Formation conditions of noble metal mineralization in sulfide cobalt-copper-nickel ores of Kamchatka (on the example of Annabergitovaya Schel ore occurrence)

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Abstract. The authors present research results, the purpose of which is to study the specifics of noble metal mineralization and its genesis in sulfide cobalt-copper-nickel ores of the Kamchatka nickel-bearing province. The paper is dedicated to one of its many ore occurrences called Annabergitovaya Schel (Annabergite Gap). The material composition of platinoids, silver, gold, bismuth and tellurium minerals, as well as sulfarsenides in the ores of this occurrence was investigated. Based on the data of mineral formation sequence and the use of geosensors, conclusions were drawn regarding the genesis of noble metal mineralization.

Formation of platinoid minerals, silver and gold at the Annabergitovaya Schel ore occurrence is mainly associated with the epigenetic effect of post-ore granitoids on ore-bearing intrusion rocks of the Dukuk complex of the cortlandite-norite formation and on syngeneic ores. An early association of noble metal minerals is represented by sperrylite, irarsite, and rare unnamed phases of Pt + Ir + Te. Irarsite and Pt + Ir + Te phases were formed at the contact-metasomatic stage. Sperrylite can be assumed to be of magmatic origin. Silver sulfides and tellurides, silver and palladium bismuth tellurides, and native gold were formed at the late, hydrothermal-metasomatic, stage. The occurrence conditions of mineral parageneses, associated with noble metal mineralization, correspond to the formation of shallow-depth metasomatic rocks (<5 km). Sub-developed quartz-feldspar metasomatites, associated with the formation of early platinoid arsenides and sulfarsenides, are in equilibrium with circumneutral solutions (pH of 4.5-6.5) at temperatures of 350-600 °C. Late hydrothermal association with Pd, Ag and Au minerals is close to propylites and was formed at pH values of 4.5-6.5 and temperature of 150-350 °C.

Key words: Kamchatka; cortlandite-norite formation; sulfide copper-nickel ores; conditions of mineral formation; platinum-group elements; gold

Introduction. Apart from being a raw material for the production of non-ferrous metals, sulfide copper-nickel ores are also an important source of platinoids [10, 13, 14, 33]. In Kamchatka, numerous ore occurrences and a medium-scale Shanuch deposit of sulfide cobalt-copper-nickel ores are located within the Kamchatka nickel-bearing province (KNP), which outlines a large block of metamorphic rocks – the Kamchatkan median massif [15, 23]. It is located in the Median (Sredniny) Range, extends over 200 km in the meridian direction and has a width of 30-40 km (Fig.1). Nickel-bearing objects are associated with the developed intrusions of the Dukuk complex of the norite-cortlandite formation (or the formation of hornblende basites, according to [7]). Sulfide ores contain noble metal minerals represented by sperrylite, irarsite, palladium telluro-bismuthides, merenskite, palladium antimonides, native platinum, isoferroplatinum and native gold [11, 13, 15, 20, 22].

The problems concerning the genesis of sulfide copper-nickel ores and the mechanisms of metal fractionation, the ratios of ore minerals, the forms of occurrence and distribution of platinum group elements (PGEs) and gold in ore bodies retain their relevance to this day [3, 4, 16, 24, 28, 29]. The genesis of cobalt-copper-nickel ores in the KNP is a debatable issue. According to most researchers, their formation is determined by magmatic processes [8, 11, 17-19, 22]. For example, for Kvinum I and II ore occurrences, it is assumed that formation of sulfide and noble metal mineralization is determined by magmatic differentiation and sulfurization under the influence of fluids. Moreover, the
productive in relation to PGEs increases with the growing role of Te, As, Sb and Bi in the fluids [11, 22]. At the same time, many researchers point to metamorphic, metasomatic and hydrothermal transformations of copper-nickel ores in the KNP [5, 12, 15].

This paper discusses one of the many KNP ore occurrences – Annabergitovaya Schel (Annabergite Gap). Its geological structure, chemical and mineral composition of ores are described in a number of publications [5, 9, 15, 20]. The ore mineral associations, identified at the Annabergitovaya Schel occurrence, were previously found at the Shanuch deposit [13] and resemble the associations of the Dambukinsky nickel-bearing area in the Upper Amur Region [21]. The composition of platinoid, silver and gold minerals is close to the noble metal mineralization in the ore body N 1 of the Shanuch deposit [13], as well as in the ore occurrences in the southern part of the KNP – Kvinum I [11] and Kvinum II [21].

The purpose of the study within the scope of this paper was to identify the specifics of material composition and genesis of noble metal mineralization at the Annabergitovaya Schel ore occurrence. New data were obtained on the composition and ratios of minerals, which made it possible to draw conclusions about the genesis of PGE, silver and gold minerals.

Research methods. In the process of studying the ores of the Annabergitovaya Schel occurrence, data on their chemical composition were obtained [20], mineralogical tests were performed using electron microscopy and X-ray diffractometry. To assess the formation conditions of noble metal minerals, authors used geothermometers, available data on thermodynamic properties of the associated minerals and results of summarizing studies [1].

Detection of Ni, Cu, Co and Au was carried out using the atomic absorption method on a Shumadzu AA-6300 spectrometer at the Scientific Research Geotechnological Centre of the Far Eastern Branch of the Russian Academy of Sciences (SRGC FEB RAS). Pt and Pd were detected by the method of atomic emission spectrometry with inductively coupled plasma on an IRIS Intrepid Duo spectrometer at the Vernadsky Institute of Geochemistry and Analytical Chemistry of the Russian Academy of Sciences. Ag was detected by means of spectral analysis at the Institute of Mineralogy, Geochemistry and Crystal Chemistry of Rare Elements.

Studies of the chemical composition of the minerals and microphotography were performed at the Institute of Volcanology and Seismology of the Far Eastern Branch of the Russian Academy of Sciences on a Tescan Vega 3 scanning electron microscope with an energy dispersive spectrometer.
Oxford Instruments X-Max 80 mm² and AZtec software. As for the reference samples, PGEs were compared to pure metals, obtained at the Moscow State Institute of Steel and Alloys, while Au, Ag, Ni and Fe were compared to the corresponding metals. Artificial compounds FeAsS and FeS₂ were used for As and S detection; synthetic minerals CdTe, CuSbS₂, Bi₂S₃, PbS, ZnS, CuFeS₂ – for Te, Sb, Bi, Pb, Zn, Cu; a Fe-Co-Ni alloy of known composition – for Co. The tests were carried out using accelerating voltage of 20 kV and absorbed current on the nickel reference sample of 700 pA.

Certain samples of ore and rock powders were studied at the SRGC FEB RAS on a Rigaku Ultima IV powder X-ray diffractometer with horizontal sample fixation. A glass cuvette was used as a sample holder. Diffraction patterns of the sample powders were recorded using a D/teX detector in the 2Θ range of 10-100°, at a speed of 1.0°/min, in standard mode. The crystalline phases were identified using the ICDD PDF-2 (Powder Diffraction File) database (version 2012) updated in 2019 and PDXL-2 software package.

**Research results.** *Geological structure of the ore occurrence, mineralogy and ore geochemistry.* Annabergitovaya Schel, an occurrence of rich sulfide cobalt-copper-nickel ores, is located in the southern part of the KNP (Fig.1, a), in the upper reaches of the river Stepanova (53°24’56.1” N and 157°43’4” E). The ore body outcrops in a narrow glen, formed along the rupture fault zone. The length of the ore outcrop is 18 m along the thalweg; the height of the western vertical edge is 11 m (Fig.1, b). The ore body is composed of massive, vein-disseminated, nested and disseminated sulfide ores. The contacts with the host rocks are tectonic, with zones of fragmentation, boudinage and foliation [20]. Massive unoxidized ores contain on average 4.82 wt.% of Ni, 1.95 wt.% of Cu and 0.13 wt.% of Co, as well as 0.036 g/t of Pt and 1.12 g/t of Pd. The Au content reaches 0.08 g/t, the Ag concentration is on average 6.66 g/t [20].

The ores are associated with the Dukuk complex intrusion – a deformed sill-like body a few tens of meters thick in the apoterrigenous schists of the Kheivan formation, metamorphosed under the conditions of greenschist and amphibolite facies [5]. The intrusion is composed of predominant highly altered ultramafites (probably, olivine websterites [15]), as well as gabbronorites, norites and diorites (Fig.1, b). These rocks and their syngenetic associates were transformed as a result of contact metamorphism and metasomatose under the influence of nearby post-ore intrusions of granitoids [12], which on a 1:1,000,000-scale geological map are attributed to the second phase of the Kola plagiogranite-graniodiorite complex [5]. At the ore occurrence, there is a wide representation of hybrid rocks, their composition varying from melanocratic diorites to biotite-hornblende plagiogranites. The ore body and the surrounding schists intersect associated with granitoids veins of quartz, quartz-feldspar, sometimes muscovite, which are a few centimeters in thickness. Quartz-feldspar areas with a micropegmatitic structure are typical for hybrid rocks, granitoids and veins.

The main ore minerals are pyrrhotite (predominantly monoclinc, 6C), chalcopryite and pentlandite. Pyrite and marcasite are common. Hypergene minerals are widespread, such as iron hydroxides, morenoisite, retgersite, malachite and azurite. As for rare minerals, it was detected that the ores contained nickel and cobalt sulfarsenides, galena, sphalerite, as well as compounds of bismuth and tellurium [20]. X-ray diffraction studies of the ores revealed the presence of cobaltpentlandite, nickel-bearing pyrite, bornite, covellite, colusite and scheelite. Minerals of noble metals are mainly represented by sperrylite [15] and irarsite. Hessite and volynskite are less developed [20]. Native gold and argentopentlandite are occasionally found [20], as well as michenerite [15].

**Mineral associations.** Ore minerals form three mineral associations (or parageneses, according to [9]), which are also typical for the Shanuch ore field [13]: pentlandite-pyrrhotite, pyrrhotite-pentlandite-chalcopryte and pyrite-marcasite ones [9]. The associations are distinguished according to the specifics of their mineral composition, taking into account quantitative ratios of ore minerals and their morphology. They partially characterize the sequence of ore formation and reflect primary zonality of sulfide ores, which is formed at the magmatic stage in the process of sulfide melt differentiation [10].
Pentlandite-pyrhotite association, the formation of which is attributed to the magmatic stage, is represented by sulfides of the early generation (I): porphyry segregations of pentlandite and pyrrhotite with reticulate pentlandite [9]. At the Shanuch ore field, the composition of this association also includes early chalcopyrite I [13]. At the studied ore occurrence, the association is weakly developed, which possibly reflects primary zonality of sulfide ores, formed in the process of magmatic differentiation of the sulfide melt.

Pyrrhotite-pentlandite-chalcopyrite association is represented by pyrrhotite with flame- and plate-like segregations of pentlandite II, chalcopyrite II, sulfarsenides of nickel and cobalt (I), sperrylite and irarsite. The association is formed as a result of syngenetic processes and subsequent transformations under the influence of post-ore granitoids of the Kola complex. Contact changes in igneous rocks of the Dukuk complex intrusion are manifested in the replacement of primary plagioclase by albite and replacement of tschermakites, pargasites, pyroxenes and olivines by magnesio-hornblende, anthophyllite, cummingtonite, as well as aggregates of these amphiboles, including segregations of sphene and rutile. As a result of contact-metasomatic transformations (sub-developed quartz-feldspar metasomatose), an aggregate of oligoclase, potassium feldspar and quartz is represented in some areas of the ore body; a replacement of phlogopite by biotite is observed, as well as replacement of early amphibole aggregates by quartz and feldspars.

Pyrite-marcasite association is formed in the process of subsequent hydrothermal-metasomatic transformation of the minerals in early sulfide associations: pentlandite is replaced by violarite, pyrrhotite – by pyrite and marcasite. Notably, pyrrhotite, which occurs in paragenesis with pyrite and marcasite, is characterized by higher sulfur content. Redistribution of chalcopyrite I and II led to the formation of its late variation (chalcopyrite III [13, 21]). Galena, native gold, volynskite, hessite, argentopentlandite and tellurium-bismuth mineralization were also formed at this stage. The silicates were replaced by clinohlore, epidote, actinolite and sphene, in some areas – by talc and carbonate.

Specifics of noble metal mineralization. Early and late noble metal mineralizations are distinguished at the ore occurrence. The early one belongs to the pyrrhotite-pentlandite-chalcopyrite association and is represented by sperrylite, irarsite and an unnamed compound of Pt, Ir and Te. Early minerals of noble metals are spatially and genetically related to sulfarsenides of the cobaltite group, which often contain isomorphic traces of platinoids. The presence of gersdorffite is confirmed by X-ray diffractometry results. The late mineralization belongs to the pyrite-marcasite association and is represented by michenerite, volynskite, hessite, argentopentlandite, native gold and Ag-containing tellurobismuthite (Table 1).

| No. | Mineral                        | Chemical formula |
|-----|--------------------------------|------------------|
|     | **Platinoid minerals**          |                  |
| 1   | Sperrylite                      | PtAs₂            |
| 2   | Irarsite                        | IrAsS            |
| 3   | Michenerite                     | PdBiTe           |
|     | **Silver and gold minerals**    |                  |
| 4   | Volynskite                      | AgBiTe₂          |
| 5   | Argentopentlandite              | Ag(Fe, Ni)₃S₈   |
| 6   | Hessite                         | Ag₂Te           |
| 7   | Native gold                     | Au, Ag          |
|     | **Minerals, containing noble metals in isomorphic form** |             |
| 8   | Sulfarsenides (cobaltite-gersdorffite series) | NiAsS, traces of Co, Fe, Ir, Pt, Rh, Pd |
| 9   | Tellurobismuthite               | Bi₂Te₃, traces of Ag, Pd |

Early noble metal mineralization. Sperrylite and irarsite form round or platy inclusions in sulfarsenides (intermediate members of the cobaltite-gersdorffite series). The size of sperrylite segregations varies from 2.5 to 2 μm, their composition is close to the theoretical formula (Table 2). The size
of irarsite inclusions ranges from 2 to 5 μm (Fig. 2, d), the mineral usually contains traces of Pt (on average 5.22 wt.% and Rh (on average 1.85 wt.%).

Sulfarsenides associated with platinooids form idiomorphic crystals up to 0.1-0.5 mm in size, which are usually confined to the grain borders of main ore-forming sulfides and often have an isometric hexagonal shape of the cross-section (Fig. 2, b, g). Small grains of sulfarsenides are usually observed in chalcopyrite, while larger grains are found in pyrrhotite. Less often their inclusions are detected in pentlandite, pyrite and violarite. The crystals of sulfarsenides in pyrrhotite are characterized by flame-like segregations of pentlandite that form along their edges (Fig. 2, a).

The ratios and the segregation sequence of sperrylite, irarsite and Fe-Co-Ni sulfarsenides were examined in an epitaxial intergrowth of these minerals, located at the border of the violarite grain (Fig. 3, a-d). Inside the hexagonal crystal of Fe-Co-Ni sulfarsenide, there is an elongated sperrylite crystal with irarsite-rimmed edge, in which an unnamed compound of Te, Ir and Pt is detected (Fig. 3, c-f). An idiomorphic, probably cubic mineral of the cobaltite-gersdorite series with platinooids (hereinafter referred to as sulfarsenide I) is naturally rimmed by sulfarsenide of the second generation (sulfarsenide II) with increased Co content. Its rhombic cross-section also has an idiomorphic outline. It can possibly be a nickel-containing gersdorite (rhombic).

On the cross-section through this intergrowth (Fig. 3, g), one can observe the absence of zonality and relatively constant compositions of sulfarsenides I and II. With sulfarsenides of both generations having almost equal contents of As and S, the differences in their compositions are manifested in varying ratios of Ni and Co with insignificant changes in the Fe content (Fig. 3, g). In addition to inclusions of platinooid minerals, sulfarsenide I may contain their traces (Table 3), probably as a result of isomorphic substitution: Rh – up to 0.83 wt.%, Pd – up to 0.53 wt.%, Pt – up to 0.54 wt.%, Ir – up to 2.56 wt.%. Moreover, sulfarsenide I is observed to contain Cu (up to 1.67 wt.%), as well as Se (up to 0.32 wt.) and Te (up to 0.32 wt.). Sulfarsenide II is characterized by the absence of platinooids, increased Co content (around 17 wt.%) and a close to constant ratio of Ni/Fe contents (Fig. 4).

**Table 2**

| No. | Pt    | Rh | Ir | Pd | As | S | Total | Empirical formula |
|-----|-------|----|----|----|----|---|-------|------------------|
| 1   | 53.88 | –  | –  | –  | 46.12 | – | 100.00 | Pt0.93As0.07     |
| 2   | 58.73 | –  | –  | –  | 41.27 | – | 100.00 | Pt1.06As0.94     |
| 3   | 56.96 | –  | –  | –  | 43.04 | – | 100.00 | Pt1.01As0.99     |
| 4   | 55.88 | –  | –  | –  | 44.12 | – | 100.00 | Pt0.98As0.02     |
| 5   | 56.15 | –  | –  | –  | 43.85 | – | 100.00 | Pt0.99As0.01     |
| 6   | 52.15 | 4.49 | –  | –  | 43.36 | – | 100.00 | Pt0.02Ir0.08As0.00 |

Average values (n = 15)

|   | Pt    | Rh | Ir | Pd | As | S | Total | Empirical formula |
|---|-------|----|----|----|----|---|-------|------------------|
| 7 | 56.00 | 0.30 | –  | –  | 43.71 | – | 100.00 | Pt0.99Ir0.01As0.01 |
| 8 | 7.57  | 1.68 | 51.13 | –  | 29.69 | 9.92 | 100.00 | (Ir0.70Pt0.11Rh0.09)0.91As1.12Se0.88 |
| 9 | 6.39  | 2.84 | 51.62 | –  | 27.94 | 11.22 | 100.00 | (Ir0.74Pt0.09Rh0.06)0.91As1.03Se0.97 |
| 10| 7.24  | 1.01 | 51.72 | 0.76 | 28.85 | 10.42 | 100.00 | (Ir0.76Pt0.08Rh0.03Pd0.02)0.91As1.08Se0.92 |

Average values (n = 10)

|   | Pt    | Rh | Ir | Pd | As | S | Total | Empirical formula |
|---|-------|----|----|----|----|---|-------|------------------|
|   | 4.82  | 2.64 | 53.16 | 0.08 | 28.36 | 10.95 | 100.00 | (Ir0.77Rh0.07Pt0.07)0.91As1.05Se0.95 |

Note. 1-6 – sperrylite; 7-10 – irarsite. Empirical formula (Pt, Rh, Ir)0.01AsS was used to calculate formula coefficients for irarsite. In the process of microprobe analysis, due to the small size of PGE phases, the beam also ”captured” the gersdorite matrix and probably thin microinclusions: gersdorite and irarsite in sperrylite, gersdorite in irarsite. When converting results to 100 wt.% and calculating the formulas, these ”captured” phases were excluded.
Fig. 2. Ratios of noble metal minerals and tellurobismuthides to sulfarsenides, main sulfides, violarite, biotite, quartz and clinohlole (all images are generated from backscattered electrons):

- **a** – flame-like segregations of pentlandite (Pn) in pyrrhotite (Po) along the faces of sulfarsenide crystal (Sf); **b** – idiomorphic crystal of sulfarsenide with a thin inclusion of irarsite (Irs) in pyrrhotite, which contains flame-like growths of pentlandite and an area replaced by violarite (Viol); **c** – magnified image of the sulfarsenide crystal edge, rimmed by thin aggregate compounds of bismuth and tellurium, probably volynskite and tsumoite (Ts + Vln), as well as native gold (Au);
- **d** – magnified fragment of the sulfarsenide crystal with an irarsite inclusion; **e** – magnified fragment of the sulfarsenide crystal edge with gold segregations; **f** – pyrrhotite (with composition close to Fe₇S₇) includes an idiomorphic crystal of sulfarsenide with a grain of tellurobismuthite (Tb), developing along one of its faces;
- **g** – chalcopyrite (Ccp) includes a sulfarsenide crystal, xenomorphic in relation to biotite (Bt) and idiomorphic in relation to chlorite (Chl) and chalcopyrite; a thin aggregate of hessite (Hes) is observed on one of sulfarsenide faces near the contact with chlorite; argentopentlandite (Pn(Ag)) develops on the contact with silicates (Sil) in chalcopyrite; **h** – inclusion of argentopentlandite in chalcopyrite (probably, along the healed fracture); **i** – michenerite (Mcn) with a thin inclusion of hessite, developing along the edge area of chalcopyrite grain; **j** – volynskite (Vln) in chalcopyrite; **k** – idiomorphic crystal of tellurobismuthite in chalcopyrite; **l** – biotite (Bt) intergrown with quartz (Q), being replaced by the clinohlole aggregate, which contains platy segregations of late generation chalcopyrite, thin inclusions of tellurobismuthite and rutile (Rt).
Fig. 3. Epitaxial intergrowth of sperrylite, irarsite and Fe-Co-Ni sulfarsenide crystals
(all images are generated from backscattered electrons):

a – proportion of the intergrowth of sulfarsenides, sperrylite and irarsite (Sf+Sprl+Irs) to chalcopyrite (Ccp), pentlandite (Pn), violarite (Viol); b – magnified fragment with the intergrowth of sulfarsenides and sperrylite, where the crystal of sulfarsenide II (Sf II, glaucodot) is rimmed by idiomorphic sulfarsenide I (Sf I, cobaltite-gersdorffite series), in the central part of which there is an intergrowth of sperrylite and irarsite (Sprl + Irs); c, d, e, f – the intergrowth in beams of cobalt (contours of sulfarsenide I, sulfarsenide II and PGE crystals), c, platinum (sperrylite, d), iridium (irarsite, e), tellurium (unnamed phase, a compound of tellurium, platinum and iridium, f); g – a cross-section through the intergrowth of sulfarsenides and sperrylite.

An image in cobalt beams was used as an underlay for the cross-section.

Contents of S, As, Fe, Co, Pt, Ni are expressed in weight percentage.

### Table 3

| No. | S  | As | Fe  | Co  | Ni | Cu | Rh | Pd | Pt | Ir | Total |
|-----|----|----|-----|-----|----|----|----|----|----|----|-------|
| 1   | 17.80 | 47.53 | 5.23 | 5.86 | 23.71 | – | – | – | – | – | 100.13 |
| 2   | 18.20 | 46.92 | 6.17 | 7.67 | 21.16 | – | – | – | – | – | 100.11 |
| 3   | 17.88 | 46.62 | 6.81 | 9.00 | 19.55 | – | – | – | – | – | 99.85 |
| 4   | 18.34 | 46.61 | 6.16 | 10.42 | 17.77 | – | – | – | – | – | 99.30 |
| 5   | 18.38 | 44.94 | 6.41 | 11.32 | 17.30 | 1.67 | – | – | – | – | 100.02 |
| 6   | 18.14 | 46.74 | 6.29 | 8.93 | 19.01 | – | 0.41 | – | – | – | 99.53 |
| 7   | 17.90 | 45.94 | 5.92 | 9.26 | 18.44 | – | 0.82 | – | – | – | 98.28 |
| 8   | 18.34 | 46.15 | 5.86 | 10.79 | 16.63 | – | 0.83 | 0.34 | 0.54 | – | 99.46 |
| 9   | 17.31 | 45.26 | 4.92 | 6.21 | 22.29 | – | 0.51 | 0.53 | – | – | 98.60 |
| 10  | 18.79 | 46.07 | 4.89 | 17.07 | 13.08 | – | – | – | – | – | 99.91 |
| 11  | 18.86 | 44.68 | 4.89 | 16.87 | 12.89 | – | – | – | – | – | 98.2 |

Note. 1-9 – sulfarsenide I (cobaltite-gersdorffite series), 10, 11 – sulfarsenide II (glaucodot).
Late noble metal mineralization (Table 4). Aggregates of bismuth and tellurium compounds, including hessite, are found in chalcopyrite and clinoclase aggregates rimming the edges of idiomorphic crystals of sulfarsenide I (see Fig. 2, b, c, f, g). The size of hessite segregations equals 1-10 μm. In addition to that, hessite is detected in the form of distributions in michenerite, a single segregation of which has an elongated shape of 120×25 μm and develops along the edge of the chalcopyrite grain (Fig.2, g).

Apart from hessite, Ag minerals are represented by argentopentlandite and volynskite. Argentopentlandite forms segregations of 15-30 μm, develops along the fractures in chalcopyrite and in the edge areas of its grains (Fig. 2, g, h). The Ag content in argentopentlandite ranges from 8.91 to 12.98 wt.%, on average it equals 11.61 wt.% (n = 8). Volynskite is represented by single small (up to 10 μm) inclusions in chalcopyrite (Fig. 2, j); it also forms thin aggregates with tsumoite (?), rimming the edge of sulfarsenide I (Fig. 2, c). A negligible amount of Ag (from 0.55 to 6 wt.%) is present in tellurobismuthite, which forms idiomorphic crystals of about 15 μm in chalcopyrite (Fig. 2, k) and inclusions of about 3 μm in chlorite (Fig. 2, l).

Table 4

| No. | S  | Fe  | Co  | Ni  | Cu  | As  | Te  | Pd  | Pb  | Bi  | Total |
|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-------|
| 1   | 31.64 | 32.93 | –   | 22.33 | 1.05 | –   | 12.03 | –   | –   | –   | 99.98 |
| 2   | 31.67 | 34.7 | –   | 23.11 | 1.75 | –   | 8.91  | –   | –   | –   | 100.14 |
| 3   | 0.32  | –   | –   | 0.85 | –   | 60.81 | 36.81 | –   | –   | –   | 98.78 |
| 4   | 4.68  | 1.81 | 1.52 | 8.13 | 49.21 | 31.17 | –   | –   | –   | –   | 100.51 |
| 5   | 1.75  | –   | 2.5 | –   | 16.3 | 42.74 | –   | –   | –   | –   | 35.87 |
| 6   | 2.11  | –   | 2.81 | 13.12 | 40.97 | –   | 1.8  | 39.49 | 100.31 | –   | 100.17 |
| 7   | 0.92  | –   | –   | –   | 0.55 | 48.61 | –   | –   | 48.95 | 99.04 | –   |
| 8   | 2.54  | –   | –   | –   | 1.53 | 46.21 | –   | 3.3  | 45.19 | 98.76 | –   |
| 9   | 2.37  | –   | –   | –   | –   | 6.00 | 47.02 | –   | 23.9 | 43.21 | 100.99 |
| 10  | –    | –   | –   | –   | –   | 31.58 | 24.42 | –   | 39.57 | 98.95 | –   |

Empirical formula: [Ag0.91Cu0.14]2.07[Fe34.81Ni3.10]-7.91Sb0.04

Note. 1, 2 – argentopentlandite; 3, 4 – hessite; 5, 6 – volynskite; 7-9 – tellurobismuthite; 10 – michenerite. Formulas of volynskite (for 4 atoms), tellurobismuthite (for 5 atoms) were calculated without iron, which is probably included in the hydroxide composition.

Individual micro-inclusions of high-fineness native gold (~ 93.6 wt.% of Au) less than 2 μm in size are found in chalcopyrite (Table 5). Gold (~80 wt.% of Au) in the form of segregations and a thin rim occurs on the edge of sulfarsenide I crystals (Table 5, Fig. 2, c, e). A segregation of relatively low-fineness gold (~ 71.5 wt.% of Au), 6-7 μm in size, was detected in violarite (Table 5).

Estimation of formation temperatures of noble metal mineralization. Formation conditions of early platinoids were estimated based on paragenetically related sulfarsenides I using the NiAsS-FeAsS-CoAsS diagram (Fig. 5, a), which shows the isotherms of solvus temperatures according to [31]. Judging from these data, sulfarsenides I were formed at the temperatures of around 400-500 °C.
Table 5

Representative compositions of native gold (wt.%)  

| Component | 1     | 3     | 4     | 5     | 6     | 7     |
|-----------|-------|-------|-------|-------|-------|-------|
| Au        | 91.10 | 79.24 | 80.93 | 70.83 | 72.12 |
| Ag        | 5.62  | 20.76 | 19.07 | 29.17 | 27.88 |
| Cu        | 3.28  | –     | –     | –     | –     |
| Total     | 100   | 100   | 100   | 100   | 100   |
| Fineness of gold, ‰ | ~ 936 | ~ 800 | ~ 715 |        |        |
| Commentary | Inclusion in chalcopyrite | Inclusion in gersdtrite | Inclusion in violarite |

Fig. 5. Geothermometer diagram for sulfarsenides in the coordinates NiAsS-FeAsS-CoAsS (a) and for chlorites in the coordinates $R^{2+}$ (sum of $Fe^{2+}$ and $Mg^{2+}$) – $Si$ (b): 

- $a$ – position of figurative points, corresponding to the compositions of sulfarsenide 1, solvus lines under different temperatures ($°C$) are presented according to [31];
- $b$ – diagram of semi-empirical chlorite geothermometer according to [26]; data from 31 tests were used, including the ones from Table 6; Sud – sudoite, Ame – amesite, Afcl – Al-free chlorite, $R^{3+}$ – trivalent cations ($Fe^{3+}$, $Al^{3+}$), vac – vacancy in the octahedral shell.

Clinohlore is observed to integrow with Te and Bi compounds and associated chalcopyrite and pyrite (see Fig. 2, l). This presents an opportunity to estimate formation conditions of the late noble metal mineralization using chlorite and pyrite-pyrrhotite geothermometers. The temperature of clinohlore formation was estimated using empirical chlorite geothermometers according to [27] and [30], and a semi-empirical one according to [26]. Geothermometers according to [27, 30] are based on the linear correlation of $°C$-$Alv$ in chlorites. Temperature estimates obtained according to [27] vary from 119 to 337 $°C$; the ones obtained according to [30] – from 124 to 335 $°C$ (Table 6).

Table 6

Compositions of clinochlore (wt.%)  

| Component | A107-1 | A113-7 | A113-5 | GR7 |
|-----------|--------|--------|--------|-----|
| SiO$_2$   | 27.51  | 28.18  | 28.30  | 30.60 | 32.17 |
| TiO$_2$   | –      | –      | –      | 0.25  | –     |
| Al$_2$O$_3$ | 20.15 | 18.81  | 20.88  | 17.42 | 16.78 | 12.77 |
| Cr$_2$O$_3$ | 0.68  | –      | 0.25   | –     | –     |
| FeO      | 17.23  | 18.80  | 13.10  | 18.00 | 17.19 | 24.21 |
### End of Table 6

| Component | A107-1 | A113-7 | A113-5 | GR7 |
|-----------|--------|--------|--------|-----|
|           | 1      | 2      | 3      | 4   | 5   | 6   |
| V_2O_3    | –      | –      | –      | –   | 0.36| –   |
| MnO       | –      | –      | –      | –   | –   | –   |
| MgO       | 21.90  | 20.57  | 24.35  | 21.54|21.30|14.49|
| NiO       | 0.50   | 0.40   | 0.30   | 0.56 |0.61 |1.34 |
| K_2O      | 0.23   | –      | –      | –   | –   | –   |
| Total     | 88.19  | 86.76  | 87.35  | 87.12|86.59|84.98|

**Number of cations in the formula with 14 atoms of O**

| Component | A107-1 | A113-7 | A113-5 | GR7 |
|-----------|--------|--------|--------|-----|
| Si        | 2.763  | 2.901  | 2.797  | 3.013|3.094|3.439|
| Ti        | –      | –      | 0.013  | –   | –   | –   |
| Al<sup>IV</sup> | 1.238 | 1.100  | 1.204  | 0.988|0.907|0.561|
| Al<sup>III</sup> | 1.154 | 1.177  | 1.230  | 1.101|1.115|1.062|
| Cr        | 0.054  | –      | 0.020  | –   | –   | –   |
| Fe<sup>II</sup> | 0.000 | –      | 0.034  | 0.034|0.100|0.265|
| Fe<sup>III</sup> | 1.470 | 1.586  | 1.045  | 1.477|1.353|1.887|
| Mn        | –      | –      | –      | –   | –   | –   |
| Mg        | 3.278  | 3.157  | 3.587  | 3.268|3.237|2.310|
| Ni        | 0.041  | 0.033  | 0.024  | 0.046|0.050|0.115|
| K         | 0.059  | –      | –      | –   | –   | –   |
| OH        | 8.000  | 8.000  | 8.000  | 8.000|8.000|8.000|
| Fe<sup>II</sup>/Fe<sup>III</sup> | 0.31  | 0.33   | 0.23   | 0.31 |0.29 |0.45 |

**T, °C [27]**

|       | 337 | 292 | 326 | 256 | 230 | 119 |
|-------|-----|-----|-----|-----|-----|-----|

**T, °C [31]**

|       | 335 | 292 | 322 | 256 | 230 | 124 |
|-------|-----|-----|-----|-----|-----|-----|

**Note.** The ratio of Fe<sup>II</sup>/Fe<sup>III</sup> and number of OH ions were calculated assuming complete filling of shells.

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**Fig. 6.** The distribution pattern of arsenides and sulfarsenides in the rim of a large inclusion, composed of pyrrhotite and pentlandite, in the mass of post-ore granitoids:

- **a** – pyrrhotite segregation with pentlandite growths (light grey area), around 5 mm in size, in the vein hosted by plagiogranite (dark grey background), composed of quartz and feldspar (prevailing minerals), as well as biotite and hornblende. A large sulfide spot is surrounded by scattered small fragments of sulfides, tenth-hundredth of a millimeter in size, as well as by biotite and hornblende;
- **b** – a magnified fragment of the contact between the large sulfide inclusion (Po + Pn) and plagiogranite, the rimming of sulfide grains by biotite (Bt) and hornblende (Hbl) is clearly visible in the main mass of quartz and feldspar (Q + Ps).
The semi-empirical chlorite geothermometer according to [26] is a diagram in the coordinates R$^2\text{+}$ (sum of Fe$^{2+}$ and Mg$^{2+}$) – Si (see Fig. 5, b). The formulas were calculated for 14 atoms of oxygen, all iron was assumed to be divalent. According to this diagram, estimated temperatures of clinohydroxy formation vary from ~ 110 to more than 300 °C.

Pyrrhotite with increased sulfur content, the composition of which is close to Fe$_7$S$_7$, forms a part of the pyrite-marcasite mineral association. It occurs in paragenesis with pyrite, which forms under equilibrium thermodynamic conditions in the process of hydrothermal transformations. There is no contradiction between this fact and the observed ratios of nickel and cobalt in pyrites and pyrrhotites, which presents an opportunity to estimate the temperatures using a geothermometer according to [2], based on Co distribution between these minerals (Table 7). The calculated temperatures, obtained using average Co content, vary from 263 to 323 °C (Table 7, average value of ~ 296 °C) [2]. In this case, the estimate for Ni is incorrect, since its concentration in pyrites exceeds 1.5 wt.%. As a result, the isotherms of Ni distribution between pyrite and pyrrhotite deviate from the straight-line relationship [2].

| Sample No. | Co content, wt.% | (Co, at.%/Fe, at.%): 10$^{-2}$ | $K_{Co}$ | T, °C |
|------------|------------------|-------------------------------|---------|-------|
|            | Pyrrhotite | Pyrite | Pyrrhotite | Pyrite |
| A113-7     | 0.38 (n = 6) | 0.70  | 0.920    | 1.700  | 0.375 | 323 |
| GR10       | 0.32 (n = 6) | 0.48 (n = 4) | 0.690    | 1.050  | 0.493 | 301 |
| GR7        | 0.52 (n = 2) | 0.45  | 1.170    | 1.010  | 0.837 | 263 |

Note. The estimation of Co content in pyrite and pyrrhotite was based on the average of values measured in different points of the same sample; n – number of measurements; $K_{Co}$ – coefficient of Co distribution between pyrite and pyrrhotite.

Discussion of results. The segregation of early noble metal minerals and paragenetically related sulfarsenide I occurs in the following sequence: sperrylite → irarsite → Pt + Ir + Te phase → sulfarsenide I (with traces of platinoids). The earliest platinum mineral is sperrylite. It is possibly of magmatic origin, and its segregates act as crystallization centers for other platinoid minerals and Fe-Co-Ni sulfarsenides. The contact-metasomatic origin of sulfarsenide I is explained by the intrusion of post-ore granitoids, which is proved by the ratios of minerals, illustrated in Fig. 6. In Fig. 6, a 5-mm pyrrhotite inclusion is enclosed in a vein hosted by plagiogranite, which is composed of an orthoclase-oligoclase-quartz aggregate with inclusions of biotite and hornblende. A large sulfide segregation is rimmed by chains of smaller, acute-aged sulfide inclusions. All the grains of ore minerals in the quartz-feldspar aggregate are surrounded by reaction rims of biotite and magnesi-hornblende (Fig. 6, b). Observed proportions indicate the fragmentation of the large sulfide grain in the process of granitoid intrusion, accompanied by the formation of dark-colored silicates around the sulfides as a result of contact-metasomatic processes.

The plot of As contents, a cross-section through the large sulfide grain and surrounding plagiogranite (Fig. 6, a) demonstrates that arsenides and sulfarsenides are attracted to accumulations of biotite and hornblende, rimming small ore grains in plagiogranite, whereas the large sulfide segregation contains no As-bearing phases. Hence, formation of sulfarsenides, as well as the rims of dark-colored minerals around sulfides, is related to contact-metasomatic processes. It is also accompanied by redistribution of PGEs, associated with syngenetic sulfide ores, which results in the formation of irarsite and the Pt + Ir + Te phase, paragenetically related to sulfarsenide I.

The absence of ore melting signs in the near-contact area of the post-ore intrusion of granitoids correlates with experimental and calculated data. The melting of sulfide mineral associations occurs in the temperature interval from 850 to 1200 °C [10]. At the same time, estimations of formation
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Stages of ore redistribution | Contact-metasomatic | Hydrothermal-metasomatic
---|---|---
| | High-temperature | Medium-temperature | Low-temperature
| Overburden pressure, GPa | ≤0.15 | | |
| Temperature, °C | 600 | 500 | 400 | 300 | 200 | 100 |
| pH | 6.5-4.5 | | | |

| Mineral associations | Minerals |
|---|---|
| Pyrrhotite-pentlandite-chalcopyrite association | Sperrylite, Irarsite, Pt + Ir + Te phase, Sulfarsenide I, Pyrrhotite (4Fe, Fe/S), Pentlandite II, Chalcopyrite II |
| Pyrite-marcasite association | Sulfarsenide II, Pyrrhotite (Fe,S), Pyrite, Marcasite, Chalcopyrite III, Chlorite, Tellurobismuthite, Volynskite, Violarite, Argentopentlandite, Michenerite, Hessite, Native gold |

Fig. 7. A chart representing the sequence and conditions of formation for noble metal minerals (in bold), minerals with isomorphic traces of noble metals (in italics), associated main and widely-occurring sulfides and chlorites at the Annabergitovaya Schel ore occurrence (using data from [13]).

1 – formation conditions are estimated using geothermometers and mineral ratios; 2 – only mineral ratios and indirect data; 3 – solid-phase transformations without resulting formation of new mineral species.

Conditions for hornblende in the petrotypical massif of granitoids of the Kola complex are less extreme: 513-552 °C at pressure of 0.17-0.21 GPa [22]. These conditions correspond to the estimated formation temperatures of sulfarsenides I and associated early platinoid sulfarsenides (400-500 °C, Fig. 5, α), as well as to the capture temperatures of fluid inclusions in quartz from the ores of the Shanuch deposit (~ 400-500 °C) [8]. Taken together, all these data are consistent with the formation conditions of quartz-feldspar metasomatites [1], which are in equilibrium with circumneutral solutions (pH of 4.5-6.5) at temperatures of 350-600 °C, depths of ≤5 km and pressure below ~ 0.15 GPa (Fig. 7).

At the late hydrothermal-metasomatic stage, the following formation sequence of noble metal minerals is observed: tellurobismuthite → volynskite (an aggregate of tsumoite (?) and volynskite) → argentopentlandite → michenerite + hessite. Formation of native gold probably did not occur before the formation of tellurides. Glaucoidot (sulfarsenide II), which is known as a high-temperature hydrothermal mineral, probably belongs to the same stage of mineral formation. Due to hydrothermal origin of glaucoidot, the diagram from [31] is inapplicable to it, since it has been developed for “dry” systems [6].

Hydrothermal-metasomatic transformations of ores, associated with late noble metal mineralization, pyrite, marcasite, and clinohore-epidote-actinolite paragenesis, are close to propylitization processes at pH values of 4.5-6.5 and temperatures of 150-350 °C at depths ≤5 km [1]. It is confirmed
by estimated formation temperatures for chlorites (from 110 to 340 °C), pyrites and late pyrrhotites (around 296 °C). The connection of clinochlore-epidote-actinolite paragenesis to Bi and Te compounds emphasizes its medium- to low-temperature nature: the formation temperatures of tellurides are less than 350 °C, usually below 250 °C [25], and hessite is one of characteristic minerals of low-temperature hydrothermal associations. The exception in this case is provided by the Pt + Ir + Te phases, segregated together with sulfarsenide I (Fig.7).

**Conclusion.** Cobalt-copper-nickel sulfide ores of the KNP, containing noble metals, were formed as a result of magmatic differentiation in the process of intrusion development in the Dukuk complex of the norite-cortlandite formation and further epimagmatic transformations. Performed studies allowed to specify the characteristics of material composition of platnoid, silver, gold, bismuth and tellurium minerals, as well as closely associated sulfarsenides in the ores of one of the KNP ore occurrences – Annabergitovaya Schel. Identification of the sequence of mineral formation and the use of geosensors made it possible to estimate the formation conditions for the minerals of PGE, silver and gold.

The formation of noble metal minerals at the Annabergitovaya Schel ore occurrence is mainly caused by epigenetic processes; magmatic origin can only be assumed for sperrylite.

The early, contact-metasomatic, stage is associated with intrusion of post-ore granitoids. It led to the development of an association of sperrylite, irarsite, rare unnamed phases of Pt + Ir + Te and sulfarsenide I. According to [31], the formation temperature of sulfarsenide I is 400-500 °C. It agrees with the estimated conditions for the formation of quartz-feldspar metasomatites – at temperatures of 350-600 °C and depths of ≤5 km (pressure of ~ ≤0.15 GPa), in equilibrium with circumneutral solutions (pH of 4.5-6.5) [1].

Argentopentlandite, volynskite, native gold, michenerite and hessite were formed as a result of late high- to medium-temperature hydrothermal-metasomatic transformations. The association of hydrothermal minerals, paragenetically related to them, is close to propylites. The conditions of propylitization – at depths of ≤5 km (~ ≤0.15 GPa), pH values of 4.5-6.5 and temperature of 150-350 °C [1] – are consistent with the temperature estimates, obtained with pyrite-pyrrhotite (about 296 °C) and chlorite geothermometers (from 110 to 340 °C).

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