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Platinum transport in chloride-bearing fluids and melts: insights from in situ X-ray

absorption spectroscopy and thermodynamic modeling

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ABSTRACT

Hydrothermal chloride-rich fluids are identified at the late stages of the formation of platinum group element (PGE) deposits in giant layered intrusions, and are considered as the PGEs transport media in Cu(-Mo,Au) porphyry systems. In order to quantify the hydrothermal mobility of Pt we performed an investigation of the speciation of Pt in hydrothermal chloridebearing fluids and dry melt by means of X-ray absorption spectroscopy (XAS). The experiments consisted in recording the Pt L_3 -edge X-ray absorption near edge structure/extended X-ray absorption fine structure (XANES/EXAFS) spectra of Pt-bearing fluids obtained by dissolution of Pt metal in KCl/HCl and CsCl/HCl fluids in the temperature range from 450 to 575 °C at pressures from 0.5 to 5 kbar. A spectrum of Pt dissolved in dry CsCl/NaCl/KCl + K₂S₂O₈ melt was recorded at 650 °C. The capillary method, when the experimental solution together with Pt_(cr) is sealed inside a silica glass capillary, was used. As was determined from the XANES spectra, in all the experimental systems Pt existed in the +2 oxidation state. Analysis of EXAFS spectra showed that Pt is coordinated by four Cl atoms with $R_{Pt-Cl} = 2.31\pm0.01$ Å independently of the *T*-*P*-compositional parameters. No evidence of the formation of complex with alkali metal cations in the second coordination sphere of Pt was found by the analysis of the EXAFS spectra of concentrated CsCl brines and melt. Our results imply that $PtCl_4^{2-}$ is the main Pt-Cl complex which predominates in hydrothermal fluids at t > 400 °C and fluid density d > 0.3 g·cm⁻³. Experimental data obtained for dry melt of alkali metal chlorides suggest that Pt-Cl complexes can dominate Pt speciation in chloride-bearing aluminosilicate melts where Cl exhibits a salt-like atomic arrangement and ionic bonding. The literature data on the Pt solubility constant, $Pt_{(cr)} + 2$ $HCl_{(aq)}^{\circ} + 2 Cl^{-} = PtCl_{4}^{2-} + H_{2}^{\circ}_{(aq)}$, are compiled and fitted to the simple density model equation $\log K_{s}^{\circ}(\text{PtCl}_{4}^{2-})=0.973-8202 \cdot T(\text{K})^{-1}-5.505 \cdot \log d(\text{w})+2223 \cdot (\log d(\text{w})) \cdot T(\text{K})^{-1})$, where d(w) is the pure water density in $g \cdot cm^{-3}$. The equation, combined with the extended Debye-Hückel equation for activity coefficients, can be used to calculate the solubility of Pt up to 1000 °C/5 kbar. It accurately predicts the solubility of Pt in concentrated chloride brine (up to 50 wt% NaCl) at parameters of magmatic-hydrothermal transition (800 °C/1.4 kbar). At fluid/vapor density below 0.3 g·cm⁻³ a neutral complex PtCl₂ $^{\circ}$ _(aq) is suggested as the dominant Pt species. Our data

demonstrate that Pt is highly mobile in high-temperature oxidized chloride-rich hydrothermal fluids. For example, at 800 °C/2 kbar the concentration of Pt can reach a few wt.% in the 1 wt% HCl/50 wt% NaCl fluid which is in equilibrium with magnetite-hematite buffer. Once a Cl-reach fluid exsolves from alumuinosilicate melt, Pt follows Cl and enriches the fluid phase where it exists mostly in the form of $PtCl_4^{2^-}$. Decrease of temperature, acidity, and fluid chlorinity results in precipitation of Pt from the fluid phase.

Keywords: platinum, hydrothermal fluids, solubility, chloride complexes, chloride melts, X-ray absorption spectroscopy

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1. Introduction

The majority of sulfide deposits of Pt and other platinum-group elements (PGEs) are related to magmatic systems - plutonic and volcanoplutonic complexes among which the layered intrusions host the largest reserves of these metals (e.g. Bushveld Complex, South Africa; Stillwater Complex, Montana, USA; deposits of Norilsk-Talnakh district, Russia). The role of hydrothermal fluids in the formation of these deposits is still debatable. Apart from magmatic hypothesis that considers formation of these deposits as a consequence of the progressive crystallization of one or more magmas or their mixtures, there is another view on the formation of the PGE deposits in the layered intrusions where important role in the concentration of PGEs is assigned to hydrothermal fluids. The possible role of these late mineralizing fluids consists in the formation of the PGEs-bearing reefs as a result of alteration and remelting of the original magmatic mineral assemblages (e.g. Boudreau, 2016; Boudreau, 2017). In all these deposits, hydrothermal fluids were chloride-bearing with the dominant role of NaCl. For example, at the Stillwater Complex the concentration of chloride salts ranged from NaCl-dominated halide melts (>82 wt% NaCl eq., t = 660-800 °C) to more complex Na-Ca-K-Fe-Mn-Ba brines of lower salinity (28-79 wt% NaCl eq., t = 480-640 °C, Hanley et al., 2008). At Merensky reef of the Bushveld Complex salinity of chloride-rich hydrothermal fluids varied from high (60-70 wt% NaCl eq.) to moderate-to-low (20 to ~7 wt% NaCl eq.) depending on the entrapment temperature which was in the range of 750-400 °C (Ballhaus and Strumpfl, 1986; Zhitova et al., 2016). Abundance of H₂O-, Cl-, and F- bearing minerals crystallized in Norilsk ores in the vicinity of the Au-Ag-PGE chalcogenides, chlorides, and intermetallic compounds shows that the ore mineralization was related to post-magmatic hydrothermal stage at temperature <350 °C (Sluzhenikin and Distler, 2015). The PGEs are often extracted as by-products in porphyry Cu(-Mo,Au) deposits where their concentration in sulfide minerals can reach a few hundred ppb level (c.f. Plotinskaya et al., 2018 and references therein). In these deposits oxidized high-temperature (>350 °C) high-salinity (50 wt% and higher) fluids are considered as the transport medium of the PGEs.

Existing experimental data on the state of PGEs in hydrothermal fluids (with emphasis on Pd) implies that Cl is the most important complexing ligand for these elements (e.g., Xiong and Wood, 2000; Boily and Seward, 2005; Boily et al., 2007; Tagirov et al., 2013, Bazarkina and Pokrovski, 2014; Mei et al, 2015). There is a number of experimental determinations of the stability of Pt-Cl complexes published in the literature, but these data are controversial, especially at hydrothermal T-P parameters. Gammons et al. (1992) and Tagirov et al. (2015) calculated the PtCl₃ stability constant using experimental data on the solubility of Pt. In these studies $PtCl_3^-$ was considered as the dominant Pt complex in chloride-bearing fluids at $t \ge 300$ °C. However, as we noted in Tagirov et al. (2015), our experimental data can be affected by interaction of the dissolved metal with the autoclave walls yielding overestimated Pt solubility. To circumvent this we performed a study of the coupled solubility of Au and Pt in chloridebearing fluids (Zotov et al., 2017). The solubility of the metals was best described by $AuCl_2^-$ and PtCl₄²⁻ complexes. These results are consistent with our recent X-ray absorption (XAS) study of the speciation of Pt in chloride-bearing solutions (Tagirov et al., 2017) where it was demonstrated that $PtCl_4^{2-}$ predominates at 20-350 °C/P_{sat.} at chlorinity from 0.2m to 4m. Experimental studies of the solubility of Pt in chloride-rich brines and coexisting vapors at the parameters of magmatic-hydrothermal transition also indicate the possibility of the formation of stable Pt-Cl complexes (c.f. Simon and Pettke, 2009). In contrast, Scholten et al. (2018), based on the results of XAS experiments performed at 400-500 °C/800 bar concluded that Pt is practically immobile in Cl-bearing fluids at HCl concentration up to 6.9m. Therefore, more experimental data are necessary to provide a basis for reliable thermodynamic model of Pt-Cl complexing in hydrothermal fluids.

The aim of this work is to study the impact of *T-P*-compositional parameters on the Pt speciation by means of synchrotron-based XAS, to combine these data with reliable literature values of the Pt solubility constant, and to quantify the hydrothermal Pt mobility in chloridebearing fluids at *T-P* parameters from ambient to those of magmatic-hydrothermal transition. The Pt L_3 -edge X-ray absorption near edge structure/extended X-ray absorption fine structure (XANES/EXAFS) spectra of HCl/KCl and HCl/CsCl fluids (up to 11*m* of total chloride

concentration) were recorded in situ in the temperature range from 450 to 575 °C at pressures of 500-5300 bar. The X-ray absorption spectrum of Pt in dry CsCl/NaCl/KCl melt was acquired at 650 °C. Combination of these data with Pt solubility constant calculated using literature data for 25-450 °C made it possible to build a model which quantitatively describes the solubility of Pt in Cl-bearing fluids in the wide range of temperatures and pressures – from 25 °C/*P*_{sat} to 1000 °C/5000 bar and at fluid salinities at least up to 50 wt% NaCl eq. The model was tested via processing experimental data of Simon and Pettke (2009) on Pt solubility in chloride-rich brine and vapor at 800 °C/1400 bar, and Scholten et al. (2018) on the solubility of Pt in HCl-rich fluid. Both sets of data are in fair agreement with our model. These results provided a basis for discussion of the mechanisms of hydrothermal Pt transport and deposition, and for the estimation of Pt speciation in Cl-bearing aluminosilicate melts,

2. Methods

2.1. X-ray absorption spectroscopy

The Pt L_3 -edge X-ray absorption spectra were recorded at the Rossendorf Beamline BM20 (ROBL) of the European Synchrotron Radiation Facility (ESRF) in Grenoble, France. The storage-ring operating conditions were 6.0 GeV and 80-100 mA. The photon energy was scanned from 11310 to 12320 eV using the Si(111) monochromator coupled to Rh-coated mirrors for the collimation and reduction of higher harmonics. Energy calibration was performed using the L_3 -edge excitation energy of Pt metal foil (11564 eV). The spectra were collected using 13-element high-throughput Ge-detector. The total energy resolution (incident energy and core – hole lifetime broadening) has been evaluated as 8.8 eV. The detected intensity was normalized to the incident photon flux.

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The experimental set-up used for the high-temperature in situ XAS experiments is described in detail in Trigub et al. (2017a,b). A small piece of Pt wire (1.2-1.5 mm long) was loaded into a Polymicro TechnologiesTM silica glass capillary: 600 μ m OD, 250 μ m ID, 12 mm length. The capillaries were filled with experimental solutions and hermetically sealed (the capillaries were water-cooled during the sealing). The redox state of the system was controlled

either by sulfite/sulfate equilibrium (experimental solution contained Na₂SO₃/H₂SO₄), or by oxygen produced due to the thermal decomposition of $K_2S_2O_8$ ($K_2S_2O_8 \rightarrow K_2SO_4 + SO_2 + O_2$). In the latter case a weighted amount (~0.2 g) of KCl + $K_2S_2O_8$ mixture was loaded into the capillary prior to the experimental solution. The HCl concentration was determined by volumetric titration against Trizma® base using methyl red as indicator, and concentration of H₂SO₄ - by densimetry. In case of dry chloride melt the capillaries were loaded with the eutectic mixture of CsCl/NaCl/KCl (30 at% NaCl/24.5 at% KCl with melting temperature of 478 °C) together with $K_2S_2O_8$. The volumetric ratio of the chloride mixture to $K_2S_2O_8$ was ~ 10/1. The capillaries with eutectic chloride mixture were evacuated down to a 10^{-4} bar pressure, sealed, and preconditioned at the experimental temperature during 24 hours. At the beamline a capillary was placed into the microtomography furnace (Bellet et al., 2003) between two plates of silver which eliminated temperature gradient. Temperature readings were calibrated before the experiments at an accuracy of ± 5 °C with a K-type thermocouple inserted directly in the place of the capillary. Pressure inside the capillary was estimated as following. First, the volumetric degree of filling of the capillary was determined with accuracy of $\pm 3\%$ using optical microscope. Then the mass of solution in the capillary was calculated using tabulated densities of H₂O-CsCl or H₂O-KCl solutions at 25 °C, 1 bar. Finally, the pressure inside the capillary at the experimental temperature was evaluated using *PVT* properties of H₂O-KCl system (Anderko and Pitzer, 1993). We estimate the uncertainty of the calculated pressures as $\pm 25\%$. This value is based on the uncertainty of the degree of filling of the capillaries. As the *PVT* properties of H₂O-CsCl system at high temperatures are not known, the utilization of the data of the system H₂O-KCl for calculation of experimental pressures of CsCl-bearing fluids also contributes to the overall pressure uncertainty. The spectra were recorded after thermal equilibration of the experimental system and the signal stabilization, which took 30-45 min at the experimental temperature. The parameters of XAS experiments and the experimental fluid compositions are listed in Table 1.

The EXAFS ($\chi_{exp}(k)$) data were analyzed using ARTEMIS program (a part of IFEFFIT software package, Ravel and Newville, 2005). Following standard procedures for pre-edge subtraction and spline background removal, the structural parameters - interatomic distances (R_i), coordination numbers (N_i), and Debye–Waller factors (σ^2_i) - were determined via the non-linear fit of the theoretical spectra to the experimental ones with the equation

$$\chi(k) = S_0^2 \sum_{i=1}^n \frac{N_i F_i(k)}{R_i^2 k} e^{\frac{-2R_i}{\lambda(k)}} e^{-2\sigma_i^2 k^2} \sin(2kR_i + \varphi_i(k)) \qquad (1)$$

The theoretical spectra were simulated using photoelectron mean free path length $\lambda(k)$, amplitude $F_i(k)$, and phase shift $\varphi_i(k)$ parameters calculated *ab initio* using the program FEFF6 (Zabinsky et al., 1995).

2.3. Thermodynamic calculations

The standard state of a pure solid phase and H_2O corresponds to a unit activity of the pure phase at a given temperature and pressure. The standard state adopted for the aqueous species is unit activity for a hypothetical one molal ideal solution. The activity coefficients of charged aqueous species were calculated using an extended Debye-Hückel equation

$$\log \gamma_{i} = -\frac{Az_{i}^{2}\sqrt{I}}{1+Ba\sqrt{I}} + \Gamma_{\gamma} \qquad (2)$$

where the ion size parameter a° was taken to be 4.5 Å for all species, *A* and *B* refer to the Debye-Hückel activity coefficient parameters, *I* is the ionic strength in molal scale, z_i is the charge of the species, and Γ_{γ} denotes the conversion factor of mol fraction to molality. For neutral species, it was assumed that $\log \gamma_n = \Gamma_{\gamma} = -\log (1 + 0.018 \cdot m^*)$ where m* is the sum of the concentrations of all solute species. Speciation calculations were performed by means of the Gibbs computer code of the HCh software package (Shvarov, 2008). Thermodynamic properties of Pt_(cr) and aqueous species Na⁺, Cl⁻, OH⁻, HS⁻, H₂S^o_(aq), HSO₄⁻, SO₄²⁻, HSO₃⁻, SO₃²⁻, S₂O₃²⁻, K⁺, NaOH^o_(aq), KOH^o_(aq), Cs⁺, CsCl^o_(aq), CsOH^o_(aq), and other oxidized sulfur species were taken from SUPCRT92 database (Johnson et al., 1992), for CsCl^o - from Sverjensky et al. (1997).

Thermodynamic properties of H₂O and HCl° from Wagner and Pruss (2002) and Tagirov et al. (1997), respectively; those of H₂°_(aq), O₂°_(aq), and SO₂°_(aq) - from Akinfiev and Diamond (2003), data for NaCl°_(aq) and KCl°_(aq) – from Ho et al. (1994) and Ho and Palmer (1997), respectively. The values of Henry constants, dissociation constants of aqueous electrolytes calculated using the aforementioned thermodynamic data (except data available at SUPCRT database) are presented in Electronic Annex together with corresponding values of Gibbs fee energies of aqueous species (Tables EA1-EA10). Thermodynamic properties of hematite Fe₂O₃, magnetite Fe₃O₄, Ni_(cr), and bunsenite NiO were taken from Robie and Hemingway (1995).

In order to compare the Pt solubility model obtained in the present study with experimental data on the solubility of Pt in coexisting vapor-brine-rhyolite melt system (Simon and Pettke, 2009), the chemical speciation of solutes and the solubility of Pt in coexisting brine and vapor phases of the H₂O - NaCl system were calculated using the method developed in Akinfiev and Diamond (2009) for computation of SiO₂ solubility in H₂O - salt systems. In this method an *effective* H₂O molar volume $V_{H_2O}^*$ (which corresponds to *effective* pressure $P_{eff.}$) is used instead of the pure water volume $V_{H_2O}^{\circ}$. The value of $V_{H_2O}^*$ is computed for given *T-P* parameters and total NaCl content from the equation

$$V_{mix} = X_{\rm H_2O} V_{\rm H_2O}^* + X_{\rm NaCl} V_{\rm NaCl}$$
(3)

where V_{mix} is the molar volume of the H₂O - NaCl fluid (cm³·mol⁻¹), and X_{NaCl} and V_{NaCl} denote the mole fraction and the intrinsic volume of the solute, NaCl, respectively. The V_{mix} was adopted from the Driesner and Heinrich (2007) H₂O - NaCl equation of state (EoS), while for pure NaCl $V_{NaCl} = 30.8 \text{ cm}^3 \cdot \text{mol}^{-1}$ was used (Akinfiev and Diamond, 2009). In some of the experiments of Simon and Pettke (2009) the experimental system contained Ag/AgCl buffer which controlled the activity of HCl. To model the Ag-bearing system, thermodynamic properties of AgCl₂⁻ (Tagirov et al., 1997), AgCl^o_(aq) (Akinfiev and Zotov, 2016), and other Ag species (Akinfiev and Zotov, 2010) were used. Activity of Ag in Pt-Ag alloy was taken equal to Ag mole fraction a(Ag) = X(Ag) = 0.7 (Okamoto, 1997).

3. Results

3.1. Quantitative analysis of XANES spectra

The Pt L_3 -edge XANES spectra are presented in Fig. 1a. Positions of the main spectral features are given in Table 2. The spectra of Pt in experimental solutions and melt differ from the spectra of Pt(IV) references – H₂PtCl₆ and K₂PtCl₆. At the same time, position of the absorption edge (or edge jump, e.j.) and white line (the first intensive peak of the spectrum, WL) of the experimental systems are identical to the Pt(II) reference K₂PtCl₄. Therefore, the formal oxidation state of Pt is +2 in all the experimental fluids and chloride melt despite high oxygen fugacity.

3.2. EXAFS spectra fitting

The Pt L_3 -edge EXAFS spectra and their Fourier Transforms are shown in Fig. 1b and c, respectively. Results of the EXAFS spectra fitting are presented in Table 3 and shown in Fig. 2 (Experiments Cap 2160 and Cap 120) and Fig. EA1 (Experiments Cap 119, Cap 125). In all studied systems the best fits of experimental spectra were achieved for the model with four Cl atoms in the 1st coordination sphere of Pt. The interatomic distances in the 1st coordination sphere are independent of the *T-P*-compositional parameters. Attempts to replace a Cl atom with O always resulted in the oxygen-free 1st coordination sphere. Therefore, only Cl presents in the first coordination sphere of Pt, and PtCl4²⁻ is the dominant Pt complex.

In order to explore the possibility of the alkali metal atom (Me) to present in the second coordination sphere of Pt and to contribute to the EXAFS signal we performed two fits of each spectra. The first model included the alkali metal atom (left panels in Table 3 and Fig. 2, Fig. EA1), whereas the second fit was performed without the Pt-Me path (right panels in Table 3 and Fig. 2, Fig EA1). In general, inclusion of the second-coordination-sphere atom had negligible effect on the fit quality. In some cases, however, the fine structure of the EXAFS spectra in the region between 3 and 4 Å (not corrected for phase shift) was better described when the Pt-Me path was included into the fit (for example, for Cap 120 experiment). This data indicate that, even when the alkali metal cation presents in the second coordination sphere of Pt, it does not

form a stable complex of rigid geometry (as in this case its contribution would be clearly distinguishable), but has a diffuse distribution similar to the Au-Cl complexation (Mei et al., 2014; Tagirov et al., 2019). As the cation has no effect on the fit results for the 1st coordination sphere, any model presented in Table 3 can be used to characterize the nearest neighbors of Pt.

The Pt-Cl coordination numbers and distances in the 1st coordination sphere (N_{Cl} = 4.0±0.7, R_{Pt-Cl} = 2.31±0.01 Å) are similar to those described in our previous study of Pt-Cl complexing (Tagirov et al., 2017), where the local atomic environment of Pt was determined at 20-350 °C in solutions of 1.5*m* HCl, 0.2*m* HCl/0.8*m* NaCl, and 0.2*m* HCl/4*m* NaCl. Therefore, PtCl₄²⁻ predominates in the wide range of system parameters, from ambient *T-P* parameters to those of magmatic-hydrothermal transition, and at system chlorinities from ca. 1*m* of total chloride concentration to anhydrous chloride melt.

3.3. Pt solubility constant, $K_s^{\circ}(PtCl_4^{2-})$, at 25 – 1000 °C, P_{sat} – 5000 bar

The speciation of Pt in chloride-bearing fluids, according to the EXAFS data, corresponds to $PtCl_4^{2^-}$. In the present section stability of this complex is evaluated using literature data on the solubility of Pt. Three sources of stability constants for $PtCl_4^{2^-}$ were used (Table 4). For low temperatures and pressures, $t \le 150$ °C, $P_{sat.}$, we adopted the Pt solubility constant

$$Pt_{(cr)} + 2 HCl_{(aq)}^{\circ} + 2 Cl^{-} = PtCl_{4}^{2-} + H_{2(aq)} \qquad K_{s}(PtCl_{4}^{2-})$$
(4)

reported in compilation of Tagirov et al. (2015). This compilation is based on the potentiometric studies of $PtCl_4^{2-}$ stability at 25 and 60 °C/ $P_{sat.}$ performed by Ginstrup (1972), and at 25-150 °C/ $P_{sat.}$ by Nikolaeva and Erenburg (1977). The partial molal volume $V^{\circ}(PtCl_4^{2-}) = 73.28$ cm³·mol⁻¹ calculated using data of Kawaizumi (1992) was included into the set of the experimental data. For elevated temperatures and pressures of 350-450 °C/500-1000 bar we used results of our recent study of the coupled solubility of Au and Pt in chloride-bearing fluids (Zotov et al., 2017). In this study it was demonstrated that at the listed above *T-P* parameters and fluid chlorinities from 0.2*m* to 1*m* the solubilities of Au and Pt are consistent with the reaction,

with reaction constant independent of the system physical-chemical parameters (pH, fluid chlorinity and redox potential),

$$\log K^{\circ}_{(\text{Pt-Au)}} = 2 \log a(\text{AuCl}_2) - \log a(\text{PtCl}_4) + \log a(\text{Pt}_{(\text{cr})}) - 2 \log a(\text{Au}_{(\text{cr})}) \quad .$$
(6)

The values of log $K^{\circ}_{(Pt-Au)}$, together with the values of AuCl₂⁻ solubility constant

$$Au_{(cr)} + HCl^{\circ}_{(aq)} + Cl^{-} = AuCl_{2}^{-} + H_{2(aq)} \qquad K_{s}(AuCl_{2}^{-}) \qquad (7)$$

from Zotov et al. (2018) were used to evaluate the Pt solubility constant $K^{\circ}_{s}(PtCl_{4}^{2^{-}})$ (Eq. 4). The values of $K^{\circ}_{s}(AuCl_{2})$ used in the calculation are given in Table EA11, and the values of Gibbs free energy (g) of the complex are listed in Table EA12. The resulting Pt solubility constants (Table 4) were fitted to a simple density model equation (Anderson et al., 1991),

$$\log K_{\rm s}^{\circ}({\rm PtCl_4}^{2-}) = 0.973 - 8202 \cdot T({\rm K})^{-1} - 5.505 \cdot \log d({\rm w}) + 2223 \cdot (\log d({\rm w})) \cdot T({\rm K})^{-1}) , (8)$$

where d(w) is the pure water density in g·cm⁻³. The calculated values of $K^{\circ}_{s}(PtCl_{4}^{2^{-}})$ are listed in Table 5, and the corresponding values of Gibbs free energy of $PtCl_{4}^{2^{-}}$ are given in Table EA13. The calculated solubility constants, $K_{s}^{\circ}(PtCl_{4}^{2^{-}})$, are plotted in Fig. 3 together with the literature experimental data. An important feature of the Pt solubility constant (log $K_{s}(PtCl_{4}^{2^{-}})$) is its proximity to linear character with respect to the reciprocal temperature. Therefore, the range of the temperatures where Eq. (8) can be used to calculate the Pt solubility was extended up to 1000

4. Discussion

4.1. Speciation of Pt in hydrothermal systems and anhydrous chloride melts

Our data demonstrate that the speciation of Pt in chloride-bearing fluids corresponds to $PtCl_4^{2-}$ in wide range of *T-P*-compositional parameters of the Earth crust and the upper mantle, from subcritical temperatures to parameters of magmatic-hydrothermal transition, and from relatively diluted acidic chloride fluids of ca. 0.2*m* total chloride to dry chloride melt. In these

fluids the solubility of Pt is described by Eq. (4), or, involving the proton activity (or pH = -log $a(H^+)$),

$$Pt_{(cr)} + 2 H^{+} + 4 Cl^{-} = PtCl_{4}^{2-} + H_{2(g)}$$
(9)

or, using oxygen as a reactant which acts as an oxidizing agent,

$$Pt_{(cr)} + 2 HCl^{\circ}_{(aq)} + 2 Cl^{-} + 0.5 O_{2(g)} = PtCl_{4}^{2-} + H_{2}O$$
(10)

Similar reaction can be written for Pt dissolution in dry chloride melt,

$$Pt_{(cr)} + 4 Cl^{-} + 0.5 O_{2(g)} = PtCl_{4}^{2-} + O^{2-}$$

or, in terms of Na-bearing species,

$$Pt_{(cr)} + 2 NaCl + 2 Cl^{-} + 0.5 O_{2(g)} = PtCl_{4}^{2} + Na_{2}O$$
(12)

(11)

Reactions (4) and (9)-(12) show that the solubility (concentration) of Pt in chloride-bearing fluids, brines, and melts increases proportionally to the 2nd power of fluid acidity, the 4th power of chloride activity (or, in rough, the total chloride salt concentration), and decreases proportionally to the hydrogen fugacity (or increases proportionally to $f(O_2)^{0.5}$). The literature data on the solubility of Pt in sulfide-bearing systems imply that the other important species that can afford the transport of Pt at subcritical temperatures and low fluid chlorinity are hydrosulfide complexes (Pt(HS)₂° and Pt(HS)₃⁻, e.g. Gammons and Bloom, 1993; Kokh et al., 2017).

Identity of the speciation of Pt in wide range of Cl concentration implies that the solubility of Pt can be calculated using reaction (4) constant, $K_s(PtCl_4^{2-})$, at any set of *T-P*-compositional parameters of the natural ore-forming fluids, from diluted low-temperature solutions to concentrated magmatic-hydrothermal fluids. In such calculations the effect of dissolved chloride salts on the solubility of Pt can be explicitly accounted for by the activity coefficients calculated by means of the extended Debye-Hückel equation even in the case of concentrated brines. In low-density fluids ($d < 0.3 \text{ g} \cdot \text{cm}^{-3}$) the PtCl₂° is expected to be the dominant Pt species (see the next section). Some examples of these thermodynamic calculations are given in the following sections.

Anhydrous chloride salt melts as a reservoir which accumulates metals at the upper part of degassing magma chamber beneath volcanoes was suggested by Shmulovich and Churakov (1998) and Shmulovich et al. (2016) to explain formation of the metal-rich fumarolic products.

For example, Pd-Pt selenide and native Pt were discovered in artificial fumarolic preciptates sampled at Kudryavy volcano (Kurile Islands) (Korzhinsky et al., 1996; Yudovskaya et al., 2006). At this volcano high PGEs concentrations were established in mineralized rocks and fumarolic gas condensates (a few tens ppb to 0.5 ppm Pt, up to 8 ppm Pd, and 1 ppm Os, Distler et al., 2002). Formation of lenses of high-salinity brine is suggested in Afanasyev et al. (2018) to explain regions of high electrical conductivity beneath active and dormant volcanoes. Results of our study imply that these melts and brines can accumulate high concentrations of Pt in the form of $PtCl_4^{2-}$ (Reactions 11, 12) which is transported to the earth surface by high-temperature low-density fumarolic gases in the form of $PtCl_2^{\circ}_{(aq)}$.

4.2. Comparison of calculated Pt solubility in magmatic-hydrothermal fluids with the literature data

Simon and Pettke (2009) determined the solubility of Pt in coexisting Cl-bearing vapor, brine, and rhyolite melt at 800 °C. The hydrogen fugacity was controlled by Ni-NiO buffer. Two series of experiments were performed: at pressure of 1400 bar the HCl activity was controlled by Ag/AgCl buffer, whereas at 1000 bar pressure the HCl activity was not buffered. The experimental data reported in Simon and Pettke (2009) were used to test our thermodynamic model of Pt solubility. Comparison of the experimental and calculated Pt solubility is given in Table 6; results of the speciation calculations are presented in Tables EA14-EA17. In the system with HCl activity buffered by Ag/AgCl couple (P = 1400 bar, Calculations A and B in Table 6) the experimental Pt solubility values agree within 0.3 log units with the concentration of $PtCl_4^{2-}$ in both brine and vapor phases. Close agreement between the experiment and the model predictions for PtCl₄²⁻ argues for high accuracy of our model of Pt speciation and solubility, as well as of thermodynamic properties of Ag-Cl aqueous complexes which affect the speciation of Cl due to high solubility of liquid AgCl at the experimental T-P parameters. High NaCl concentration in dense brine results in higher Pt content compared to the low density vapor with much lower NaCl concentration (the difference is ca. 30 times at 1400 bar, Calculations A and B). In the case of unbuffered with respect to HCl brine (P = 1000 bar, Calculation C) the HCl

concentration of 2.8m (mol·(kg H₂O)⁻¹) estimated by Simon and Pettke (2009) yields the dissolved Pt concentration 160 times higher than the measured value. To get the correct value of Pt solubility in unbuffered with respect to HCl activity brine the HCl concentration of 0.22m should be used (Calculation E). Calculation F shows that the measured concentration of Pt in the low-density vapor phase is much higher than the $PtCl_4^{2-}$ concentration calculated using Eq. 8 for log $K_s^{\circ}(\text{PtCl}_4^{2-})$. This means that another Pt-Cl complex becomes dominant in the low-density phase which results in increase of dissolved Pt concentration. As the concentration of NaCl in the vapor phase is much lower than that in brine, and neutral species become much more important due to enhancement of ionic association at low densities, it is expected that the neutral $PtCl_2^{\circ}_{(aq)}$ complex predominates in the vapor phase. The value of Gibbs energy $g(PtCl_2^{\circ}_{(aq)}) = -$ 298.7 kJ·mol⁻¹ fits accurately the experimental Pt solubility in the low-density vapor (Calculation F). Assuming that the partial molal volume $V^{\circ}(PtCl_{2}^{\circ}(aq)) \sim V^{\circ}(PdCl_{2}^{\circ}(aq))$ (Table 6 in Tagirov et al., 2015) we got the equilibrium concentration in the coexisting brine $m(PtCl_2^{\circ}_{(aq)}) \sim 3.8 \cdot 10^{-7}$ mol·(kg H₂O)⁻¹ which is negligible in comparison with the measured Pt concentration of $5.0 \cdot 10^{-1}$ ${}^{5}m$ accounted for by the presence of PtCl₄²⁻. This calculation yields the Pt solubility constant for 800 °C/1000 bar,

$$Pt_{(cr)} + 2 HCl_{(aq)}^{\circ} = PtCl_{2}^{\circ}{}_{(aq)} + H_{2(aq)} \qquad \qquad K^{\circ}{}_{s}(PtCl_{2}^{\circ}{}_{(aq)}) = 10^{-6.4}$$
(13)

or, when reaction involves the gas phase H₂,

$$Pt_{(cr)} + 2 HCl^{\circ}_{(aq)} = PtCl_{2}^{\circ}_{(aq)} + H_{2(g)} \qquad K^{\circ}_{s} * (PtCl_{2}^{\circ}_{(aq)}) = 10^{-5.0}$$
(14)

Further extrapolation of $g(PtCl_2^{\circ}_{(aq)})$ to the system with Ag/AgCl buffer results in $PtCl_2^{\circ}_{(aq)}$ concentration in vapor phase commensurable with the concentration of $PtCl_4^{2-}$ (last column in Table 6, Calculation B). We note, however, that the values of $g(PtCl_2^{\circ}_{(aq)})$ and the calculated concentrations of $PtCl_2^{\circ}_{(aq)}$ are the rough estimates only. The accurate determination of the $PtCl_2^{\circ}_{(aq)}$ stability needs further experimental investigation.

Scholten et al. (2018) determined the solubility of Pt using results of X-ray fluorescence measurements. A glassy carbon cell was used in the experiments. It was determined that at 500 °C/800 bar 12.8 ppm Pt ($8 \cdot 10^{-5}m$) was dissolved in 6.86*m* HCl. At lower HCl concentration of 1*m*, as well as in equilibrium with sperrylite (PtAs₂), the Pt concentration was below the

detection limit. To model the Pt solubility experiment of Scholten et al. (2018) we performed two calculations. The *T-P* parameters were set as 500 °C/1000 bar. In the first calculation the system was unbuffered with respect to oxygen fugacity, whereas in the second one the $f(O_2)$ was controlled by reactions of dissolution of graphite with the formation of $CO_{(aq)}$, $CO_{2(aq)}$, and $CH_{4(aq)}$ (thermodynamic properties of all these substances were taken from SUPCRT92 database). The calculated $PtCl_4^{2-}$ concentration in $f(O_2)$ unbuffered system was $6 \cdot 10^{-5}m$ which is in excellent agreement with results of Scholten et al. (2018) ($8 \cdot 10^{-5}m$). In equilibrium with graphite the $PtCl_4^{2-}$ concentration decreased by 3 orders of magnitude due to decrease of the redox potential. Therefore, we can conclude that the equilibrium with graphite was not attained in the experiments of Scholten et al. (2018), and the measured solubility of Pt was accounted for by the formation of $PtCl_4^{2-}$. Experiments of Scholten et al. (2018) were performed in the salt-free system where the solubility of Pt is low because $HCP_{(aq)}$ is the dominant form of chloride (in 6.9 *m* HCl at 500 °C/1000 bar the concentration of free Cl⁻ ion is 0.1*m*). In the presence of chloride salts the concentration of Pt increases sharply (Eqs. 9, 10).

4.3. The solubility of Pt in chloride-bearing fluids and Au/Pt ratio

The solubility of Pt is shown in Fig. 4a in terms of the concentrations of individual complexes. As follows from the reaction (9) stoichiometry, the concentration of $PtCl_4^{2-}$ increases proportionally to the second power of fluid acidity (at constant chloride concentration). Therefore, the slope of the lines corresponding to the concentration of $PtCl_4^{2-}$ vs. pH in Fig. 4a is close to -2. The concentration of the Pt-Cl complex increases sharply with increasing temperature. For example, at pH = 2 the concentration of $PtCl_4^{2-}$ increases by 11 orders of magnitude with increase of the temperature from 200 to 400 °C and reaches 0.01 ppb in 1*m* NaCl (6 wt%) in equilibrium with Ni-NiO redox buffer. The concentration of Pt-HS complexes, in contrast, depends weakly on the temperature. Experimental data available on the solubility of $PtS_{(cr)}$ (cooperite) in hydrothermal fluids are consistent with the presence of two Pt-HS complexes: $Pt(HS)_2^{\circ}$ and $Pt(HS)_3^{-}$. The concentration of pH, whereas the concentration of the latter

has a maximum at near-neutral pH with pH = pK_{H_2S} . Stability of Pt(HS)₃⁻ decreases with increasing temperature. Therefore, the maximum on the solubility curve is clearly distinguishable only at 200 °C and disappears at 300 °C. At the concentration of reduced sulfur of 0.01*m* (320 ppm) and temperature between 200 and 450 °C the concentration of Pt-HS complexes varies between 10 ppt and 0.5 ppb. As the effect of temperature on the stability constant is much more pronounced for PtCl₄²⁻, at *t* > 400 °C Pt-HS complexes in acidic fluids become negligible compared to PtCl₄²⁻. As follows from Fig. 4a, the concentration of Pt at parameters of magmatic-hydrothermal transition (*t* > 600 °C) can be very high even in relatively diluted fluids with reduced redox state imposed by the Ni-NiO buffer. For example, at 800 °C in near-neutral fluids the concentration of PtCl₄²⁻ is 1 ppm, and decrease of pH by one log unit yields 10² fold increase of the Pt solubility. Accordingly, decrease of temperature, redox potential, and acid neutralization result in precipitation of dissolved Pt. Besides, precipitation of Pt can result from the increase of activity of reduced S and other chalcogens (Se, Te) and pnictogens (As, Sb, Bi) which form stable solid phases with Pt.

Dependence of the solubility of Pt on the fluid acidity, concentration of NaCl and redox state is illustrated in Fig. 4b for parameters of magmatic-hydrothermal transition (800 °C, 2 kbar). The concentration of Pt in fluid which contains 50 wt% NaCl and 1 wt% HCl increases from 150 ppm to 2.5 wt% with $f(O_2)$ change from reducing (Ni-NiO) to oxidizing (Fe₃O₄-Fe₂O₃) conditions.

The effect of the fluid chlorinity, redox state and temperature on the solubilities of Au and Pt are compared in Fig. 5a,b. The slope of the solubility curves as a function of fluid chlorinity is controlled by the composition of the dominant complexes: AuOH°/AuCl₂⁻ and PtCl₄²⁻. The extended flat part of Au solubility curve is explained by predominance region of AuOH°. Since there is no data on the stability of Pt-OH complex, we did not include it in the calculation and, therefore, the Pt solubility curve starts to increase at lower fluid chlorinity. The solubility of Pt increases much more sharply with increase of NaCl concentration as the number of Cl ligands in PtCl₄²⁻ complex is twice as much as that in the AuCl₂⁻. Increase of the redox potential from Ni-NiO to Fe₃O₄-Fe₂O₃ buffer results in increase of the solubility of ca. 10^2 times

for Pt and 10 times for Au. The difference between the concentrations of Au and Pt decreases with increasing temperature, especially in concentrated chloride brines (contrast to the low-density fluids, see below). At 800 °C/2000 bar and C(NaCl) = 50 wt% the concentration of dissolved Pt is only 2.5 log units lower than the concentration of Au, whereas at C(NaCl) = 0.1 wt% the difference is 7 log units.

Figure 6 illustrates the effect of *T-P* parameters on the Au/Pt ratio in chloride-bearing fluids. The factors that decrease the Au/Pt ratio, or, in other words, lead to Pt enrichment of the fluid and deposited ores are:

(i) an increase of the temperature in the region of high to moderate fluid densities. However, in the fluids of low density (the pure water density below 0.4 g·cm⁻³, this region corresponds to t > 600 °C in Fig. 6) the dependence of the Au/Pt ratio vs. *T* inverses and the ratio increases with increasing temperature;

(ii) an increase of the NaCl concentration.

4.4. Speciation of Pt in chloride-bearing magmas

Another important consequence of the present study concerns the state of Pt in silicate melts. Experiments on the solubility of Pt in Cl-free silicate melts of different composition (Borisov and Palme, 1997) demonstrated that the dissolved Pt concentration increases proportionally to the $f(O_2)^{0.5}$,

$$Pt_{(cr)} + 0.5 O_{2(g)} = PtO_{(melt)}$$
 (15)

which means that the "formal" oxidation state of Pt in silicate melts is +2. At the same time, the silicate melts contain halogens and sulfur which are important complexing ligands for Pt and other PGEs. The positive oxidation state of Pt in Cl-free melts implies the possibility of the formation of Pt²⁺-Cl⁻ complexes in Cl-bearing systems. The concentration of Cl in magmas varies from a few ppm to 1-2 wt% depending on the melt composition and history (i.e., partial melting and the concentration of Cl at the source of melting, fractional crystallization, magma mixing and so on, see review of the magmatic halogen chemistry in Webster et al., 2018). Spectroscopic studies of the speciation of Cl in aluminosilicate glasses indicate that Cl is mostly

coordinated by network-modifying cations (Na, Ca, Mg, Ba and so on), without clear contribution of network-forming Al and Si (Webster et al., 2018). Results of XANES study of the speciation of Cl in silicate glasses reported in Evans et al. (2008) showed that "there are similarities between spectra from Ca-bearing melts and the spectra of hydrated CaCl₂·2H₂O, suggesting that the Ca-Cl_x species could have a salt-like atomic arrangement and ionic bonding." Investigation of quench products of KCl-bearing carbonate-silicate melts by means of Raman and XANES spectroscopies demonstrated that Cl predominantly presents as (K_xNa_{1-x})Cl complexes with a distinct contribution of free Cl⁻ ion (Safonov et al., 2017). These data suggest that the speciation of Cl in aluminosilicate melts, at least to some extent, is similar to the state of Cl in anhydrous chloride melts and aqueous brines. Therefore, our experimental data on the speciation of Pt also can be used to constrain the Pt speciation model in Cl-bearing aluminosilicate melts.

Experimental determination of the solubility of Au in Cl- and S-bearing hydrous melts of rhyodacitic to andesitic compositions demonstrated direct correlation between the dissolved Au concentration and the concentrations of Cl and S (Botcharnikov et al., 2010). Increase of concentration of dissolved Cl to 1 wt% lead to increase of the solubility of Au by a factor of five compared to the Cl-free system. At the same time, no effect of H_2O content on the Au solubility was observed. These results lead the authors to the conclusion that Au may form complexes with Cl and S in silicate melts. Results of our study confirm this finding and extend it to the case of Pt and, probably, the other PGEs. Both Au and Pt in dry alkali metal chloride melts exist in the form of chloride complexes $AuCl_2^-$ (Tagirov et al., 2019) and $PtCl_4^{-2-}$ (this study). Comparison of our data with experimental results of Botcharnikov et al. (2010) for the solubility of Au in Cl-bearing silicate melts, and Evans et al. (2008) and Safonov et al. (2017) for the speciation of Cl in quench products of silicate melts lead to conclusion that Au and Pt chloride complexes can dominate the speciation of these metals not only in chloride-rich aqueous fluids and anhydrous chloride melts, but in chloride-bearing silicate melts as well. In this case the solubility of Pt will be controlled by activity (concentration) of Cl and the redox state (oxygen fugacity) according to

Reactions (11) and (12). As the concentration of Cl in silicate melts is low, formation of Pt-Cl complexes with lower number of ligands is possible (for example, $PtCl_3^-$ and $PtCl_2^\circ$).

Another possible important species which can dominate the speciation of Pt, other PGEs and Au in silicate melts are complexes with sulfur-bearing ligands. For example, Botcharnikov et al. (2011) found that Au can be mobilized in the sulfur-bearing silicate melts at redox conditions of sulfide-sulfate transition (when the concentrations of S^{2-} and S^{6+} are close to each other). In such a case the concentration of Au exhibits a maximum when it is complexed with S^{2-} ion or with other S-bearing ligands, including species in transitional oxidation states like S_3^- which is abundant component of hydrothermal fluids (Pokrovski and Dubessy, 2015; Barré et al., 2017) and can also present in silicate melts. Similar complexes can be formed by Pt.

At parameters of magmatic-hydrothermal transition the partition coefficient $D_{\text{Cl}}^{\text{fluid/melt}} >$ 1 (Botcharnikov et al., 2015), i.e. Cl partitions to the fluid phase. Our data imply that Pt follows Cl and enriches the fluid phase where it exists mostly in the form of PtCl_4^{2-} or PtCl_2° (the latter can dominate at fluid/vapor density $d < 0.3 \text{ g} \cdot \text{cm}^{-3}$). As the temperature decreases, the Pt-HS complexes also contribute to hydrothermal transport of Pt (Fig. 4a). However, the available thermodynamical data on the stability of Pt-HS species are insufficient to quantify their impact on the hydrothermal transport of Pt: determination of the stability of Pt-HS complexes is ongoing study in our laboratory.

5. Conclusions

In the present study we recorded Pt L_3 -edge X-ray absorption spectra of aqueous chloride-bearing fluids (7*m* HCl/4*m* CsCl and 4*m* HCl/6*m* KCl) at temperatures 450-575 °C and pressures up to 5000 bar. X-ray absorption spectrum of dry CsCl/NaCl/KCl melt was recorded at 650 °C. The capillary method, where the experimental solution together with a solid phase (Pt) is sealed inside a silica glass capillary, was used for the experiments. Interpretation of the EXAFS spectra showed that in all the experimental systems PtCl₄²⁻ predominates the Pt speciation. The distance between Pt and the nearest-neighbor Cl atoms is 2.31 ± 0.01 Å. The fact that this complex presents in high concentration in dry melt of the alkali metal chlorides indicates the

possible important contribution of Pt-Cl complexing in Pt transport by chloride-bearing aluminosilicate melts, where Cl could have a salt-like atomic arrangement and ionic bonding. Available literature data on the Pt solubility constant with the formation of PtCl₄²⁻ (25-450 °C, $P_{\rm sat}$ -1000 bar) were compiled and fitted to a simple density model equation. The equation, combined with the extended Debye-Hückel equation for activity coefficients of ionic species, can be used to predict Pt solubility in chloride-bearing fluids up to 1000 °C at pressures up to 5000 bar. Comparison of the calculated Pt solubility with the experimental data published so far demonstrates that the solubility of Pt can be accurately calculated in the wide range of fluid chlorinities including concentrated brines (50 wt% NaCl) at T-P parameters of magmatichydrothermal transition. Thermodynamic calculations performed using the obtained results showed that the formation of $PtCl_4^{2-}$ can account for efficient hydrothermal transport of Pt at t > t400 °C. For example, at temperature of 800 °C the solubility of Pt in oxidized chloride-rich fluids and brines can reach a few tens ppm to one wt% level depending on the acid concentration. At t < 400 °C, in sulfide-bearing fluids with moderate salt concentration of a few wt% NaCl eq. hydrosulfide complexes become the dominant Pt species which concentration varies from a few ppt to a few tenths of ppb.

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Table 1. Compositions of experimental solutions and equilibrium concentrations of aqueous species $(mol \cdot (kg H_2O)^{-1})$. Calculations were performed using thermodynamic data available in the literature (sources of thermodynamic data are listed in section 2.3) and calculated in the present study $(PtCl_4^{-2})$.

Capillary	cap1	19		cap120	cap125					
Solute		7.25 <i>n</i>	n HCl/4.23m Cs	sCl/		3.76m HCl/6.46m KCl/				
concentr.		0.56 <i>m</i> H	I ₂ SO ₄ /0.12 <i>m</i> N	a_2SO_3		0.35 <i>m</i>	$K_2S_2O_8$			
d, g·cm ⁻³	0.8	б		1.01	0.96					
<i>t</i> , °C/	500 °C/	575 °C/	450 °C/	525 °C/	575 °C/	450 C°/	525 °C/			
P , bar	1300 bar	2100 bar	3300 bar	4500 bar	5300 bar	500 bar	1300 bar			
		Calcul	ated concentrat	tions, mol·(kg H	$(I_2O)^{-1}$					
H^+	1.60E-01	6.46E-02	1.36E+00	8.05E-01	5.39E-01	2.91E-02	2.59E-02			
OH	1.36E-09	5.55E-09	1.09E-09	3.58E-09	7.76E-09	7.38E-09	8.71E-09			
H_2S°	2.39E-08	1.79E-07	6.04E-08	1.44E-07	3.96E-07	2.34E-07	6.04E-10			
HSO ₃ ⁻	1.44E-06	2.22E-06	2.15E-06	2.24E-06	2.22E-06	4.99E-06	3.45E-06			
HSO_4^-	7.57E-02	2.67E-02	2.22E-01	7.11E-02	2.97E-02	2.07E-01	3.62E-01			
HSO ₅ ⁻	5.83E-16	1.06E-14	1.77E-17	8.08E-16	4.78E-15	3.44E-17	4.49E-14			
S_2^-	6.04E-27	2.44E-25	1.89E-26	6.13E-26	2.99E-25	1.99E-22	3.22E-28			
$S_2O_3^{}$	2.44E-16	3.71E-16	1.43E-15	4.96E-16	3.68E-16	1.46E-12	1.67E-16			
$HS_2O_3^-$	1.79E-13	3.14E-13	3.94E-12	1.92E-12	1.62E-12	3.96E-12	1.70E-14			
$H_2S_2O_3^\circ$	1.69E-11	1.16E-11	3.22E-10	1.12E-10	6.95E-11	3.80E-10	3.46E-13			
$S_2O_4^{}$	2.15E-20	7.20E-20	4.39E-20	5.95E-20	7.82E-20	1.54E-17	6.53E-20			
HS_2O_4	3.48E-17	1.16E-16	1.74E-16	2.88E-16	4.00E-16	1.38E-16	1.50E-17			
$H_2S_2O_4^\circ$	3.02E-15	3.93E-15	1.24E-14	1.51E-14	1.57E-14	1.67E-14	2.91E-16			
$S_2O_5^{}$	1.31E-17	1.64E-17	2.85E-17	2.09E-17	1.57E-17	5.02E-15	9.72E-17			
$S_2O_5^{}$	1.72E-15	1.04E-15	3.06E-15	1.59E-15	8.02E-16	2.90E-13	3.45E-14			
$S_2 O_0^{}$	6 89E-22	2.10E-21	2.08E-23	1.77E-22	2.98E-22	2.53E-21	3 49E-19			
$S_2 = S_2^{-1}$	4 01E-32	2.05E-30	9 39E-31	1.56E-30	6.81E-30	6 72E-27	9 34E-35			
$S_{2}O_{1}^{}$	1 19E-22	9 57E-23	2.05E-21	5.99E-22	2 84F-22	1 16F-19	1.04F-22			
S	2 16E-37	1.41E-35	2.05E 21	3 20E-35	1.26E-34	1.10E 19	2 22F-41			
54 S(0)	1.85E-19	9.64E-20	2 17F-17	1.52E-18	4 22E-19	2.09F-15	6.17E-21			
S406	5 29E-15	9.04E-20 8.80E-15	2.17E-17 8 77E-15	9.14E-15	9.63E-15	2.05E-13 7.06E-12	8.61F-14			
SO	1.06E-06	1.07E-06	1.83E-06	9.14E-13	$3.17E_{-07}$	8 19E-03	1.46E-04			
504 НS ⁻	4.00L-00 6 18E 15	7.20F 14	1.03L-00	7.47E-07 4.21E-14	1 22E 13	0.17E-03	7.03E 16			
C^{1-}	6.16E-13	5.90E-01	1.74L-14	4.211-14 1 $4/1E+00$	1.22E-13	1.54E+00	8 20F-01			
	5.84E+00	5.78E+00	1.002+00 5.12E+00	5.18E+00	1.27E+00 5.32E+00	1.54E+00 1.57E+00	2.01E+00			
Na ⁺	1.13E.01	1.08F.01	1.25F 01	1.26E.01	1.26F 01	1.5712+00	2.01L+00			
	6.08E 11	1.06E-01	1.25E-01 1.24E 11	1.20E-01 2 70E 11	1.20E-01 4.33E-11	-	-			
NaOI1 NaClº	0.96E-11	1.80E-10	1.24L-11 1 12E 01	2.70E-11	4.33E-11	-	-			
K ⁺	1.2215-01	1.2712-01	1.12L-01	1.092-01	1.092-01	- 3 37E+00	- 2.51E+00			
K KOHo						3.37E+00	2.312 ± 00			
KC1°						2.00E-08	1.99E-08			
C_{s}^{+}	1 30E+00	1 /0F+00	1 28E+00	1.53E+00	1.67E±00	3.07L+00	4.J/L+00			
	1.391 ± 00	1.492 ± 00	7 20E 11	1.55 ± 00	1.071 ± 00					
CsOI1	4.20L-10	1.31L-09	7.29L-11	2.10L-10	3.321-10					
	2.73E+00	2.03E+00	2.09E+00	2.01E+00	2.47E+00	1 10E 06	2 08E 07			
Π_2	4.07E-07	2.55E-00	9.20E-08	4.70E-07 2.60E-17	1.40E-00	1.19E-00 7.17E-10	2.98E-07			
O_2	7.20E-17	5.02E-13	1.23E-19	5.00E-17	0.20E-10	/.1/E-19	4.36E-13			
\mathbf{D}_{1}	J.91E-01	0.39E-01	4.40E-UI	J.90E-01	0.30E-01	4./3E-UI	5.50E-01			
r(a)	4./4E-01	3.22E-01	3.30E-01	4./8E-01	3.19E-01	0.1/E-UI	0./0E-UI			
	2.1 1.6	2.2	5.1	2.9	2.9	4.2	5.2 2.5			
рн	1.0	1.9	0.4	0.6	0.8	3.0	2.5			
$E\Pi$	U.I 12.0	0.0	0.2	0.2	U.I	-U.I 1 <i>C F</i>	0.0			
$f(O_2)$, bar	-13.9	-12.0	-13.7	-13.2	-11.9	-10.5	-12.1			
ionic suelly	ionic strength.									

Table 2. Positions of the edge jump (e.j.), white line (WL), and the next to WL feature (B) of Pt L_3 -edge XANES spectra (as determined by IFEFFIT program). Uncertainty of the energy values is ± 0.5 eV.

Experimental system	Feature	Position, eV	
H ₂ PtCl ₆	e.j.	11565.5	
	WL	11568.0	
	В	11580.0	
K ₂ PtCl ₆	e.j.	11565.5	
	WL	11567.7	
	В	11579.7	
K ₂ PtCl ₄	e.j.	11563.5	
	WL	11566.0	
	В	11581.2	
Cap 2160, melt	e.j.	11564.0	
$t = 650 \ ^{\circ}\mathrm{C}$	WL	11566.5	
CsCl/NaCl/KCl	В	11578.5	
45.5at.%/30 at%/24.5 at%			
Cap 125	e.j.	11563.5	
$t = 450 ^{\circ}\text{C}, P = 500 \text{bar}$	WL	11566.5	
3.76m HCl/ $6.46m$ KCl + K ₂ S ₂ O ₈	В	11578.5	
Cap125	e.j.	11563.5	
$t = 525 ^{\circ}\text{C}, P = 1300 \text{ bar}$	WL	11566.5	
	В	11578.5	
Cap 120	e.j.	11563.5	
$t = 450 ^{\circ}\text{C}, P = 3300 \text{bar}$	WL	11566.5	
$7.25m \text{ HCl}/4.23m \text{ CsCl} + \text{H}_2\text{SO}_4/\text{Na}_2\text{SO}_3$	B	11578.5	
Cap 120	e.j.	11563.5	
$t = 525 ^{\circ}\text{C}, P = 4500 \text{bar}$	WL	11566.5	
	В	11578.0	
Cap 120	e.j.	11563.5	
$t = 575 ^{\circ}\text{C}, P = 5300 \text{bar}$	WL	11566.5	
	В	11578.5	
Cap 119	e.j.	11563.5	
$t = 500 ^{\circ}\text{C}, P = 1300 \text{bar}$	WL	11566.5	
$7.25m \text{ HCl/}4.23m \text{ CsCl} + \text{H}_2\text{SO}_4/\text{Na}_2\text{SO}_3$	В	11578.5	
Cap 119	e.j.	11563.5	
$t = 575 ^{\circ}\text{C}, P = 2100 \text{ bar}$	WL	11566.5	
	В	11578.5	

Table 3. Results of EXAFS spectra fitting (ARTEMIS program, $S_0^2 \sim 0.80$). Experimental spectra fits were performed in *k*-space. *Left* – Model with the second coordination sphere cation. *Right* – Model without the second coordination sphere cation.

	Model with the second coordination sphere cation						Model wit	thout the sec	cond coordinati	ion sphere	e cation
			EXAFS						EXAFS		
mo	N R, Å			Fi	t quality	mc				Fit	quality
Ate		σ^2 , Å ⁻²	$E^0 $ V	Red chi ²	Ate	N	<i>R</i> , Å	σ^2 , Å ⁻²	$E^0 \rightarrow V$	Red chi ²	
			E, ev	(R-factor)					E, ev	(R-factor)	
					CsCl/NaCl/I	KCl n	nelt				
	Cap 2160 (<i>k</i> -range $3 - 13$), $t = 650$ °C.										
Cl	3.89±0.50	2.30 ± 0.01	0.005 ± 0.001		20914.4	CI	3 85+0 47	230+0.01	0.005+0.001		10007 3
Cs	1	3.72±0.06	0.006 ± 0.006	6.3±1.5	(0.054)	MS	3.85	2.30 ± 0.01 4 61+0.06	0.003 ± 0.001 0.018+0.009	6.2 ± 1.4	(0.057)
MS	3.89	4.59±0.06	0.017 ± 0.009		(0.051)	NID.	5.05	1.01±0.00	0.010_0.009		(0.057)
$\mathbf{HCl/CsCl} \ \mathbf{aqueous} \ \mathbf{fluid} \ (7.25m \ \mathrm{HCl/4.23m} \ \mathrm{CsCl/0.56m} \ \mathrm{H_2SO_4/0.12m} \ \mathrm{Na_2SO_3})$											
	1		Car	p 119 (<i>k</i> -r	ange 3 – 11), <i>t</i>	= 500	$^{\circ}C, P = 13$	00 bar.	1	1 1	
Cl	4.01±0.62	2.32 ± 0.01	0.004 ± 0.002	9.5±1.7	9.5 \pm 1.7 8735.7 (0.060)	CI	4 01+0 58	2 32+0 01	0 004+0 002	9.3±1.5	7806 7
Cs	1	3.68±0.11	0.009 ± 0.014			(0.060) MS	4.01	4.65±0.05	0.001 ± 0.002 0.014 ± 0.008		(0.063)
MS	4.01	4.65±0.07	0.016±0.011								~ /
			Car	p 119 (<i>k</i> -r	ange $3 - 12$), <i>t</i>	= 575	°C, $P = 21$	00 bar.	1		
Cl	3.95±0.76	2.32 ± 0.02	0.005 ± 0.002	0.0.01	7201.6	Cl	3.93±0.71	2.31±0.01	0.005 ± 0.002	0 (1 0	6393.2
		3.65±0.14	0.010 ± 0.017	8.8±2.1	(0.091)	MS	3.93	4.62 ± 0.07	0.016 ± 0.011	8.6±1.9	(0.095)
INIS	5.95	4.01±0.09	0.019±0.013								

Table 3 – continued

	Cap 120 7.25 <i>m</i> HCl/4.23 <i>m</i> CsCl + H ₂ SO ₄ /Na ₂ SO ₃ (<i>k</i> -range 3 – 12), $t = 450$ °C, $P = 3300$ bar.										
Cl Cs MS	3.90±0.68 1 3.90	2.31±0.01 3.51±0.08 4.64±0.08	0.004±0.002 0.007±0.009 0.015±0.012	8.8±1.9	7031.6 (0.080)	Cl MS	3.94±0.64 3.94	2.31±0.01 4.63±0.06	0.004±0.002 0.013±0.010	8.7±1.7	6084.5 (0.083)
			Caj	120 (k-r	ange 3 – 12), t	= 525	$5 ^{\circ}\text{C}, P = 45$	00 bar.			
Cl Cs MS	3.74±0.57 1 3.74	2.31±0.01 4.12±0.05 4.67±0.09	0.003±0.001 0.002±0.004 0.020±0.015	8.3±1.7	4604.5 (0.063)	Cl MS	3.73±0.55 3.73	2.30±0.01 4.63±0.06	0.003±0.001 0.014±0.009	8.1±1.6	4268.5 (0.069)
Cap 120 (<i>k</i> -range 3 – 11.5), $t = 575$ °C, $P = 5300$ bar.											
Cl Cs MS	3.74±0.65 1 3.74	2.31±0.01 3.70±0.06 4.61±0.08	0.004±0.002 0.005±0.006 0.019±0.015	8.4±1.9	8051.9 (0.070)	Cl MS	3.68±0.62 3.68	2.31±0.01 4.63±0.07	0.004±0.002 0.016±0.012	8.2±1.8	7644.7 (0.079)
HCl/KCl aqueous fluid $(3.76m \text{ HCl/}6.46m \text{ KCl/}0.35m \text{ K}_2\text{S}_2\text{O}_8)$											
			Ca	p 125 (<i>k</i> -1	range 3 – 11), t	= 45	$0 ^{\circ}\mathrm{C}, P = 50$)0 bar.			
Cl K MS	3.64±0.46 1 3.64	2.31±0.01 3.44±0.13 4.62±0.04	0.003±0.001 0.016±0.021 0.006±0.006	8.7±1.3	5157.5 (0.033)	Cl MS	3.64±0.44 3.64	2.31±0.01 4.63±0.04	0.003±0.001 0.006±0.005	8.5±1.2	4650.2 (0.036)
			Cap	125 (k-ra	nge 3 – 11.5), t	= 52	5 °C, $P = 13$	300 bar.			
Cl K MS	3.83±0.67 1 3.83	2.30±0.01 3.85±0.07 4.63±0.06	0.005±0.002 0.003±0.009 0.012±0.009	7.4±1.8	16124.9 (0.071)	Cl MS	3.85±0.60 3.85	2.30±0.01 4.62±0.06	0.005±0.002 0.013±0.008	7.7±1.7	12965.5 (0.069)
Multi	ple scatterin	g MS = Pt-C	11-Pt-Cl2.								
	Multiple scattering MS = Pt-Cl1-Pt-Cl2.										

<i>t</i> , C	P, bar	$\log K^{\circ}{}_{\rm s}$	Source						
25	1	-26.50±0.24	а						
25	1000	-26.71	b						
50	1	-24.29±0.12	а						
100	$P_{\rm sat}$.	-20.86 ± 0.05	а						
150	$P_{\rm sat.}$	-18.41±0.07	а						
350	1000	-12.39±0.3	с						
450	500	-9.53±0.16	с						
450	1000	-9.42±0.27	с						
^a adopt	ed from Tag	irov et al. (201							
are l	are based on the experimental								
potenti	ometric resu	ilts of Nikolae	eva and						
Erenbu	rg (1977) an	d Ginstrup (19	72);						
^b calcul	ated using t	he value of V°	$(\operatorname{PtCl_4}^{2-})$						
from K	awaizumi (1	1992) and the	value of						
$\log K^{\circ}_{s}$	for 25°C, 1	bar from this ta	able;						
^c calcu	lated using	data of Zoto	v et al.						
(2017)	(2017) and Zotov et al. (2018) as								
describ	ed in section	ı 3.3.							

Table 4. The Pt solubility constants, $K^{\circ}_{s}(PtCl_{4}^{2-})$, of the reaction $Pt_{(cr)} + 2 HCl^{\circ}_{(aq)} + 2 Cl^{-} = PtCl_{4}^{2-} + H_{2(aq)}$ fitted to the density model equation.

Table 5. The Pt solubility constant, $K_s^{\circ}(PtCl_4^{2-})$, of the reaction $Pt_{(cr)} + 2 HCl_{(aq)}^{\circ} + 2 Cl^{-} = PtCl_4^{2-} + H_2^{\circ}_{(aq)}$ as a function of temperature and pressure. Calculations were performed using Eq. (8) obtained in the present study.

t °C	P, bar							
<i>l</i> , C	$P_{\rm sat.}$	500	1000	1500	2000	5000		
25	-26.54	-26.52	-26.51	-26.49	-26.48	-26.42		
100	-21.02	-21.01	-21.01	-21.01	-21.00	-20.99		
200	-16.31	-16.33	-16.34	-16.34	-16.35	-16.38		
250	-14.58	-14.61	-14.63	-14.65	-14.67	-14.72		
300	-13.10	-13.16	-13.20	-13.23	-13.25	-13.34		
350	-11.72	-11.88	-11.96	-12.01	-12.04	-12.16		
400		-10.69	-10.86	-10.94	-10.99	-11.15		
450		-9.41	-9.86	-9.98	-10.06	-10.26		
500		-8.09	-8.91	-9.11	-9.22	-9.48		
550			-8.00	-8.30	-8.45	-8.78		
600			-7.16	-7.55	-7.74	-8.15		
650			-6.38	-6.85	-7.09	-7.58		
700			-5.69	-6.21	-6.48	-7.05		
750				-5.61	-5.92	-6.57		
800				-5.06	-5.40	-6.13		
850				-4.56	-4.92	-5.72		
900				-4.09	-4.47	-5.33		
950				-3.66	-4.05	-4.97		
1000				-3.26	-3.67	-4.64		

Table 6. Experimental data on the solubility of Pt in coexisting vapor-brine-rhyolite melt system at 800 °C, 1000 and 1400 bar (Simon and Pettke, 2009), and results of thermodynamic modeling of the experiment.

Calculation	Type of	$P_{\rm eff.}^{a}$,	$d*_{\rm H_2O}^{\rm b}$,	$d_{H_2O}^{*}$, NaCl concentration		HCl concentration,	Pt concentration, $mol \cdot (kg H_2O)^{-1}$				
	inclusion	bar	g⋅cm ⁻³	Experiment ^c	Calculated ^d	$mol \cdot (kg H_2O)^{-1}$	Experiment	$\begin{array}{c} Calculated \\ PtCl_4^{2-e} \end{array}$	Calculated $PtCl_2^{\circ}{}^{(aq)}$		
P = 1400 bar	r, Ag-AgCl bu	iffer of HCl a	ctivity								
Α	Brine	3279	0.575	~ 43	50.1	2.37 ^f	$1.7 \cdot 10^{-4}$	$2.9 \cdot 10^{-4}$	$4.8 \cdot 10^{-5}$		
В	Vapor	1632	0.365	~ 9	8.5	3.13 ^f	$5.1 \cdot 10^{-6}$	$1.1 \cdot 10^{-5}$	$3.3 \cdot 10^{-5}$		
P = 1000 bar, without buffer of HCl activity											
Concentratio	ns of HCl in b	rine and vapo	r phases are a	dopted from Sin	non and Pettke	(2009)					
С	Brine	3189	0.567	~ 63	66.4	2.8 ^g	$5.0 \cdot 10^{-5}$	$8.1 \cdot 10^{-3}$	-		
D	Vapor	1034	0.239	~ 2	2.0	0.5 ^g	$1.9 \cdot 10^{-6}$	$4.6 \cdot 10^{-10}$	-		
Concentratio	n of HCl in br	ine is calculat	ed in the prese	ent study using t	he experimenta	ıl Pt solubility dat	ta from Simon	and Pettke (2009	\mathcal{P}) and \log		
$K_{\rm s}^{\circ}({\rm PtCl_4}^{2-})f$	rom the preser	1t study (Eq.8)	1	. <u> </u>							
Ε	Brine	3189	0.567	~ 63	66.4	0.22 ^h	$5.0 \cdot 10^{-5}$	$5.0 \cdot 10^{-5}$	$3.8 \cdot 10^{-7}$		
F	Vapor	1034	0.239	~ 2	2.0	0.83 ⁱ	$1.9 \cdot 10^{-6}$	$1.3 \cdot 10^{-9}$	$1.9 \cdot 10^{-6j}$		
^a effective pr	essure, see sec	tion 2.3 for ca	lculation detai	ls; ^b pure water	density which	corresponds to $P_{\rm e}$	_{ff.} , this value w	as used to calcul	ate the Pt		
solubility rea	solubility reaction constant with Eq. 8; ^c wt% NaCl eq.; ^d calculated using Driesner and Heinrich (2007) EoS for H ₂ O-NaCl binary system; ^e										
calculated using thermodynamic properties of PtCl ₄ ²⁻ obtained in the present study; ^f calculated for equilibrium with the Ag-AgCl buffer assuming											
a(Ag) = X(A)	g) =0.7 in Pt-A	Ag alloy; ^g – Si	mon and Pettk	te (2009) estima	tion; ⁿ estimate	ed using experime	ental Pt concent	tration determine	ed in brine		
given in the r	next column; ¹	calculated using	ng our estimati	on of the conce	ntration of HC	l in brine (see ^g) a	nd thermodyna	unic data for HC	'l° _(aq) ; ^J		
calculated us	ing $g^{\circ}(\text{PtCl}_{2}^{\circ}{}_{(a)}$	$_{aq}) = -298.7 \text{ kJ}$	$1 \cdot \text{mol}^{-1}$ at these	: T-P _{eff.} paramet	ters (800 °C, 10)34 bar).					





Fig. 2



Fig. 6