, 1-3 175 spectra per sample were taken for PGM grains, and from 1 to 41 spectra per point measurement 176 were taken for pentlandite. No beam damage was observed during repeated scans on the same 177 178 point of a sample. The XANES spectrum of a Pd foil was measured 1-2 times every 24 hours during the beamtime in order to follow an energy shift of the incident beam. No energy shift was 179 observed. The absolute energy of the spectra is referenced to the maximum of the Pd⁰ (Pd foil) 180 derivative taken to 3173.8 eV. The precision of the energy measurement is \pm 0.1 eV. The 181 XANES spectra of all PGM grains including a PGM grain from Norilsk samples were corrected 182

for the self-absorption effect. The relative energies between spectra were established by a careful comparison of the first derivatives. The XANES spectra were reduced, normalized and corrected for self-absorption with the Athena software (Ravel and Newville, 2005).

Concentrations of the major and trace elements in pentlandite were determined by an 186 JEOL JXA 8200 EPMA at the "IGEM-Analitika" analytical centre. The EPMA is equipped with 187 five wave-length spectrometers. Measurements were operated at 20 kV accelerating voltage, 20 188 nA beam current per Faraday cylinder, and 1 μ m beam diameter. The counting time for Fe (Ka, 189 190 LIF), S (K α , PET), Ni (K α , LIF), Cu (K α , LIF), and Co (K α , LIF) was 10 s for the peak and 5 s 191 for the backgrounds. Natural CuFeS₂ was used as a reference for Cu, Fe, and S determination; 192 and NiSbS was utilized for analyzing Ni and Co. Pure synthetic Pd was used as reference 193 materials for Pd (L α , PET) determination with 20 s counting time for the peak and 10 s for the backgrounds. Pure native Pt was used as a reference material for Pt (L α , LIF) determination at 60 194 195 s counting time for the peak and 30 s for the backgrounds. Native Ag was used as a reference 196 material for Ag (L β , PET) determination at 30 s counting time for the peak and 15 s for the backgrounds. Synthetic GaAs was used for As (La, TAP) determination with a counting time of 197 198 30 s for the peak and 15 s for the backgrounds. The analytical conditions provided detection limits (3 standard errors) of around (wt %) 0.08 for Cu, 0.07 for Pt, 0.06 for Co, 0.05 for As, 0.04 199 200 for Ag, and 0.02 for Pd. The uncertainty of the concentrations was estimated from the 201 reproducibility of home standard analysis as <5 % rel. (2 standard errors).

Homogeneity and distribution of crystallographic orientations within the abovementioned Pd-rich pentlandite grains were studied with EBSD using Hitachi S-3400 SEM equipped with Oxford NordLysNano EBSD detector and Oxford X-max 20 energy-dispersive X-Ray analyzer (EDX) ("Geomodel" resource center, St Petersburg University (SPbU)).

Acquisition conditions were as follows: 30 kV accelerating voltage, 5 nA beam current, step size 0.5-1 µm, averaging of 3 patterns per point. All the patterns were processed automatically using Oxford AztecHKL software, the maps were created with the Channel5 software. The reference structure for pentlandite was taken from Inorganic Crystal Structure Database (ICSD, number 61021). Prior to EBSD studies, Ar ion polishing was performed (Oxford IonFab 300, 10 min exposure, 500 V accelerating voltage, 200 mA beam current, at "Nanophotonics" resource center, SPbU) to remove the mechanically deformed layer from the surface.

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Results and Discussion

214 Distribution of Pd and element correlations

Elemental mapping by synchrotron-based µXRF was employed to investigate the 215 distribution of Pd and its correlations with other elements, notably Ni, Cu, S, and Pt. The µXRF 216 mapping shows a heterogeneous distribution of Pd in sulfides. Figure 1a provides the elemental 217 distribution maps of Pd, Ni, Cu and S analyzed on sample RM 30 3. The association of Pd with 218 Ni shown on the elemental map documents that Pd is localized only in pentlandite and not in the 219 Cu-rich intermediate sulfide solid solution (ISS) or Fe-rich MSS. The Pd hot spot in the sample 220 has a size of $45 \times 30 \text{ }\mu\text{m}^2$ and is the biggest among detected Pd-rich areas in all samples. Two Pd-221 rich spots ($2 \times 3 \mu m^2$) to the left and bottom of the main Pd-rich region in pentlandite might 222 represent Pd-bearing PGM. However, due to the small size, it was not possible to measure a 223 XANES or µXRF spectrum at these locations. Similar elemental correlations and the same 224 elemental distribution are observed on elemental maps of samples RM 29 1 and RM 29 3 225 226 (Supplementary Figures S2, S3, and S4). No Pt fluorescence was observed in the scanned areas. Four Pd-rich domains in the three sulfide mineral samples are related to pentlandite. In contrast, 227 one Pd-rich region was found in sample RM 10 3, which has other elemental correlations (Figure 228

1b). The Pd in sample RM 10 3 correlates positively with Pt. No Ni or Cu are detected in the Pd-

rich zone. The detected Pd hot spot represents a Pd-bearing PGM grain with a size of $10 \times 6 \,\mu\text{m}^2$.

Micro-XRF spectra recorded in the hot spots of Pd are presented in Figure 2. Four μ XRF 231 spectra taken in Pd hot spots of samples RM 29 and RM 30 are very similar to each other and 232 indicate the occurrence of Pd in pentlandite, which in turn was determined based on the 233 occurrence of the Ni peak at 0.848 keV (Figure 2a). Figure 2b shows µXRF spectra of pentlandite 234 from sample area RM 30 3, a PGM grain from sample RM 10 3, and pentlandite and ISS matrix 235 236 in sample RM 29 1. The four spectra are evidently different. The μ XRF spectra in the matrix of pentlandite and ISS were recorded to demonstrate that no intense Pd La peak (2.838 keV) is 237 observed in the matrix material. A low intensity noisy peak at the energies of Pd La is observed 238 239 in the spectra and attributed to the entrance of Pd fluorescence from the Pd hot spot in the proximity of the matrix area. Therefore, the Pd fluorescence in all μXRF spectra is attributed to 240 the Pd-rich pentlandite and PGM. Platinum, identified by the high-intensity Ma peak at 2.047 241 242 keV is clearly visible in the spectrum of PGM grain. Whereas no Pt peak is observed in the spectrum of pentlandite and matrix regions. Other peaks that can be identified in the µXRF 243 spectra are Fe (L α = 0.704 keV), S (K α = 2.309 keV), Cu (L α = 0.928 keV), Si (K α = 1.740 244 245 keV), Al (K α = 2.309 keV). Iron and sulfur are major elements in both, pentlandite and ISS 246 matrix, and they are often detected in different Pd-bearing PGM (e.g., Sluzhenikin et al., 2020). 247 Copper is a major element in ISS. The appearance of fluorescence peaks of Si and Al in the spectrum of the sample RM 30 3 is explained by the presence of small holes common for the 248 samples and containing Si- and Al-bearing debris from polishing. In the case of sample RM 30 3 249 250 the holes are located near the Pd hotspot and are therefore recorded in the µXRF measurements.

The above observations based on μXRF measurements confirm the occurrence of Pd in at 251 252 least two forms in the studied samples from Norilsk. Palladium within pentlandite grains and Pd 253 within the PGM grain. The relationship between Pd and Ni in pentlandite determined by μXRF was additionally examined by EPMA. The Pd concentrations of up to 4.84 wt.% were detected. 254 Ouantitative EPMA measurements (Table S1) indicated a negative correlation between the two 255 256 elements with a Pearson correlation coefficient of -0.97. In addition, EPMA analyses demonstrated a positive correlation between Pd and Fe with a Pearson correlation coefficient of 257 0.65 (Figure 3). 258

EBSD study of the Pd-rich pentlandite grains

Diffraction contrast maps show that the pentlandite crystallites exhibit the same quality 260 261 over the entire grain surface and thus the existence of nanocrystalline Pd inclusions is improbable (Figure 4). Grain boundaries divaricate to the Pd-rich and Pd-poor zones contact (Figures 4e, 4f, 262 and 4g) and do not correlate with Pd content (Figure 4c), nor any defect accumulation is observed 263 between Pd-rich and Pd-poor zones. The orientation maps correspond to a polycrystalline growth 264 from a common border. The patterns acquired in the areas with different Pd content within the 265 266 same crystallite do not differ in either the bands positions or bandwidth. The resemblance between the patterns suggests that the presence of Pd in the structure does not lead to a noticeable 267 distortion of the structure of pentlandite (Figures 4a and 4b), which means Ni is substituted by Pd 268 269 in the structure. The additional EBSD data are presented in the Supplementary material (Figures S6 and S7). 270

271 Oxidation state of Pd in PGM

The Pd L₃-edge XANES spectra of the six PGM and the Pd metal reference are shown in 272 Figures 5a and 5b. The six spectra of the PGM grains differ from each other, which is consistent 273 with the different atomic coordination of Pd in these six minerals. The PGM spectra are shifted to 274 higher energy relative to Pd foil (metallic Pd⁰). The energy positions of the white lines, feature 275 marked as a', of the PGM and the reference spectra are listed in Table 2. The energy positions of 276 277 the white lines for the PGM spectra vary from 3174.7 eV for vasilite to 3175.6 eV for paolovite. The difference in white line energy for the vasilite and paolovite spectra compared to the white 278 line of metallic Pd is 0.9 and 1.8 eV respectively. The PGM spectra are shifted towards higher 279 energies compared to the spectrum of metallic Pd demonstrating a higher oxidation state of Pd in 280 the PGM grains. The energy positions of the white lines are 3174.4 eV for PdCl₂, 3174.7 eV for 281 PdSO₄, and 3175.3 eV for PdS. These energies are in the same range as the energies of the PGM. 282 This indicates that Pd in the studied PGM grains has an oxidation state of +2. An oxidation state 283 of +4 for Pd is excluded since the energy position for Pd^{4+} compounds is expected to be at much 284 higher energies, 3176.8 eV for Zn₂PdO₄ (Kim et al., 2002), 3184.6 eV for [PdCl6]²⁻ (Barton, 285 2013), 3188.3 eV, $[N_4PdCl_2]^{2+}$ (Barton, 2013). The difference in energy between the measured 286 spectra varies from 3174.4 eV for PdCl₂ to 3175.6 eV for paolovite. This can be interpreted as an 287 288 electronegativity effect of neighboring ligands. The white line of the PdCl₂ spectrum has the lowest energy due to the high electronegativity of Cl (3.16) compared to other ligand atoms 289 (electronegativity of S = 2.58; Bi = 2.02; As = 2.18; Sn = 1.96; Ni 1.91; Cu = 1.9). The higher the 290 291 electronegativity of a ligand atom, the lower the energy position of the white line.

The XANES spectrum of the PGM grain from sample RM 10_3 is shown together with the spectra of the PGM named above in Figure 5a and with the Pd references in Figure 5b. The energy position of the RM 10_3 PGM spectrum is 3175.1 eV, almost identical to that of atokite.

Moreover, the second spectral feature, indicated as b' in Figure 5a, of the RM 10 3 PGM 295 296 coincides with that of atokite. Based on the presence of the intense Pt peak on the μXRF 297 spectrum for the RM 10 3 PGM grain and the resemblance of its XANES spectrum to the spectrum of atokite, we propose that the PGM in sample RM 10 3 can be identified as atokite. 298 Atokite is a common Pd-bearing mineral and is often described in sulfide mineral assemblages of 299 300 Norilsk ores (Barkov et al., 2000; Sluzhenikin et al., 2020), including Mt. Rudnaya MSS-ISS ores 301 (Brovchenko et al., 2020). Nanometer-sized atokite has also been identified as one of the most favored forms of Pd occurrence in pentlandite (Junge et al., 2015). Thus, the detection of atokite 302 in our samples is valuable for the comparison of various possible forms of Pd in minerals. 303

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Oxidation state and bonding of Pd in pentlandite

305 The four Pd hot spots in pentlandite depicted on µXRF maps (Figure 1 and Supplementary material) were selected for Pd L₃-edge XANES measurements. The XANES 306 spectra in these four pentlandite grains from four different sample areas are identical (Figure S8 307 in Supplementary material), suggesting that there is only one form of Pd identified in the Mt. 308 Rudnaya pentlandite. The Pd hot spot in sample RM 30 3 is the biggest among the detected Pd-309 310 rich pentlandite grains. Forty-one spectra were collected on the same position of this Pd hot spot and averaged to get a spectrum with better statistics. The obtained averaged spectrum is plotted 311 together with XANES spectra of PGM and Pd references in Figures 5a and 5b. The position of 312 313 the white line of the pentlandite spectrum is in the range of energies for all measured PGM and Pd references, based on that this energy position is assigned to Pd²⁺. The white line of the 314 pentlandite coincides with the energy of the white line of vasilite. This match can be explained by 315 the bonding of palladium with sulfur in the two sulfide minerals, pentlandite, and vasilite. The 316 position of the white line of the PdS reference is 0.3 eV higher than that of pentlandite. Such a 317

difference could be explained by the fact that a "mixed electronegativity" is present in pentlandite
due to the different elements (Fe, Ni, and S) involved in the bonding compared to the exclusive
"pure" bonding of Pd and S in PdS.

Figures 5a and 5b demonstrate that the white line of the pentlandite spectrum is more 321 intense compared to those of PGM, metallic Pd, and PdS. The interpretation of white line 322 intensities is challenging because several effects can influence the white line intensities of the L₃-323 edge XANES. Given small uncertainty due to the normalization of the spectra and correction of 324 325 the PGM spectra to self-absorption, these effects can be due to nanoparticle size, adsorbate effect, and alloying effect (Koningsberger et al., 2000; Zheng et al., 2011; Todorovic, 2012). We 326 exclude the contributions of size and adsorbate effect since they are not relevant to our samples. 327 328 It is known that the intensity of the white line of L₃-edges in XANES for 4d metals is correlated with the number of unoccupied d states (Muller et al., 1982; Sham, 1985). In pure metals and 329 their alloys, d states are full and the white line is less pronounced (Sham, 1985). Alloying of Pd 330 331 with other metals that can be present in the minerals and compounds showed here results in changes to the electronic structure of Pd, in particular occupancy of the d state, and consequently 332 333 decreases the intensity of the white line (Sham, 1985; Kim et al., 2002; Witjens et al., 2004). The 334 high intensity of the L_3 -edge white line of the pentlandite spectrum thus can be explained by the 335 absence of alloying of Pd in the mineral. In contrast, lower white line intensities of the PGM 336 spectra demonstrate a possible alloying of Pd with other metals in the mineral structures. Indeed, the PGM studied here contain metals (Pt, Cu, Ni) or metalloids (Sn, Te, Bi) that form short 337 bonding distances with Pd (Mihalik et al., 1975; Matkovic et al., 1976; Bayliss et al., 1990; 338 339 Evstigneeva et al., 2000) contributing to the alloying of Pd.

- The shape of XANES spectra is usually indicative of the atomic configuration. The different shape of the pentlandite spectrum compared to PGM and Pd references, therefore, confirms that Pd in pentlandite does not occur in micro-inclusions of the present Pd compounds, but has its own atomic coordination.
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Pd substitution in pentlandite

The µ-XRF and XANES results of this study confirm the existence of two forms of Pd in 345 the Mt. Rudnaya MSS-ISS ores, Pd associated with PGM inclusions and Pd incorporated in 346 pentlandite. XANES analysis demonstrates that palladium in pentlandite and PGM is represented 347 by Pd^{2+} . Following the presented results, we conclude that Pd related to pentlandite is 348 incorporated into the structure of the mineral. This is the first report of Pd incorporation in the 349 350 lattice of pentlandite from Norilsk, which is based on direct spectroscopic measurements. Considering the positive charge of Pd, we propose that Pd occupies a cationic position in the 351 pentlandite structure. Iron and nickel in the pentlandite crystal structure evenly occupy octahedral 352 and tetrahedral sites (Rajamani and Prewitt, 1975). As noted by Junge et al. (2015), the electronic 353 configuration of Pd is similar to that of Ni (dsp²) and not to Fe, which can be an argument for the 354 preferential occupation of Ni sites by Pd. Kalugin et al. (2021) also noted a preferable 355 substitution of Ni sites by Pd. And although ionic radii of Pd²⁺ for tetrahedral (0.64 Å) and 356 octahedral (0.86 Å) coordination is closer to ionic radii of Fe^{2+} (0.64 Å, 0.78 Å) than to those of 357 Ni²⁺ (0.55 Å, 0.69Å) (Shannon, 1976), our findings of the negative correlation between Pd and 358 Ni and positive correlation between Pd and Fe in pentlandite suggests that palladium prefers to be 359 substituted on the place of Ni. A resemblance of the ionic radii of Pd, Ni, and Fe accounts for an 360 361 absence of any structure distortion of pentlandite noticed on EBSD orientation maps. Whereas, in contrast, Ag addition into pentlandite (argento pentlandite (Fe₅Ni₃)AgS₈; Mandziuk and Scott, 362

363	1977) causes distortion of the pentlandite structure due to the difference of the ionic radius
364	between Ag (1.15 Å for Ag^{1+} in an octahedral configuration) and the substituted Fe and Ni
365	(Shishkin, 1972). The silver ionic radius is almost twice as much as Pd, while another isomorphic
366	component, typical for pentlandite, is Co which has a similar ionic radius (0.58 Å, 0.74 Å) to Pd.
367	It has been demonstrated that Co incorporation into pentlandite leads to a decrease in the cell size
368	(Riley, 1977) and growth of temperature range for pentlandite stability (Vaasjoki et al., 1974).
369	Riley (1977) showed that Co substitutes either Fe and Ni making two different series with
370	octahedral or tetrahedral Co atoms coordination in pentlandite structure. In the pentlandites form
371	the MSS-ISS ores of Mt. Rudnaya, Co (as from 0.29 to 0.53 wt%) negatively correlates with the
372	Fe/Ni ratio displaying Co accommodation mostly in the Fe site with a Pearson correlation
373	coefficient between Fe and Co in aphu of -0.4 (Table S1). Co and Pd contents in pentlandite from
374	J-M reef ores, Stillwater deposit, correlate positively (Godel and Barnes, 2008), although these
375	components do not correlate with each other in Pd-rich pentlandite from Talnakh deposit
376	(Kalugin et al. 2021). The Pearson correlation coefficient for Co and Pd in pentlandite from Mt.
377	Rudnaya MSS-ISS ores (in apfu) is -0.45 (Table S1), which can be described by the additional
378	substitution of Co by a minor part of Pd atoms in the Fe site.

379

Origin of Pd-rich pentlandite

As mentioned above, Pd-rich pentlandite can be formed in two ways. One way is exsolution from MSS below 450 °C (Etschmann et al., 2004, Helmy et al., 2021). Concentration of Pd dissolved in MSS at 900 °C can reach 11 wt. %. Upon cooling, Pd solubility in MSS decreases, and its concentration drops to 0.4 wt. % at 500 °C (Makovicky et al., 1986). Contact pentlandite starts to grow at 450 °C as the flame-like nano exsolutions from MSS (Helmy et al., 2021). Pd partitions into these exsolutions formed at the rims of MSS grains by diffusion.

Whereas the diffusion rate of pentlandite in MSS shows its maximum at 400 °C and drops below 386 this temperature (Etschmann et al., 2004). The second scenario for Pd-rich pentlandite formation 387 388 involves a peritectic reaction between Ni-poor MSS and Ni-rich residual sulfide melt at 870-800 °C, followed by eutectic reaction directly from the melt at ~750 °C (Kitakaze et al., 2016). 389 Although literature data on Pd solubility in high-temperature pentlandite is not available, it is 390 391 reported that pentlandite associated with Pd-rich phases contains up to 4 at.% of Pd at 550 °C in 392 the Pd-Ni-Fe-S phase system (Makovicky and Karup-Møller, 2016). Pd partitions into peritectic pentlandite from residual sulfide melt which is far more enriched in Pd than MSS. This causes 393 directional enrichment trends in the pentlandite grains: from ISS, more compatible with Pd, to 394 MSS, less compatible with Pd (Liu and Brenan, 2015; Mansur et al., 2019). 395

396 When both types of pentlandite are present, exsolved Pd-poor pentlandite can equilibrate with early-formed peritectic Pd-rich pentlandite at a grain boundary, resulting in gradual trends of 397 Pd enrichment in one grain (Barnes S.-J. et al., 2020). The crystallographic orientation of the 398 399 subsolidus pentlandite can be inherited from early-formed peritectic pentlandite. Pd-rich zones in pentlandite from Mt. Rudnaya MSS-ISS ores are confined to either rims or central parts of grains. 400 401 Pd-rich pentlandite appears as 10-100 µm loop-like and coarse grains at the contact between MSS 402 and ISS, and as up to 50 µm disseminated euhedral grains in ISS. Pd-rich zones in disseminated 403 pentlandite are surrounded by ISS and present only in the central parts of grains. In contrast, Pd-404 rich zones in pentlandite from the contact between MSS and ISS (studied in this work) are confined to the contact with ISS or the inner parts of grains, with a maximum also located closer 405 to ISS. In the scenario of pentlandite exsolution from MSS, the most enriched in Pd zones must 406 407 be at the edges of grains and disappear gradually towards the contact with MSS, because Pd, as a non-coherent element, rather enriches the rims of MSS than random parts. Pentlandite which is 408

later formed by exsolution must feed on the broken MSS at the MSS-ISS boundaries and have the 409 410 same trends of enrichment. However, the majority of Pd-rich zones are found in the inner parts of 411 pentlandite grains. More probably, this type of Pd enrichment indicates the peritectic origin of Pd-rich pentlandite when both MSS and sulfide liquid can be the feeding environments, and the 412 directional distribution of other trace and major elements confirms this (Brovchenko et al., 2020). 413 414 The disseminated pentlandite grains among ISS had an eutectic origin. However, formation of Pd-rich pentlandite exsolved from MSS was also probable and we can suggest that this 415 pentlandite could be formed after the peritectic pentlandite. As a result, these Pd-poor pentlandite 416 exsolutions equilibrated with the peritectic Pd-rich pentlandite, setting the directional trend of Pd 417 enrichment. 418

419

Implications

For the first time, Pd L₃-edge XANES has been measured in natural Pd-rich pentlandite and various natural PGM. The combined μ XRF and XANES results demonstrate that Pd incorporates in the crystal structure of pentlandite. By comparing XANES data for Pd in pentlandite and various natural PGM and synthetic Pd, PdS, PdCl₂, and PdSO₄ we exhibit a difference between Pd spectra in these common Pd-bearing phases.

The form of Pd in pentlandite and the formation of Pd-rich pentlandite from PGE ore deposits are still debated. Nano and micro-inclusions of various PGM were described in different sulfide minerals (Wirth et al., 2013), including pentlandite (Junge et al., 2015). Junge et al. (2015) also demonstrated the presence of Pd solid solutions in the lattice of pentlandite from the Bushveld complex. However, the contents of Pd in pentlandite from Junge et al. (2015) are on average 390 ppm. High concentrations of Pd in Pn from the Talnakh deposit (Kalugin et al., 2021) and J-M Reef (Li and Ripley, 2006) were achieved by the hydrothermal secondary

redistribution leading to additional incorporation of Pd predominantly in the Ni site (Kalugin et 432 al., 2021). Here, we show the first evidence of incorporation of Pd in pentlandite from the non-433 434 altered Mt. Rudnaya MSS-ISS ores as up to 4.84 wt%. Our findings leave no doubt about the possibility of Pd entering the crystal structure of pentlandite as a solid solution. The coexistence 435 of primary magmatic high-temperature sulfide phases such as MSS, ISS, and pentlandite with a 436 437 high Pd content as well as the exceptionally high bulk PGE content in ores (up to 350 ppm), support the idea of the primary magmatic origin of ores from the fractionated residual Cu-Ni-438 PGE enriched sulfide liquid described in Brovchenko et al. (2020). The rapid quenching, 439 therefore, ensured the preservation of high-temperature MSS and ISS, and the heterogeneous 440 distribution of trace and major elements. Also, the recent X-ray microtomography study by 441 Sittner et al., (2022) declares a specific globular three-dimensional texture of these ores. It seems 442 to be supportive of an idea of a possible sulfide-sulfide liquid immiscibility between Cu-richer 443 and Cu-poorer liquids in the exceptionally Cu and Ni enriched sulfide systems. The most 444 enriched in Pd grains of pentlandite (studied in the current work) are confined to these globular 445 textures, namely at the contact between Cu-richer ISS globules and Cu-poorer MSS-ISS fine-446 grained matrix. The fact that palladium could be incorporated into the structure of pentlandite 447 448 with such high concentrations in not-altered primary magmatic ores can potentially reveal that the solubility of Pd in pentlandite extends with temperature and encourages the determination of new 449 constraints on the physical-chemical conditions of ore-forming processes. The ability of BMS to 450 451 dissolve PGE at high temperatures has also been demonstrated for MSS and Pt (Makovicky et al., 1986; Majzlan et al., 2002), for MSS and Pd (Ballhaus and Ulmer, 1995) and pyrrhotite and Pt 452 (Filimonova et al., 2021). Our data confirm the possibility of Pd solubility in pentlandite. 453 Therefore, PGE could be concentrated in BMS as solid solutions indicating that the behavior of 454

455 these elements is preliminary controlled by sulfide melt. It signifies that the role of PGM 456 nanonuggets in the formation of Cu-Ni-PGE deposits has been overestimated.

457 Our results illustrate that the combination of μ XRF, XANES, and EBSD provides 458 complementary and full information on the occurrence of Pd in pentlandite. Such combined 459 investigations have great potential in understanding the mechanisms of other noble metals 460 incorporation in ore minerals.

461

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684 **Figure legends**

Figure 1. (a) Micro-XRF map of pentlandite (Pn) and two PGM grains (PGM₁ and PGM₂) RM 685 30 3 9b. Map size: $170 \times 69 \ \mu\text{m}^2$, pixel size: 3 μm ; (b) μ XRF map of PGM₃ grain RM 10 3 10a. 686 Map size: $20 \times 20 \ \mu\text{m}^2$, pixel size: 2 μm . Pd – red, Ni – green, Cu – blue. 687 Figure 2. (a) Micro-XRF spectra of four Pd "hot" spots in different pentlandite (Pn) grains. 688 Micro-XRF spectra for areas RM 29 1 8a, RM 29 3 7a, and RM 30 3 9b were collected with 689 3174 eV incident beam and 300 s collection time. µXRF spectrum for the area RM 29 1 1a was 690 obtained by summing 192 Pd-rich pixels of the µXRF map RM 29 1 1a recorded at 3174 eV 691 with 1 s per point; (b) Micro-XRF spectra of Pd-rich pentlandite in the area RM 30 3 9b, PGM 692 grain in the area RM 10 3 10a, Pd-poor pentlandite (matrix Pn) and ISS (matrix ISS). Micro-693 694 XRF spectra for areas with Pd-rich pentlandite and PGM were obtained with 3174 eV incident beam and 300 s collection time. µXRF spectra for Pn matrix and ISS matrix were obtained by 695 summing 150 and 140 pixels respectively on the μ XRF map of the area RM 29 1 recorded at 696 3174 eV with 1 s per point. 697

Figure 3. (a) BSE photo of the contact pentlandite located between ISS globule and MSS with
the EPMA profile. The EPMA data from the profile are presented in apu on (b, c) – Ni vs Pd and
Fe vs Pd correspondingly.

Figure 4. EBSD data collected from heterogeneous Pd-rich pentlandite grain, (a) and (b) correspondingly - EBSD patterns collected from Pd-rich and Pd-free areas; (c) - EDX mapping of the corresponding grain in Ni and Pd characteristic X-rays; (d) - diffraction quality map; (e) -Euler-colored orientation map and corresponding inverse pole figure, (f) - local misorientation map depicting local strains and deformations and local disorientation graph; (g) - map of

- 706 disorientation from average depicting grain modularity and disorientation directions and
- 707 corresponding coloring legend.
- **Figure 5.** Pd L₃-edge XANES spectra for Pd-rich pentlandite from the area RM 30 3 9b (Pn),
- PGM from the area RM 10_3_10a (PGM) compared to the spectra of PGM grains (a), and Pd
- references (**b**). The white line is indicated as a', the second spectral feature is indicated as b'.

712	Table 1.	List	of inv	estigated	samples.
			· · · · · ·		

Name	Sample/ inventory number	General formula	Origin	Geological setting	[Pd] wt%		
Pentlandite and PGM	RM 10, RM 29, RM 30	(Fe,Ni) ₉ S ₈ pentlandite	Mt. Rudnaya, Norilsk, Russia	magmatic	up to 4.84 ^{a,c} pentlandite		
Atokite	75793	$Pd_{2.25}Pt_{0.75}Sn$	Oktyabrsky mine, Norilsk, Russia	magmatic	47.47 ^b		
Cabriite	80761	Pd ₂ CuSn	Oktyabrsky mine, Norilsk, Russia	magmatic	46.19 ^c		
Paolovite	80761	Pd ₂ Sn	Oktyabrsky mine, Norilsk, Russia	magmatic	64.20 ^b		
Sobolevskite	80762	Pd(Bi,Te)	Oktyabrsky mine, Norilsk, Russia	magmatic	31.31 ^c		
Vasilite	80272	$Pd_{15.93}Pt_{0.18}Cu_{0.21}S_{6.68}$	Miass river, Ural, Russia	alluvial	87.9 ^c		
Majakite	75810	PdNiAs	Oktyabrsky mine, Norilsk, Russia	magmatic	44.34 ^b		
^a concentration based on LA-ICP-MS analyses							

^b concentration based on empirical mineral formula ^c concentration based on EPMA analysis

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	WL		
Sample	position,		
	eV		
Pd foil	3173.8		
PdCl ₂	3174.41		
PdS	3175.3		
PdSO ₄	3174.7		
atokite	3175		
cabriite	3175.3		
paolovite	3175.6		
sobolevskite	3175.3		
vasilite	3174.7		
majakite	3175.3		
pn in RM 30_3_9b	3174.7		
pn in RM 29_1_8a	3174.69		
pn in RM 29_3_7a	3174.69		
pn in RM 29_1_1a	3174.7		
PGM in RM 10_3_10a	3175.1		

Table 2. White line (WL) energy positions for Pd L₃-edge XANES spectra. pn – pentlandite.

716

718 Figure 1



719

721 Figure 2



722

724 Figure 3

725



728 Figure 4



729

731 Figure 5



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