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Gold transport in hydrothermal chloride-bearing fluids: Insights from in situ x-ray absorption spectroscopy and ab initio molecular dynamics

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1	The chemistry of gold in chloride hydrothermal fluids and meits by A-ray absorption
2	spectroscopy
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ABSTRACT

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Chloride-rich fluids are widespread in the Earth's interior from low-temperature subsurface conditions to deep lithosphere. The concentration of chloride salts vary from diluted aqueous solutions peculiar to epithermal deposits to concentrated chloride brines with salinity up to 70 wt% NaCl eq., which coexist with low density vapor during the formation of porphyry systems, and anhydrous (dry) chloride melts at the upper part of degassing magma chamber beneath volcanoes. Here we report an investigation of the speciation of Au in hydrothermal chloride fluids and anhydrous melt by means of X-ray absorption spectroscopy (XAS). The experiments consisted in registration of Au L₃-edge X-ray absorption near edge structure/extended X-ray absorption fine structure (XANES/EXAFS) spectra of Au-bearing fluids in the temperature range from 350 to 575 °C at pressures 0.5-4.5 kbar. The experimental aqueous systems consisted of HCl-, NaCl-, KCl-, or CsCl- dominated fluid (0.02 to 7 mol·(kg H₂O)⁻¹) inside a silica glass capillary together with small piece of Au, and SO₂/SO₃ buffer or K₂S₂O₈ as a source of oxygen. Spectra of Au dissolved in dry CsCl/NaCl/KCl + K₂S₂O₈ melt were recorded at 650 °C. The measured concentrations of Au dissolved in hydrothermal fluids were in close agreement with the values calculated using thermodynamic data available for AuCl₂. High concentration of Au dissolved in dry chloride melt implies that it can be an effective Au scavenger in natural systems with low H₂O content. It was determined that Au is coordinated by two Cl atoms ($R_{\text{Au-Cl}} = 2.25 - 2.28 \text{ Å}$) and 0.2 - 0.6 alkali metal atoms (Me) in the distant coordination sphere of Au at $R_{\text{Au-Me}} = 3.3 - 4.1 \text{ Å}$. The presence of the alkali metal cations in the vicinity of Au-Cl complex was confirmed by the XANES spectra registered in High Energy Resolution Fluorescence Detection (HERFD) mode: the white line (WL) intensity decreased substantially in NaCl-dominated fluid and dry melt in comparison with HCldominated fluid. The observed decrease of the WL intensity stems from the partial compensation of the positive charge located on Au due to the presence of the distant-coordination-sphere alkali metal cations. The interatomic distances between Au and atoms localized in the first and distant coordination spheres exhibit contrasting behavior with respect to fluid pressure. Increasing pressure

results in weak contraction of the first coordination sphere of Au, whereas conjugated increase of the dielectric constant causes expansion of the distant coordination sphere. These results can be explained by the difference in the chemical bond nature. The Cl⁻ ions in the nearest coordination sphere of Au are bonded to the central atom via the strong (predominantly covalent) chemical bonding, whereas the distant-coordination-sphere cations are weakly bonded to the negatively charged AuCl₂⁻ complex mostly via the electrostatic interaction. This means that in thermodynamic calculations the dissolution of Au in natural ore-forming fluids, at least of moderate to high densities (> 0.5 g·cm⁻³), can be explicitly described by reactions which involve the formation of AuCl₂⁻, with mandatory consideration of the effect of weak outer sphere coulomb interactions via the calculation of activity coefficients.

1. INTRODUCTION

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Chlorides are the most abundant salt components of hydrothermal fluids in the Earth's crust. The concentration of NaCl, the main electrolyte salt, varies in wide limits, with the average value close to ~ 0.5 -1 m (see, for example, review of hydrothermal fluid compositions in Seward et al., 2014). In particular, in hydrothermal fluids which formed the volcanogenic massive sulfide (VMS) deposits the chloride concentration ranged from few to 20 wt% NaCl eq. (e.g., Large, 1977; Simonov et al., 2006; Vikent'ev et al., 2009, 2012; Karpukhina et al., 2013). Similar concentrations of chloride salts, with the average value of 4-8 wt% NaCl eq. and maximum concentration up to 26 wt% NaCl eq., were established in hydrothermal vent fluids of black smokers – modern analogues of VMS deposits (Douville et al., 2002; Bortnikov et al., 2004, 2014; Schmidt et al., 2011). Chlorinity of the high-temperature ($t\sim600-800$ °C) metal-rich fluid rising from a magma chamber in porphyry systems is about 10 wt% NaCl eq. Lowering of the temperature and pressure induces phase separation that results in formation of a low density fluid with salinity of a few wt% NaCl eq. and a brine with salinity of 30-70 wt% NaCl eq. (Heinrich et al., 2004; Sillitoe, 2010; Gregory, 2017). Note that many VMS and porphyry systems belong to the World-class gold deposits with Au resources of more than 100 t. Gold alloys with platinum group elements (PGE: Pt, Pd, Ir, and Rh) are frequent in giant PGE deposits of magmatic origin - deposits of Norilsk, Bushveld, and Stillwater. In all these deposits hydrothermal fluids participated in the formation of PGE-Au mineralization. In the Stillwater Complex the concentration of chloride salts ranged from NaCldominated halide melts (>82 wt% NaCl eq., $t = 660-800^{\circ}$ 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). Chloride-rich fluids discovered as fluid inclusions in quartz associated with sulfide minerals at Merensky reef (Bushveld complex) were entrapped in the temperature range 750-400 °C. Their salinity varies from high (60-70 wt% NaCl eq.) to moderate-to-low (20 to ~7 wt% NaCl eq.) depending on the entrapment temperature (Ballhaus and Strumpfl, 1986; Zhitova et al., 2016). Abundance of H₂O-, Cl-, and F- bearing minerals crystallized in Norilsk ores in the vicinity of Au chalcogenides, chlorides, and intermetallic compounds suggested that the Au mineralization was related to the post-magmatic hydrothermal stage which took place at temperature of ~350 °C and below (Sluzhenikin and Distler, 2015).

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Native gold crystals and particles are often formed during volcanic eruptions and fumarolic activity. Crystals of native gold were found in products of volcanic eruptions which were formed during 1975-1976 and 2012-2013 eruptions of Tolbachik volcano, Kamchatka (see Chaplygin et al., 2015, and references cited). Micron size native gold particles were discovered on the surface of sulfide minerals (wurtzite-greenockite, (Zn,Cd)S) formed in active fumaroles of Kudriavy volcano, Iturup Island (Chaplygin et al., 2007). High-temperature fumarole gases at these volcanoes, contained both Au (few ppb to few tens ppb) and chloride (as high as 1.2 mol% HCl at Tolbachik volcano, Chaplygin et al., 2015). Association of Au-bearing phases with oxychlorides and chlorides observed in tube precipitates collected at Kudriavy volcano also suggests elevated Au and Cl concentrations in the gas phase (Yudovskaya et al., 2006). Almost dry (anhydrous) chloride salt melt as a reservoir which accumulates metals at the upper part of degassing magma chamber beneath a volcano was suggested by Shmulovich and Churakov (1998) to explain the formation of the metal-rich gas sublimates. High capacity of chloride melts to accumulate Au is demonstrated by their effective usage as a salt flux for the synthesis of crystals (cf. Chareev et al., 2016) and is confirmed in the experiments on the dissolution of Au in (Na,K)Cl melts (Shmulovich et al., 2016). The speciation of Au in melted chloride salts, as well as the dissolution mechanism is, however, unknown.

The chemistry of Au in chloride systems has been experimentally studied via measurements of Au solubility at 300-600 °C and pressure up to 1800 bar (e.g., Zotov and Baranova, 1989; Zotov et al., 1989, 1991, 1995, 2017; Archibald et al., 2001; Stefánsson and Seward, 2003; Guo et al., 2018, and references therein). It was determined that in chloride-bearing acidic fluids Au can be effectively transported in the form of Au-Cl complexes with predominant role of AuCl₂-, even though some inconsistencies still remain regarding the complex stability. The Au speciation model

was confirmed by X-ray absorption spectroscopy (XAS) (Pokrovski et al., 2009). Mei et al. (2014) studied the complexing of Au with chloride in NaCl-rich aqueous fluids by means of ab initio molecular dynamics (AIMD) simulations performed at conditions from ambient to 1000 °C, 1500 bar. It was determined that in concentrated NaCl solutions AuCl₂ is associated with Na⁺ ion which is weakly bonded in the outer coordination sphere. The authors proceeded with the calculation of thermodynamic properties of NaAuCl2°(aq) complex and found that its stability rapidly increases with decrease of the hydrothermal fluid density. These data imply that neutral ionic associates predominate at relatively low densities ($d < 0.7 \text{ g/cm}^3$), whereas charged complexes dominate the Au speciation in higher density fluids. At the same time, the dissolved Au concentration measured in the cited above Au solubility studies in the wide range of NaCl/KCl concentrations and fluid densities, including d < 0.7 g/cm³, can be accurately described without the formation of $Na(K)AuCl_2^{\circ}{}_{(aq)}$. In the works of Frank et al. (2002) and Zajacz et al. (2010) the solubility of Au was measured in high-temperature fluids of contrast densities (dense brines with total chloride content of 70 wt% at 800°C, and low density fluids at 1000 °C/1500 bar, respectively). Positive correlations between HCl and NaCl/KCl concentrations with the solubility of Au was observed and interpreted in terms of the formation of HAuCl₂° (Frank et al., 2002) and NaAuCl₂° (Zajacz et al., 2010). However, complex system compositions and the absence of the reliable thermodynamic data on the system's components did not allow the authors to determine the dominant Au complex composition and calculate its stability constant unambiguously.

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The aim of this work is to quantify the impact of temperature, pressure, and the system compositions on the chemistry of Au in chloride-rich hydrothermal fluids by means of synchrotron-based XAS. Gold L_3 -edge X-ray absorption near edge structure/extended X-ray absorption fine structure (XANES/EXAFS) spectra were recorded in situ for HCl-, NaCl-, KCl-, and CsCl-dominated fluids in the temperature range from 350 to 575 °C at pressures of 0.5-4.5 kbar. The X-ray absorption spectra of dry CsCl/NaCl/KCl melt were acquired at 650 °C. Most of the experiments were performed using the Total Fluorescence Yield (TFY) detection method while the

XANES spectra of two solutions (NaCl- and HCl- dominated), and the dry chloride melt were registered in the High Energy Resolution Fluorescence Detection (HERFD) mode (Glatzel and Bergman, 2005). The presence of cation was detected in the second coordination sphere of Au for alkali metal – dominated fluids and melts. Despite the fact that geometry of the distant coordination sphere is highly disordered, we found that the outer-sphere cation affects the charge state of Au. The contrast effect of pressure was determined on interatomic distances between Au and atoms localized in its nearest and distant coordination spheres.

2. METHODS

2.1. X-ray absorption spectroscopy

X-ray absorption experiments were performed at the European Synchrotron Radiation Facility (ESRF) in Grenoble, France. The Au L_3 -edge TFY XAS (XANES/EXAFS) spectra were recorded at the Rossendorf Beamline BM20 (ROBL). The storage-ring operating conditions were 6.0 GeV and 80-100 mA. The photon energy was scanned from 11620 to 12670 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 Au metal foil (11919 eV). The TFY XAS 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.

The Au L_3 -edge HERFD-XANES spectra were collected at the high-brilliance X-ray absorption/X-ray emission spectroscopy undulator beamline ID26 (Gauthier et al. 1998). The ring current was varied between 150 and 200 mA. The incident energy was selected using the <111> reflection from a double Si crystal monochromator. Rejection of higher harmonics was achieved by three Pd mirrors positioned at an angle of 2.5 mrad relative to the incident beam. The incident X-ray beam had a flux of approximately $2 \cdot 10^{13}$ photons s⁻¹ on the sample position. The spectra were measured in the HERFD mode using an X-ray emission spectrometer (Glatzel and Bergman 2005;

Kvashnina and Scheinost 2016). The sample, analyzer crystal and photon detector (silicon drift diode) were arranged in a vertical Rowland geometry. The Au L_3 -edge HERFD-XAS spectra were obtained by recording the intensity of the Au $L_{\alpha 1}$ emission line (9713 eV) as a function of the incident energy. The emission energy was selected using the <555> reflection of four spherically bent Ge crystal analyzers (1 m curvature radius) aligned at 78° Bragg angle. A combined (incident convoluted with emitted) energy resolution of 1.5 eV was determined as the full width at half maximum of the elastic peak. The intensity was normalized to the incident 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) and will be only briefly outlined here. The heating experiments were performed using a capillary technique initially developed for Raman spectroscopic studies (Chou et al., 2008, 2012; Ding, 2010). A small piece of Au wire (1.2-1.5 mm long) was loaded into a Polymicro TechnologiesTM silica glass capillary: 600 μm OD, 250 μm ID, 12 mm length (BM20 beamline), or 600 µm OD, 400 µm ID, 15 mm length (ID26 beamline). 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 (a solution with known concentration of SO₂/H₂SO₄ or Na₂SO₃/H₂SO₄ was loaded into the capillary), 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 (~0.02-0.2 g) of KCl + K₂S₂O₈ mixture was loaded into the capillary prior to an experimental solution. The HCl concentration was determined by volumetric titration against Trizma® base using methyl red as indicator, the concentration of SO₂ - by iodometric titration, and the 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 (Fullam, 1971) together with K₂S₂O₈. 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 (BM20 beamline) or stainless steel (ID26 beamline) 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 from the degree of filling of the capillary using *PVT* properties of H₂O-NaCl system (Driesner, 2007; Driesner and Heinrich, 2007) or, in case of HCldominated fluids, H₂O-HCl (Bach et al., 1977). We estimate the uncertainty of the calculated pressures as $\pm 10\%$ for HCl- and NaCl-dominated fluids, and $\pm 25\%$ for KCl- and CsCl- dominated ones. The spectra were recorded after thermal equilibration of the experimental system and the signal stabilization. Usually ca. 30-45 min at the experimental temperature was necessary to attain the stationary state. The parameters of XAS experiments and the experimental system compositions are listed in Table 1.

2.2. EXAFS spectra fitting and wavelet transforms

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

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

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).

Wavelet transform (WT) analysis of the EXAFS spectra was performed as described in Penfold et al. (2013), Timoshenko and Kuzmin (2009), Funke et al. (2007). The WT consists in replacement of the infinitely expanded periodic function in Fourier transform by a local function -

the wavelet. The WT allows analyzing the experimental EXAFS spectra in two-dimensional k/R space and enables separation of the contributions from atoms with different Z. The WT of a given signal $\chi(k)$ is defined as

$$W_f^{\psi}(a,k') = \frac{1}{\sqrt{a}} \int_{-\infty}^{+\infty} \chi(k) \psi^* \left(\frac{k-k'}{a}\right) dk \qquad , \qquad (2)$$

where the scalar product of the EXAFS signal and the complex conjugate of the wavelet (ψ^*) is calculated as a function of a and k'. The a parameter is related with R as $a = \frac{\eta}{2R}$, and k' conforms to localization of the wavelet function in k-space. In this work we used the WT based on the Morlet wavelet functions

$$\psi(k) = \frac{1}{\sqrt{2\pi}\sigma} e^{i\eta\eta} e^{-k^2/2\sigma^2} \qquad , \tag{3}$$

where parameters σ and η are width and frequency of the wavelet function, respectively. These parameters should be adjusted to get appropriate resolution in k- and R-space. For providing better quality of WT images we used modified WT functions (Timoshenko and Kuzmin, 2009).

2.3. Ab initio molecular dynamics (AIMD) simulations

The AIMD simulations are aimed to facilitate the interpretation and modeling of XAS results. Details of the AIMD simulations are described in Trigub et al. (2017b). Born–Oppenheimer AIMD simulations were performed with the aid of CP2K program package (VandeVondele et al., 2005). A hybrid Gaussian plane-wave method (GPW) (Lippert et al., 1997) was used. The Goedecker–Teter–Hutter (GTH) norm-conserving pseudo-potentials (Goedecker et al., 1996; Hartwigsen et al., 1998; Krack, 2005) were applied to replace core electrons. The electronic density cut off was set at 500 Ry. For all the atoms the Double-zeta valence plus polarization (DZVP) basis set optimized for molecular geometries was used. The spin polarized general gradient approximation by Perdew–Burke–Enzerhof (PBE) (Perdew et al., 1996) augmented with a correction of the van der Waals (vdW) interactions was applied as exchange correlation functional,

the nonlocal rVV10 vdW functional was used (Vydrov et al., 2009; Sabatini et al., 2013). The b parameter which determines the nonlocal behavior of the functional was set to 9.3 according to Miceli et al. (2015). The simulations were conducted in the canonical NVT ensemble, with the simulation time step of 0.5 fs and periodic boundary conditions. The AIMD trajectory was accumulated for 20 ps and 10 ps for solution and melt, respectively. Each AIMD calculation was preceded by equilibration phase of 5 ps. The temperature was controlled via a Nosé–Hoover chain thermostat (Nosé, 1984a, 1984b). Starting configurations for AIMD calculations were generated by classical MD method with the aid of GROMACS software (Berendsen et al., 1995). The system contained $AuCl_2^{-1}$ or $Cat[AuCl_2]$ complex (Cat = H, Na, K, Cs) which geometry was optimized with Gaussian 09 suite of programs (Frisch et al., 2016) (the Møller-Plesset Perturbation Theory (MP2) was applied; solvation effects were accounted for by means of the Conductor-like Polarized Continuum Model (CPCM), see Pokrovski et al. (2009) for calculation details). Details of the chemical system compositions, box sizes, and temperatures used in the simulations are given in Table 2.

An additional calculation was performed to check the effect of basis set size on results of the AIMD calculation. Calculation H (Table 2) was identical to calculation E in terms of the atomic system parameters (system composition, box size), but was performed with triple-zeta valence basis set (TZV2P) applied for all atoms. This calculation was performed only for one system because the calculation time increased substantially (from 25 s to 40 s for one time step).

Results of AIMD calculations are presented in the form of plots of atomic densities and coordination numbers. The radial density $\rho(R)$ of surrounding atoms is related to coordination numbers via expression:

$$N(R) = \int 4\pi r^2 \rho(R) dr . (4)$$

2.4. EXAFS and XANES spectra simulations using AIMD data

Every 50 fs of AIMD simulation snapshots of atomic configurations were extracted and used to calculate Au L_3 -edge EXAFS/XANES spectra (hereafter the AIMD-EXAFS/AIMD-XANES notation will be used for the calculated spectra). The configuration-averaged Au L_3 -edge EXAFS spectra were calculated using FEFF8.5L program (Ankudinov et al., 1998). The Au-Cl distances were corrected by a factor of 0.97 to account for differences between experimental (EXAFS) and theoretical (AIMD) radii of the first coordination sphere. The multiple scattering (MS) contributions to EXAFS were accounted up to the fifth order within a cluster of 6 Å radius around the absorbing atom. The calculation of the cluster potential was performed for every geometry configuration. The self-consistent complex exchange-correlation Hedin–Lundqvist potential and default values of mulfin-tin radii as provided within the FEFF8.5L code were applied. The amplitude of the theoretical EXAFS spectrum was multiplied by the $S_0^2 = 0.84$ factor as determined from the Au₂S EXAFS spectra fitting.

Theoretical calculations of the Au L_3 -edge HERFD-XANES spectra were performed using the finite difference method (FDM) by means of FDMNES program (Joly, 2001; Guda et al., 2015). Relativistic self-consistent field (SCF) FDMNES calculations were carried out with the exchange-correlation part of potential in a local density approximation (Hedin and Lundqvist, 1971). The final electronic states were calculated in a full core hole screening. Atomic clusters inside the spheres with radii of 6 Å were chosen for both SCF and FDM XANES calculations. Many-body effects and core-hole lifetime broadening were accounted for by means of arctangent convolution (Bunău and Joly, 2009).

2.5. Bader charges calculation

The atomic clusters obtained by means of AIMD simulations were used to study the charge state of Au. First, the electron density of each cluster was calculated by means of Quantum Espresso software package (Giannozzi et al., 2009). At this stage we applied a projector-augmented wave (PAW) method with norm-conserving pseudopotentials and the Perdew-Burke-Ernzerhof (PBE)

exchange-correlation functional. The SCF method was applied with a 100 Ry kinetic energy cutoff for the plane waves and an SCF tolerance better than 10^{-9} for the electronic structure calculations.

Atomic charges were determined according to quantum theory of atoms in molecules (QTAIM, Bader, 1990, 1991). The local atomic charges of Au were calculated by integrating the charge density within a Bader volume around the atom using Critic2 software (Otero-de-la-Roza et al., 2009, 2014).

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3. RESULTS AND DISCUSSION

3.1. Quantitative analysis of XANES spectra and dissolved Au concentrations

Gold L_3 -edge HERFD-XANES spectra are shown in Fig. 1, and the TFY-XANES spectra are presented in Fig. 2a,b. Positions of the main spectral features are given in Table 3. In all considered systems the oxidation state of Au is +1 despite high oxygen fugacity, especially in the experiment with chloride melt. The HERFD-XANES spectra of chloride melt, NaCl- and HCl-rich solutions differ in the height of the white line (WL, the first, most intense feature). The most intense WL was observed in experiment cap28 with HCl-dominated fluid (3m HCl/0.02m NaCl, Fig. 1). The WL's intensity of NaCl-rich fluid in cap30 experiment (3m NaCl) and dry chloride melt (cap5407) are considerably lower. The Au L_3 -edge absorption is related to the 2p-5d dipole-allowed transitions: $2p_{3/2} \rightarrow 5d_{5/2}/5d_{3/2}$. Therefore, the WL intensity reflects the number of empty states (holes) in $5d_{5/2}$ and $5d_{3/2}$ orbitals above the Fermi level. Low WL intensity of NaCl-rich fluid and chloride melt indicates the reduced unoccupied density of states (DOS) of Au 5d orbitals, or, in other words, larger number of electrons localized at Au 5d orbitals for salt-rich systems (cap30 and cap5407) than in the HCl-dominated system (cap28). This fact can be quantitatively explained by interaction of positively charged ions with AuCl₂ complex in concentrated salt fluid and melt. The presence of positively charged ions in the vicinity of AuCl₂ complex results in electron transfer from ligands to metal, which gives rise to decrease of the number of holes in Au 5d orbitals (or decrease of the positive charge localized on Au) and, therefore, decrease of the WL intensity. The

decrease of the positive charge is confirmed by the Bader partial atomic charges calculation which is discussed below (section 3.5).

The WL intensities of the TFY XANES spectra of hydrothermal fluids (cap122, cap118, cap117, cap110), all of which were recorded for salt-rich systems, are identical (Fig. 2a). At the same time, the WL of the TFY XANES spectrum of salt melt (cap2156) is more intense than WL of K- and Cs-bearing aqueous fluids. This fact can be explained by more effective compensation of Au positive charge in aqueous fluids which contain high concentrations of heavy alkaline metal cations in comparison with chloride salt melt. Besides, according to the AIMD calculations, the positive charge on Au in chloride melt increases due to the presence of CI^- ions in the distant coordination sphere at 2.6 - 3 Å, whereas no CI^- ions were detected in the second coordination sphere of Au in aqueous fluids (see section 3.3). We note, however, that the effect of fluid composition on the electronic density distribution within the Au complex and on the Au valence state remains open.

Figure 2b compares XANES spectra of chloride melt (cap2156) recorded at high and ambient temperature. The spectra are different: the crystallized melt spectrum exhibits several additional features which are absent in the high-temperature spectra. This difference is connected with appearance of long-range atomic order in crystallized melt. The edge jump (e.j.) and WL positions of these spectra are also shifted relative to each other (Table 3).

The concentrations of dissolved Au calculated from the amplitude of the absorption edge height of the transmission spectra are listed at the bottom of Table 1. They are in close agreement with Au solubilities calculated using thermodynamic properties of $AuCl_2$ reported in Akinfiev and Zotov (2010). Good agreement between results of our experiments and calculated solubilities argues for high quality of the thermodynamic model of Akinfiev and Zotov (2010), as well as the experimental data which were used by the authors to calculate thermodynamic properties of $AuCl_2$. Note that relatively high uncertainty of Au concentrations determined in the present study, which we estimate as \pm 0.3 log units, stem from the spherical shape of the capillary, possible shift of the beam from the center of the capillary, uncertainty in the solute concentrations due to weighting of

small masses of solid reagents added directly to the capillaries, and due to possible loss of volatile components during the sealing of the capillary. These uncertainties can account for the underestimated experimental Au solubilities compared to those calculated using database of Akinfiev and Zotov (2010).

We estimate the concentration of Au dissolved in chloride melt as ~ 1 wt%. This concentration (in mol fraction scale) is ca. 10 times lower than the concentration of oxygen produced by decomposition of potassium persulfate. However, it is more than likely that the equilibrium Au concentration has not been attained due to slow kinetics of Au dissolution in chloride melt: we were not able to detect dissolved Au during one hour XAS experiment preformed at 650 °C with capillaries which were not subjected to preconditioning. Therefore, we can conclude that the solubility of Au in chloride melts can be very high, but the quantitative thermodynamic description of the Au solubility needs further experimental examination.

3.2. EXAFS spectra fitting

Roughly speaking, temperature, pressure, and the experimental system composition have little effect on the shape of Au L_3 -edge EXAFS spectra and their Fourier transforms (at least at $k < 10 \text{ Å}^{-1}$ and R < 3 Å, Fig. 3a and b, respectively). Results of EXAFS spectra fitting preformed by means of ARTEMIS computer program are shown in Fig. 4 and collected in Table 4. In general, these data are consistent with the results of X-ray absorption spectroscopy, quantum chemical, and AIMD calculations previously reported in the literature (Pokrovski et al., 2009; Mei et al., 2013). In chloride-bearing fluids two Cl⁻ ions are the nearest neighbors of Au with $R_{\text{Au-Cl}} = 2.25 - 2.28 \text{ Å}$. Results of the present study, however, reveal some important features of Au chemistry which were not manifested previously. It follows from data of Table 4 that Au-Cl distance in the first coordination sphere weakly decreases with increasing pressure. These small systematic changes were monitored in all the studied systems. The most pronounced change in the $R_{\text{Au-Cl}}$ of 0.02 Å was observed between experiments cap118 and cap117 (the pressure increased from 2100 to 4500 bar

for fluids of identical composition). The maximum value of $R_{\text{Au-Cl}} = 2.28 \text{ Å}$ was monitored in dry chloride melt when pressure did not exceeded one bar. Therefore, one can conclude that increase of pressure induces contraction of the first coordination sphere of Au.

In all the experiments the presence of a heavy Me atom (Me = K or Cs) was observed in the second coordination sphere of Au (the contribution of Au-Me-Au path to the Fourier transform of EXAFS spectra is shown in Fig. A1). The coordination numbers and Debye-Waller parameters of cations were estimated with high uncertainty, but inclusion of Au-Me-Au path in the fits of the EXAFS spectra improved considerably the fit results in the 2.5 - 4 Å range (not corrected for phase shift) in comparison with the model without the distant-coordination-sphere cation (Table A1, Fig. A2). When it was possible to calculate the coordination number of Me atoms $(N_{\rm Me})$ in the second coordination sphere of Au the $N_{\rm Me}$ fell within 0.2 - 0.6. In contrast to the observed contraction of the first coordination sphere of Au with increase of pressure, the opposite effect was monitored for the Me atoms in the second coordination sphere. The increase of the aqueous fluid pressure resulted in the increase of the Au-Me distance. The effect of pressure is of sharp focus in experiments cap118 $(R_{\text{Au-Me}} = 3.31 \pm 0.05 \text{ Å}, P = 2100 \text{ bar}, d = 0.68 \text{ g} \cdot \text{cm}^{-3})$ and cap117 $(R_{\text{Au-Me}} = 4.14 \pm 0.03 \text{ Å}, P = 4500 \text{ m})$ bar, $d = 1.04 \text{ g} \cdot \text{cm}^{-3}$). We note here that, despite high uncertainty, decrease of the $R_{\text{Au-Me}}$ is of systematic character. It was observed, at a quantitative level, in all the experimental systems even though the exact change of the $R_{\text{Au-Me}}$ as a function of TP-parameters can not be determined via the experimental spectra fitting.

The observed changes of the $R_{\text{Au-Me}}$ can be explained by the effect of dielectric constant on the weakly bound distant-coordination-sphere ion. The decrease of density results in the decrease of the fluid dielectric constant which promotes electrostatic interaction of the negatively charged AuCl_2^- complex and positively charged Me^+ ion. As a result, $R_{\text{Au-Me}}$ decreases with decrease of pressure (density, dielectric constant) due to the enhancement of coulomb interaction.

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3.3. Results of AIMD simulations

Simulation results are given in Table 2 (calculation parameters), Table 5 (geometry of Au species), and shown in Fig. 5. All the simulations for aqueous fluids were performed for 450 °C in order to compare the effect of the system composition on the Au-Cl complexing and the X-ray absorption spectra, and for 650 °C in the case of chloride melt. The first coordination sphere of Au comprises two chloride atoms in all the simulations but simulation F (chloride melt). In the latter case the presence of Cl⁻ ions at 2.6 - 3 Å induces an asymmetric shape of Cl atomic density function (Fig. 5). The distant coordination sphere of Au is composed of H₂O, Cl, and alkaline metal cations. The atomic density curves for Na, K, and Cs is of diffuse character with maxima located between 3 and 6 Å. This shape of the density curves confirms our conclusion based on the analysis of EXAFS spectra about weak electrostatic interaction of AuCl₂ with the distant-coordination-shell cations. In this case the cation does not have a definite position, but the $R_{\text{Au-Me}}$ fluctuates within ~3 - 6 Å range. The $R_{\text{Au-Me}}$ depends on the cation type and increases with increasing cation atomic weight ($R_{\text{Au-Na}}$ = 2.8 - 4.6 Å, $R_{\text{Au-Cs}} = 3.3$ - 6 Å). The coordination number N_{Me} is not a constant but increases with increasing cationic weight and salt concentration (Table 5), which is unlike for a stable aqueous complex which has a definite composition. Computation G, which was performed for the same system composition as the computation E, but with extended atomic basis set, yielded similar results. In both calculations Cs atoms fluctuate at a distance of 3 - 6 Å with the maximum of atomic density at ~ 4 Å. Therefore, the DZVP basis set is accurate enough for AIMD investigation of hydrothermal Au complexing.

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3.4. AIMD-EXAFS spectra and comparison with experimental data

EXAFS spectra simulated using atomic configurations produced by AIMD calculations are shown in Fig. A3 (chloride melt) and Fig. A4 (hydrothermal fluids). The contribution of cation in the second coordination sphere of Au to the Fourier Transform (FT) of calculated EXAFS signal increases with increasing cation atomic number and concentration, but has a significant input only on the spectrum of chloride melt (Fig. A3). In case of chloride fluids (Fig. A4) the contribution of

the distant-coordination-sphere cation calculated from the AIMD atomic configurations is underestimated relatively to the experimental spectra fitting. Nonetheless, the AIMD-EXAFS calculations enabled us lo explore the cation contribution to the EXAFS spectra using wavelet transform (WT) diagrams.

Figure 6a shows contributions of scattering atoms (Cl, O, Cs) and multiple scattering contribution Cl-Au-Cl to the WT of calculated AIMD-EXAFS spectra. The maxima of Cl contribution to the 1st coordination sphere is located at $k \sim 6\text{Å}^{-1}$, the Cl-Au-Cl multiple scattering contribution is located at $k \sim 5.5 \text{ Å}^{-1}$, and the maxima of O contribution is at $k \sim 4.5 \text{ Å}^{-1}$ (the other local maxima of the WT of EXAFS in Fig. 6a correspond to the distant coordination sphere atoms and multiple scattering paths). The contribution of Cs is clearly seen in Fig. 6b which shows the WT of EXAFS signal of experiment cap118 – the high-temperature fluid of relatively low density (575 °C/2100 bar, $d = 0.68 \text{ g·cm}^{-3}$). Here the contribution of Cs in the second coordination sphere occurs at $k \sim 9.5 \text{ Å}^{-1}/R \sim 3 \text{ Å}$. The WT of EXAFS of fluid of the same composition at higher temperature and pressure (experiment cap 117, 450 °C/4500 bar, $d = 1.04 \text{ g·cm}^{-3}$) does not exhibit a notably heavy atom contribution except, probably, a small increase of the signal at $k \sim 9.7 \text{ Å}^{-1}/R \sim 4.5 \text{ Å}$. The observed decrease of the heavy atom contribution to EXAFS signal, which takes place with increase of the fluid density, is in line with experimental EXAFS spectra fittings performed using ARTEMIS program (section 3.2, Table 4).

3.5. AIMD-XANES spectra

The Au L_3 -edge HERFD-XANES spectra calculated using atomic configurations obtained by means of AIMD simulations are shown in Fig. 7. We calculated only the spectra of CsCl/NaCl/KCl melt, HCl- and CsCl-dominated fluids in order to reveal, on a quantitative level, the effect of the cation type (the lightest H⁺ vs. the heaviest Cs⁺) and cation concentration on the shape of the HERFD-XANES spectra. It can be seen in the figure that the WL intensity depends on the system composition. The most intense WLs correspond to systems with low salt concentration or pure HCl fluid (spectra AuCl₂Cs and AuCl₂H, correspondingly). The WL intensity of 4*m* CsCl and CsCl/NaCl/KCl melt (spectra AuCl₂Cs₄ and melt, correspondingly) is lower. The decrease of the WL intensity in salt-rich systems is similar to the one observed in our HERFD-XAS experiments discussed in section 3.1. At the same time, the shift of the e.j. and the WL to lower energies in salt-rich systems predicted by the simulations were not observed in the experiments. We have to note that the energy positions of the spectral features in the *ab initio* calculations can be subjected to substantial uncertainty.

The observed peculiarities of the WL intensity can be explained by the effect of cation on the electronic density distribution within $AuCl_2$ complex. Figure 8a demonstrates that the probability of finding a second-coordination-shell cation of Au has a maximum at a distance of ca. 4 Å at Cs-Au-Cl angle of 65°. As an alkali metal cation approaches Au the positive charge of Au decreases (Fig. 8b) due to the decrease of 5d unoccupied DOS. This behavior of Au charge state yields the WL intensity decrease. Thus, the presence of an alkali metal cation in the distant coordination sphere of Au accounts for the observed decrease of the WL intensity of Au L_3 -edge HERFD-XANES spectra of salt-rich systems.

3.6. Speciation and concentration of Au in natural chloride fluids and melts

Salinity (chlorinity) of natural hydrothermal fluids varies in wide limits from almost dry melts to diluted aqueous solutions. Results of our study demonstrate that, independently of fluid chemistry (cation type and chloride salt concentration), AuCl₂ is the dominant Au species in fluids of moderate to high density. The Au dissolution in aqueous fluids can be described by the reaction,

$$470 Au_{(cr)} + HCl_{(aq)} + Cl = AuCl_{2} + 0.5 H_{2(g)} , (4)$$

or, involving the proton activity (or pH = $-\log a(H^+)$),

$$Au_{(cr)} + H^{+} + 2 Cl^{-} = AuCl_{2}^{-} + 0.5 H_{2(g)}$$
(5)

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

$$Au_{(cr)} + HCl_{(aq)}^{\circ} + Cl + 0.25 O_{2(g)} = AuCl_{2} + 0.5 H_{2}O$$
 (6)

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

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$$Au_{(cr)} + 2 Cl^{2} + 0.25 O_{2(g)} = AuCl_{2} + 0.5 O^{2},$$
 (7)

or, in terms of neutral species,

$$Au_{(cr)} + 2 NaCl + 0.25 O_{2(g)} = NaAuCl_2 + 0.5 Na_2O$$
 (8)

These reactions imply that the solubility (concentration) of Au is governed by (i) the oxygen fugacity, (ii) the fluid acidity, and, (iii) the chloride concentration. It follows from Eq. (4-8) that the dissolved Au concentration in equilibrium with native gold increases proportionally to the first power of fluid acidity, the second power of chloride salt concentration, and decreases as a square root of hydrogen fugacity (or increases proportionally to $f(O_2)^{0.25}$). Predominance field of AuCl₂-falls within acidic range of pH. In sulfide-rich fluids the Au solubility is controlled by hydrosulfide complex Au(HS)₂-in the pH range from weakly acidic to alkaline, even in the presence of chlorides (e.g., Renders and Seward, 1989; Shenberger and Barnes, 1989; Benning and Seward, 1996; Stefánsson and Seward, 2003; Barnes, 2015; Trigub et al., 2017, see also references cited in these papers), or by hydroxide complex Au(OH)° at high (> 400 °C) temperatures (Zotov et al., 1985; Stefánsson and Seward, 2003). The dissolved Au concentration in dry chloride melt is mostly controlled by the oxygen fugacity and increases as a linear function of $f(O_2)^{0.25}$ (Eq. 7, 8).

Results of our spectroscopic experiments revealed the difference between the first and second coordination spheres of Au complexes. The Cl⁻ ions in the nearest coordination sphere of Au are bonded to the central atom via the formation of strong chemical bond whose nature can be described in terms of interaction between the metal and ligand atomic orbitals with the formation of molecular orbitals (cf. Figgs and Fitchman, 2000). In contrast, cations (H⁺, Na⁺, K⁺, Cs⁺ in the case of the present study) are weakly bonded to the negatively charged AuCl₂⁻ complex in the distant coordination sphere of Au mostly via the electrostatic interaction. The difference in the chemical bond properties is clearly demonstrated by the contrasting behavior of the atoms in the nearest and distant coordination spheres depending on the experimental *TP*-parameters. Increasing pressure results in weak contraction of the first coordination sphere of Au, whereas conjugated increase of

the dielectric constant of hydrothermal fluid causes expansion of the distant coordination sphere composed of alkali metal cations. The number of cations in the distant coordination sphere of Au also depends upon pressure, cation type, and salt concentration. From the point of view of thermodynamics this means that the dissolution of Au can be explicitly described by the equilibrium constants of chemical reactions which involve the formation of AuCl_2 (Eq. 4-8), taking into account the influence of weak outer sphere coulomb interactions via the activity coefficients. This approach is valid for fluids with density of $d > 0.5 \text{ g} \cdot \text{cm}^{-3}$. In low-density fluids ($d < 0.3 \text{ g} \cdot \text{cm}^{-3}$), where interaction between charged species becomes much more pronounced and neutral species predominate, formation of ionic associates of the type of MeAuCl₂° and HAuCl₂° is hampered by the change of predominant Au complex from charged AuCl₂° to neutral AuCl°. The same conclusion might be done with regard to other 1st group *d*-elements (Cu, Ag) which form stable chloride complexes of similar composition. However, determination of composition and stability of chloride complexes of these metals can only be done via accurate solubility measurements.

The Au solubilities monitored in the present study are in close agreement with thermodynamic model of Akinfiev and Zotov (2010), which is based on the consideration of $AuCl_2^-$ complex. We note, however, that there is still a problem of disagreement between the values of $AuCl_2^-$ stability constants reported in the key studies on hydrothermal Au-Cl complexing of Zotov et al. (1991) and Stefánsson and Seward (2003). The Au solubility constants obtained in Zotov et al. (1991) are more than one log unit higher than those determined in Stefánsson and Seward (2003) for similar TP-parameters. The reason of this deviation is unclear. This problem can be resolved by additional measurements of Au solubility with rigorous in situ control of the most important physicochemical parameters (e.g., $f(H_2)$), and in situ experimental solution sampling. This is an ongoing study in our laboratory.

4. CONCLUSIONS

In the present study we recorded Au L₃-edge X-ray absorption spectra of aqueous chloridebearing fluids at temperatures 350 – 575 °C and pressures up to 4500 bar. The composition of the experimental fluids varied from HCl – dominated solutions to concentrated KCl and CsCl brines. X-ray absorption spectra of dry CsCl/NaCl/KCl melt were recorded at 650 °C. Interpretation of EXAFS spectra showed that, independently of the experimental fluid composition, AuCl₂ complex dominates the dissolved Au speciation. The distance between Au and the nearest-neighbor Cl atoms varies between 2.25 - 2.28 Å. This interatomic distance is pressure-dependent and decreases with increasing pressure from 2.28 Å (dry melt) to 2.25 Å (7.3m HCl/4.2m CsCl fluid at 4500 bar). The outer coordination sphere of Au is composed of cations located at a distance of 3.3 – 4.1 Å. In contrast to the inner coordination sphere, the outer coordination shell expands with increasing the fluid pressure (dielectric constant). The number of cations located in the outer coordination sphere increases with increasing cation atomic number and concentration. Such a contrasting behavior of atoms located in the nearest and distant coordination spheres of Au reveals the difference in the nature of the chemical bonds. Chloride ions are connected to Au by means of strong chemical bonding, whereas weak coulomb interactions between AuCl₂ and cations are responsible for the formation of the disordered distant coordination sphere. The presence of cations in the distant coordination sphere of Au is confirmed by the change of the shape of Au L_3 -edge HERFD-XANES spectra in salt-rich brine and dry melt with respect to the HCl-dominated fluid. The results of our study imply that the Au solubility in hydrothermal fluids in the wide range of TP-conditions and chlorides concentration can be described in terms of the formation of AuCl₂ complex using an appropriate model for the activity coefficients.

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Table 1. Compositions of experimental systems, experimental Au solubilities, and equilibrium concentrations of aqueous species calculated using thermodynamic data available in the literature: $AuCl_2^-$ - Akinfiev and Zotov (2010), CsCl $^\circ$ - Sverjensky et al. (1997), HCl $^\circ$ - Tagirov et al. (1997), other aqueous species, H₂O, and $Au_{(cr)}$ – SUPCRT92 (Johnson et al., 1992).

(//	111	, , ,	(01)		,	,					
Capillary No.	28	30	110	110	117	118	122	122	122	122	
110.	3.8 <i>m</i> HCl/	3.4 <i>m</i> NaCl/	3.8 <i>m</i> HCl/	3.8 <i>m</i> HCl/	7.3 <i>m</i> HCl/	7.3 <i>m</i> HCl/	3.8 <i>m</i> HCl/	3.8 <i>m</i> HCl/	3.8 <i>m</i> HCl/	3.8 <i>m</i> HCl/	
Solute	0.02 mNaCl/	$0.7m\mathrm{H}_2\mathrm{SO}_4$	0.57mKCl/	0.57mKCl/	4.2 <i>m</i> CsCl/	4.2 <i>m</i> CsCl/	7.2 <i>m</i> KCl/	7.2 <i>m</i> KCl/	7.2 <i>m</i> KCl/	7.2 <i>m</i> KCl/	
Concentr.	$0.5m\mathrm{H}_2\mathrm{SO}_4/$	$0.6m H_2SO_3$	$0.03m\mathrm{K}_2\mathrm{S}_2\mathrm{O}_8$	0.03mK ₂ S ₂ O ₈	$0.56m\mathrm{H}_2\mathrm{SO}_4/$	$0.56m\mathrm{H}_{2}\mathrm{SO}_{4}/$	$0.4m\mathrm{K}_2\mathrm{S}_2\mathrm{O}_8$	$0.4m\mathrm{K}_2\mathrm{S}_2\mathrm{O}_8$	$0.4m\mathrm{K}_2\mathrm{S}_2\mathrm{O}_8$	$0.4mK_2S_2O_8$	
- 3	$0.5m \text{ H}_2\text{SO}_3$				$0.12m\text{Na}_2\text{SO}_3$	$0.12m\text{Na}_2\text{SO}_3$					
d, g·cm ⁻³	0.53	0.67	0.71	0.71	1.04	0.68	0.96	0.96	0.96	0.96	
<i>t</i> , °C/	485 °C/	485 °C/	400 °C/	450 °C/	450 °C/	575 C°/	350 °C/	400 °C/	450 °C/	525 °C/	
P , bar	600 bar	700 bar	850 bar	2100 bar	4500 bar	2100 bar	150 bar	500 bar	1100 bar	1900 bar	
Calculated concentrations, mol·(kg H ₂ O) ⁻¹											
\mathbf{H}^{+}	5.73E-01	1.43E-02	4.43E-01	4.55E-01	2.21E+00	6.83E-02	3.01E-01	1.89E-01	1.04E-01	3.52E-02	
OH ⁻	1.03E-10	7.10E-09	2.72E-10	8.84E-10	1.44E-09	5.92E-09	1.93E-10	9.86E-10	2.87E-09	1.53E-08	
O_2	5.33E-10	1.62E-14	9.74E-19	1.05E-17	1.11E-19	4.37E-13	1.43E-16	6.73E-16	2.89E-15	1.48E-13	
H_2	3.48E-09	2.62E-07	8.69E-09	8.01E-08	2.91E-07	3.07E-06	1.53E-11	6.29E-10	1.10E-08	2.25E-07	
HSO ₃	8.89E-09	1.73E-06	1.90E-08	7.52E-08	1.40E-06	9.07E-07	8.91E-10	2.07E-08	2.34E-07	1.83E-06	
HSO ₄	3.94E-01	6.69E-01	4.04E-02	2.55E-02	9.33E-02	3.81E-02	7.43E-01	7.70E-01	7.47E-01	6.00E-01	
$\mathrm{SO_2}^\circ$	5.82E-01	6.15E-01	5.62E-03	2.04E-02	2.52E-01	6.27E-01	1.09E-04	2.91E-03	2.77E-02	1.86E-01	
SO_4^{2-}	8.30E-06	5.27E-04	8.20E-07	3.33E-07	1.24E-06	7.77E-07	1.18E-04	2.06E-04	1.13E-04	8.59E-05	
$\mathrm{H_2SO_4}^{\circ}$	1.20E-02	8.74E-04	1.39E-02	1.40E-02	3.30E-01	1.17E-03	5.63E-02	2.70E-02	2.42E-02	8.12E-03	
Cl	1.79E-02	3.15E-01	6.96E-01	7.55E-01	3.07E+00	6.03E-01	1.61E+00	1.93E+00	1.91E+00	1.75E+00	
HCl°	3.40E+00	6.11E-01	3.27E+00	3.21E+00	4.63E+00	5.87E+00	2.64E+00	2.77E+00	2.79E+00	2.38E+00	
Na ⁺	1.49E-02	1.01E+00			1.27E-01	9.45E-02					
NaCl°	4.89E-03	2.35E+00			1.11E-01	1.41E-01					
\mathbf{K}^{+}			3.65E-01	4.27E-01			2.85E+00	3.32E+00	3.40E+00	3.49E+00	
KCl°			2.65E-01	2.03E-01			5.15E+00	4.68E+00	4.59E+00	4.46E+00	
Cs ⁺					1.09E+00	1.50E+00					
CsCl°					3.08E+00	2.62E+00					
AuCl ₂	1.76E-01	4.39E-02	7.12E-02	1.01E-01	2.65E-01	1.02E+00	8.00E-01	8.06E-01	8.55E-01	1.17	
Au _{exp.} (a)	n/d	n/d	3E-02	5E-02	2E-01	7E-01	2E-01	2E-01	2E-01	2E-01	
<i>I</i> ^(b)	0.588	1.029	0.808	0.881	3.426	1.66	3.153	3.507	3.507	3.522	
pН	1.386	2.864	0.881	0.797	0.154	1.87	1.366	1.586	1.753	2.19	
Eh	0.34	-0.039	0.28	0.235	0.255	0.017	0.378	0.279	0.186	0.042	
a 1: 1 A										-1-4 10.2	

^a dissolved Au concentration determined from the amplitude of the absorption edge height of Au L_3 -edge transmission spectra, estimated uncertainty ± 0.3 log units; ^b ionic strength.

Table 2. Results of *ab initio* molecular dynamics (AIMD) simulations performed for systems containing Au complexes of different compositions. Uncertainties correspond to the standard deviations.

AIMD calculation	Composition of simulation box	Box size, Å	Starting Au species ^a	T,K	Density ρ , g/cm ³ (without Au)	Density ρ , g/cm ³ (with Au)	Simula- tion time, ps
A	Au ⁺ , H ⁺ , 2Cl ⁻ , 55H ₂ O	14.7	Au(Cl)2	722 (46)	0.55	0.65	23
В	Au ⁺ , Na ⁺ , 2Cl ⁻ , 55H ₂ O	14.7	Au(Cl)2	722 (45)	0.56	0.67	23
С	Au ⁺ , K ⁺ , 2Cl ⁻ , 55H ₂ O	14.7	Au(Cl)2	722 (45)	0.57	0.67	25
D	Au ⁺ , Cs ⁺ , 2Cl ⁻ , 55H ₂ O	14.7	Au(Cl)2	722 (46)	0.62	0.72	23
Е	Au ⁺ , 4Cs ⁺ , 5Cl ⁻ , 55H ₂ O	14.7	Au(Cl)2	722 (48)	0.83	0.98	24
F	Au ⁺ , 48Cs ⁺ , 32Na ⁺ , 27K ⁺ , 5Cl ⁻ ,	16.3		923 (53)	2.13	2.17	15
G	Au ⁺ , 4Cs ⁺ , 5Cl ⁻ , 55H ₂ O	14.7	Au(Cl)2	723 (49)	0.83	0.98	15

^a starting system configurations were prepared using GROMACS software and included the Au complexes of indicated composition (see section 2.4 for calculation details). Starting geometry for F calculation was prepared using CsCl crystal lattice by substituting Cs atoms by Au, Na and K atoms, and correction of lattice constant to get the specified density.

Table 3. Position of edge jump (e.j.), white line (WL), and the next to WL feature (B) of Au L_3 -edge HERFD- and TFY-XANES spectra (as determined by IFEFFIT program). Uncertainty of the energy values is ± 0.5 eV.

Experimental system	Feature	Position, eV
HERFD-XANES	$5 (\pm 0.2 \text{ eV})$	
cap 5407	e.j.	11919.5
Melt CsCl/NaCl/KCl	WL	11921.0
<i>t</i> = 650 °C	В	11930.8
45.5at.%/30 at%/24.5 at%	ΔB -WL a	9.8
cap 28	e.j.	11919.3
$t = 485 ^{\circ}\text{C}, P = 600 \text{ bar}$	WL	11920.7
$d = 0.53 \text{ g·cm}^{-}$	В	11931
3.8 <i>m</i> HCl/0.02 <i>m</i> NaCl	ΔB -WL ^a	10.3
$0.5m \text{ H}_2\text{SO}_4/0.5m \text{ H}_2\text{SO}_3$	△D-WL	10.0
cap 30	e.j.	11919.6
$t = 485 ^{\circ}\text{C}, P = 700 \text{ bar}$	WL	11920.8
$d = 0.67 \text{ g} \cdot \text{cm}^{-1}$	B	11930.5
3.4 <i>m</i> NaCl	ΔB -WL ^a	9.8
$0.7m \text{ H}_2\text{SO}_4/0.6m \text{ H}_2\text{SO}_3$	△D-WL	7.0
0.7m H ₂ SO ₄ /0.0m H ₂ SO ₃ TFY-XANES ((+ 0.5 aV)	
cap 2156	i ·	11919.5
Melt CsCl/NaCl/KCl	e.j.	
45.5at.%/30 at%/24.5 at%	WL	11922.5
	В	11935.5
t = 650 °C	. :	11010
cap 2156	e.j.	11918
Crystallized melt CsCl/NaCl/KCl	WL	11920
Ambient temperature	В	11934
Cap110	e.j.	11919
$t = 400 ^{\circ}\text{C}, P = 850 \text{bar}$	WL	11922
$d = 0.71 \text{ g} \cdot \text{cm}^{-3}$	В	11931/11933
3.8 <i>m</i> HCl/0.57 <i>m</i> KCl/		
$0.03 \ m \ K_2S_2O_8$		
Cap110	e.j.	11919
$t = 450 ^{\circ}\text{C}, P = 2100 \text{bar}$	WL	11922
	В	11931/11933
Cap117	e.j.	11919
$t = 450 ^{\circ}\text{C}, P = 4500 \text{bar}$	WL	11922
$d = 1.04 \text{ g} \cdot \text{cm}^{-3}$	В	11931/11933
7.3 <i>m</i> HCl/4.2 <i>m</i> CsCl		
0.56m H ₂ SO ₄ /0.12m Na ₂ SO ₃		
Cap118	e.j.	11919
$t = 575 ^{\circ}\text{C}, P = 2100 \text{ bar}$	WL	11922
$d = 0.68 \text{ g} \cdot \text{cm}^{-3}$	В	11932
7.3 <i>m</i> HCl/4.2 <i>m</i> CsCl		
0.56m H ₂ SO ₄ /0.12m Na ₂ SO ₃		
Cap122	e.j.	11918.5
$t = 350 ^{\circ}\text{C}, P = 150 \text{ bar}$	WL	11921.5
$d = 0.96 \text{ g} \cdot \text{cm}^{-3},$	В	11931/11933
3.8 <i>m</i> HCl/7.2 <i>m</i> KCl/		
$0.4m \text{ K}_2\text{S}_2\text{O}_8$		
Cap122	e.j.	11918.5
$t = 400 ^{\circ}\text{C}, P = 500 \text{bar}$	WL	11921.5
, 2 500 our	В	11930/11933
Cap122	e.j.	11918.5
$t = 450 ^{\circ}\text{C}, P = 1100 \text{bar}$	WL	11916.5
7 - 750 C, 1 - 1100 bai	B	11921.3
Cap122	e.j.	11930/11933
$t = 525 ^{\circ}\text{C}, P = 1900 \text{bar}$	WL	11918.5
i = 525 C, i = 1700 bai	B	11921.3
a energy difference between maxima		
chergy unrefered between maxima	oi b aliu WI	Licatures.

Table 4. Results of EXAFS spectra fitting (ARTEMIS program, $S_0^2 = 0.84$, AuCs₂Cl₆ was accepted as initial model structure, Matsushita and Ahsbahs, 1998). Experimental spectra fits were performed in *k*-space if not indicated otherwise.

	EXAFS					
Atom	N R, Å	σ^2 , Å ⁻²	Fit quality			
			E°, eV	Red. chi ² (R-factor)		
		Cap 2156, melt CsCl/N	NaCl/KCl (k-space: 3 – 1	3), $t = 650 ^{\circ}\text{C}$		
Cl	2	2.28±0.007	0.004±0.001			
Cs	0.19±0.21	4.03 ± 0.05	0.002 ± 0.001	8.8±1.0	(0.033)	
MS^1	2	4.60 ± 0.05	0.021 ± 0.009			
	Cap 2156,	crystallized melt CsCl/	NaCl/KCl (k-space: 3 –	13), ambient ten	nperature	
Cl	1.88±0.32	2.29±0.009	0.001±0.001		(0.060)	
Cl	0.96±0.79	2.94 ± 0.03	0.003 ± 0.006			
Cs	1	3.26±0.08	0.012±0.011	8.0±1.7		
MS^1	1.88	4.58±0.05	0.004 ± 0.007			
MS^1	0.96±0.79	5.08 ± 0.05	0.003 ± 0.022			
	Cap110 (fit in <i>R</i> –space, <i>R</i> -range: 1.3-4.4; <i>k</i> -range: 3 – 11), $t = 400 ^{\circ}\text{C}$, $P = 850 ^{\circ}\text{bar}$ $d = 0.71 ^{\circ}\text{g} \cdot \text{cm}^{-3}$, $3.8m ^{\circ}\text{HCl}/0.57m ^{\circ}\text{KCl}/0.03 ^{o}\text{m} ^{\circ}\text{K}_2\text{S}_2\text{O}_8$					
Cl	2.27±0.13	2.27±0.005	0.006±0.001			
K	1	4.15±0.02	0.003±0.002	7.5±2.5	0.5 (0.007)	
MS^1	2.27±0.13	4.53±0.02	0.016±0.004			
	Cap110 (fit	in R-space, R -range: 1	1.3-4.4; <i>k-range</i> : 3 – 10),	$t = 450 ^{\circ}\text{C}, P =$	2100 bar	
Cl	2.08±0.23	2.26±0.009	0.004±0.001			
K	1	4.12±0.07	0.004±0.06	7.2±1.2	3.0 (0.013)	
MS^1	2.08±0.23	4.50±0.04	0.006±0.005			
Cap117 (fit in <i>R</i> –space, <i>R</i> -range: 1.3-4.4; <i>k</i> -range: 3 – 13), $t = 450 ^{\circ}\text{C}$, $P = 4500 ^{\circ}\text{bar}$ $d = 1.04 ^{\circ}\text{g} \cdot^{-3}$, 7.3 <i>m</i> HCl/4.2 <i>m</i> CsCl (+0.56 <i>m</i> H ₂ SO ₄ /0.12 <i>m</i> Na ₂ SO ₃)						
Cl	2.02±0.13	2.25±0.005	0.004±0.001		1.3 (0.013)	
Cs	0.59±0.86	4.11±0.03	0.005±0.007	4.6±0.8		
MS ¹	2.02±0.13	4.48±0.03	0.020±0.008			
Cap118 (fit in <i>R</i> –space, <i>R</i> -range: 1.3-4.4; <i>k</i> -range: 3 – 11), $t = 575$ °C, $P = 2100$ bar $d = 0.68 \text{ g} \cdot \text{cm}^{-3}$, 7.3 m HCl/4.2 m CsCl (+0.56 m H ₂ SO ₄ /0.12 m Na ₂ SO ₃)						
Cl	1.76±0.26	2.27±0.01	0.003±0.001			
Cs	0.30±0.63	3.31±0.05	0.003±0.013	6.6±1.6	13.7 (0.030)	
MS ¹	1.76±0.26	4.52±0.08	0.019±0.012			

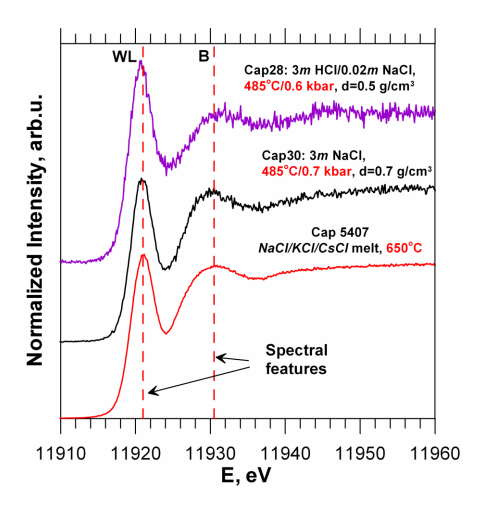
Ca	Cap122 (fit in <i>R</i> –space, <i>R</i> -range: 1.3-4.4; <i>k</i> -range: 3 – 13), t = 350 °C, P = 150 bar (possible presence of vapor bubble) $d = 0.96 \text{ g·cm}^{-3}$, 3.8 m HCl/7.2 m KCl/0.4 m K ₂ S ₂ O ₈				
Cl	2.04±0.18	2.28±0.006	0.003±0.001		
K	0.53±1.49	4.08±0.06	0.003±0.016	8.8±1.1	7.1 (0.022)
MS ¹	2.04±0.18	4.54±0.08	0.018±0.009		
	Cap122 (fi	t in R-space, R -range:	1.3-4.4; <i>k-range</i> : 3 – 10)	$t = 400 ^{\circ}\text{C}, P =$	500 bar
Cl	1.85±0.24	2.27±0.01	0.003±0.001	8.0±1.4	1.4 (0.026)
MS^1	1.85±0.24	4.54 ± 0.04	0.010 ± 0.006	0.U±1.4	
	Cap122 (<i>k-range</i> : $3 - 13$). $t = 450$ °C, $P = 1100$ bar				
Cl	2.11±0.19	2.27 ± 0.007	0.004 ± 0.001		3.6 (0.024)
K	1	4.06 ± 0.04	0.005 ± 0.004	8.1±1.1	
MS ¹	2.11±0.19	4.49 ± 0.04	0.016 ± 0.008		
	Cap122 (<i>k-range</i> : $3 - 12$). $t = 525$ °C, $P = 1900$ bar				
Cl	1.93±0.19	2.27±0.008	0.004 ± 0.001		
K	1	3.97 ± 0.02	0.014 ± 0.016	7.5±1.2	3.9 (0.026)
MS ¹	1.93±0.19	4.53±0.07	0.023±0.013		
$\frac{1}{MS}$ – 1	¹ MS – multiple scattering, Au-Cl1-Au-Cl2.				

Table 5. Geometries of Au aqueous complexes obtained by the *ab initio* molecular dynamics (AIMD) simulations (simulation details are given in Table 2). Uncertainties correspond to the standard deviations. Cation coordination number N corresponds to number of cations inside a sphere with radius of 6 Å.

AIMD calculation	Fluid composition	Complex	Au-Cl, Å	Cl-Au- Cl, °	Cation N
A	2m HCl	Au(Cl) ₂	2.35 (0.078)	167 (6.2)	-
В	2m NaCl	Au(Cl)2	2.35 (0.065)	169 (5.6)	0.17
С	2m KCl	Au(Cl)2	2.35 (0.074)	168 (5.8)	0.36
D	2m CsCl	Au(Cl)2	2.35 (0.087)	169 (6.1)	0.41
Е	4m CsCl	Au(Cl)2	2.35 (0.078)	165 (8.0)	1.31
F	CsCl/NaCl/ KCl	Au(Cl ₂)	2.40 (0.14)	159 (16.0)	8.32
G	4m CsCl	Au(Cl ₂)	2.34 (0.076)	168 (7.0)	1.63

Figure captions

- **Fig. 1.** Gold L_3 -edge HERFD-XANES spectra of Au-bearing solutions (Cap28, 30) and melt (Cap5407). Vertical dashed lines indicate the main spectral features.
- **Fig. 2.** Gold L_3 -edge TFY-XANES spectra of: (a) Au-bearing solutions (Cap110, 117, 118, 122) and melt (Cap2156), and (b) melt (Cap2156) at 650 °C and after crystallization at ambient temperature.
- **Fig. 3.** Comparison of experimental Au L_3 -edge EXAFS spectra (*left*) and their Fourier Transforms (*right*).
- **Fig. 4.** Results of Au L_3 -edge EXAFS spectra fits performed by means of ARTEMIS program. (a) CsCl/KCl/NaCl melt (Cap2156). (b-d) Aqueous fluids cap110, cap117, 118, cap122. Top -normalized k^2 -weighted EXAFS spectra; Bottom corresponding Fourier Transforms (not corrected for phase shift). Thick black line experimental spectra, thin dotted red line fit. Experimental TP-parameters and fluid compositions are indicated at the top of each figure.
- **Fig. 5.** Atomic densities and coordination numbers of Au complexes obtained using *ab initio* molecular dynamics (AIMD) simulations (see Table 2 for system compositions).
- **Fig. 6.** EXAFS signal and wavelet transform (WT) of the EXAFS signal (distance R (Å) vs. wavevector k (Å⁻¹)). (a) contributions of different scattering atoms to the EXAFS spectra, AIMD calculation E (4m CsCl). (b) experimental spectra cap117 and cap118. Not corrected for phase shift. The WT intensity scales are shown at the right side of each WT image.
- **Fig. 7.** Gold L_3 -edge HERFD-XANES spectra calculated using atomic configurations generated via AIMD simulations. Spectrum AuCl₂H corresponds to calculation A, AuCl₂Cs calculation D, AuCl₂Cs₄ calculation E, melt calculation F. Compositions of simulation boxes are given in Table 2.
- **Fig. 8.** Position of Cs calculated by AIMD (calculation E) (a), and positive Bader partial atomic charge of Au as a function of Au-Cs distance (b). Each point in Fig. (b) corresponds to average AIMD configuration for ±0.12 Å interval around the indicated Au-Cs distance.



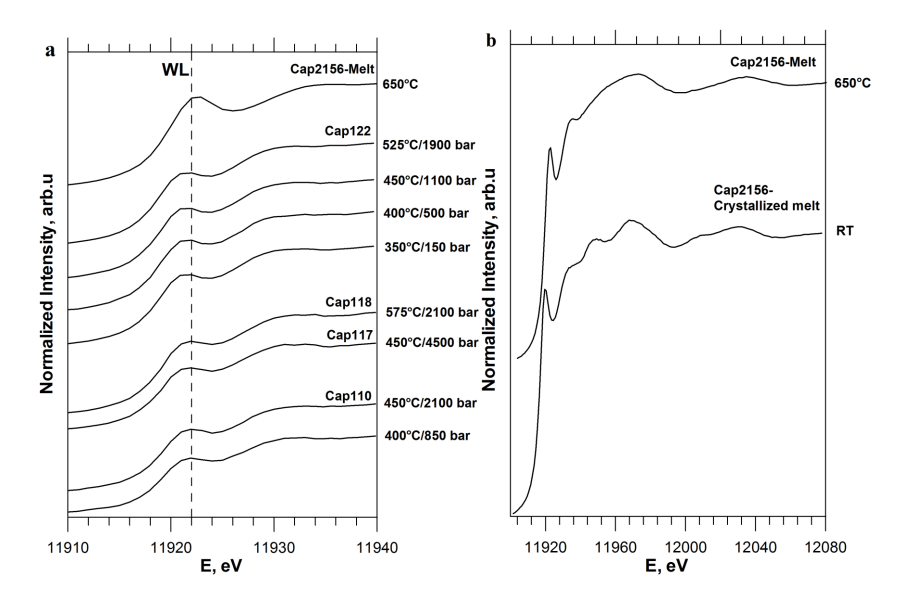


Fig. 2

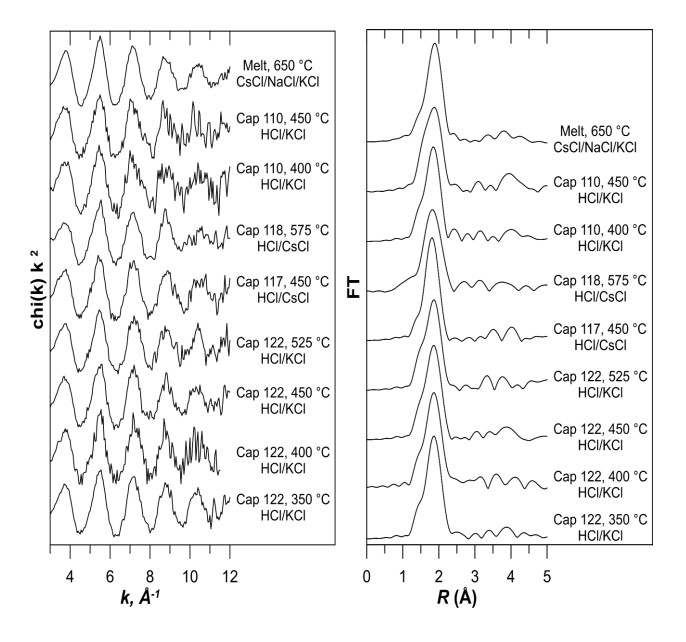
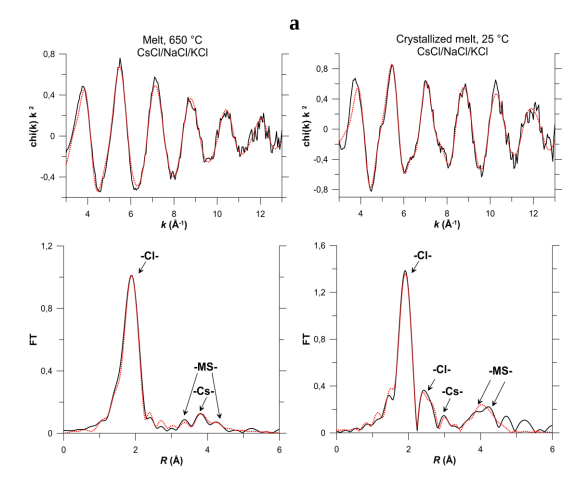


Fig. 3



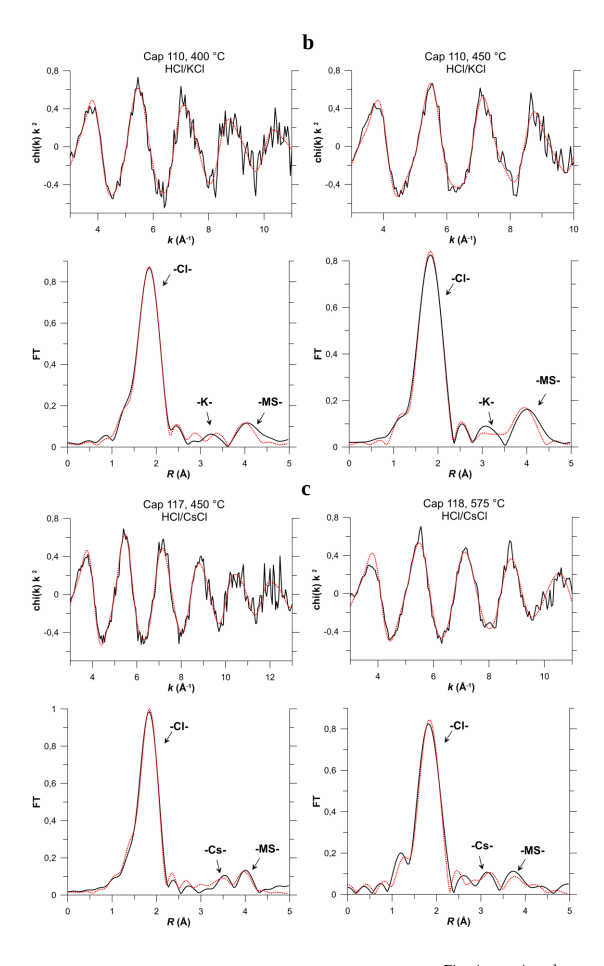


Fig. 4 - continued

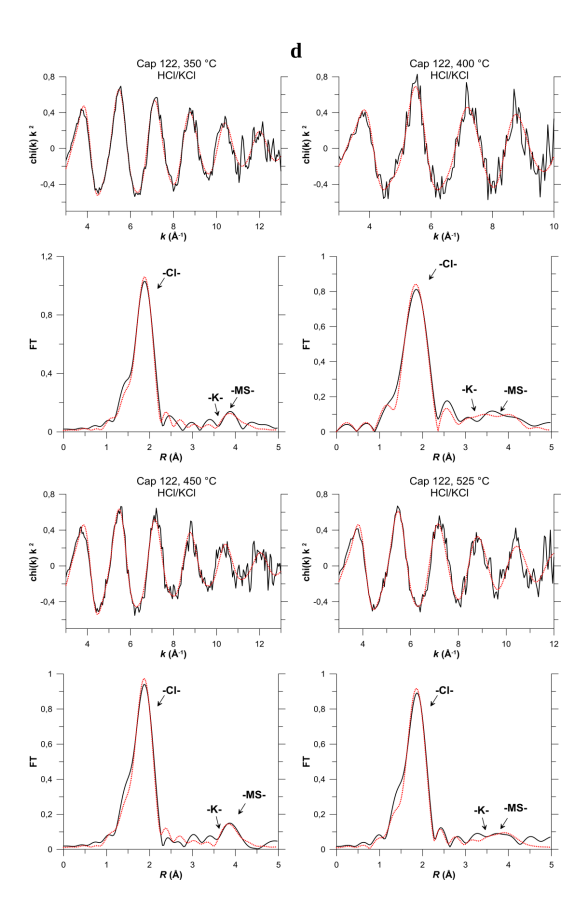


Fig. 4 – continued

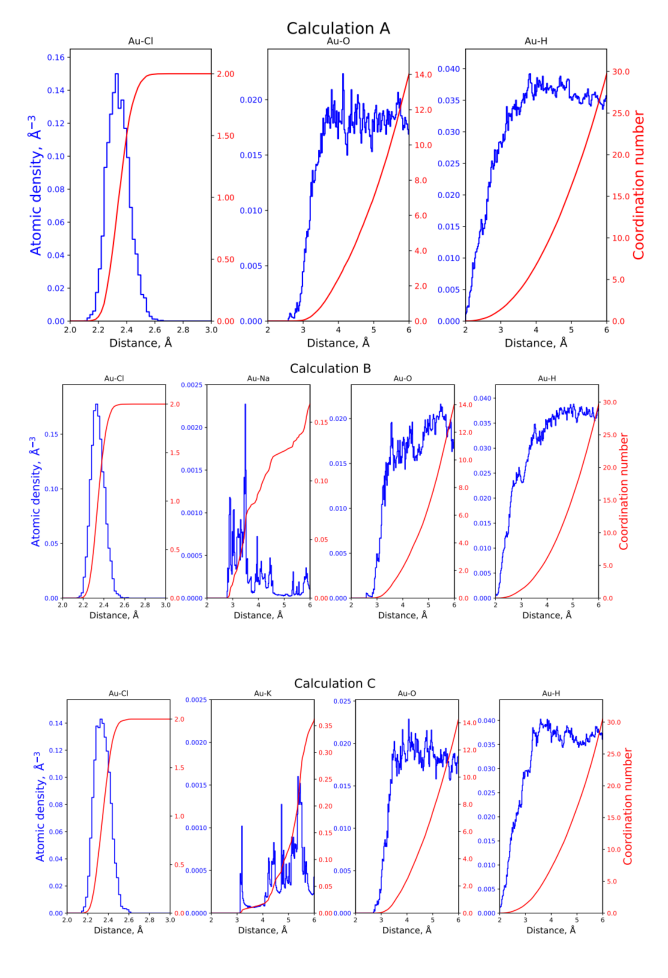


Fig. 5

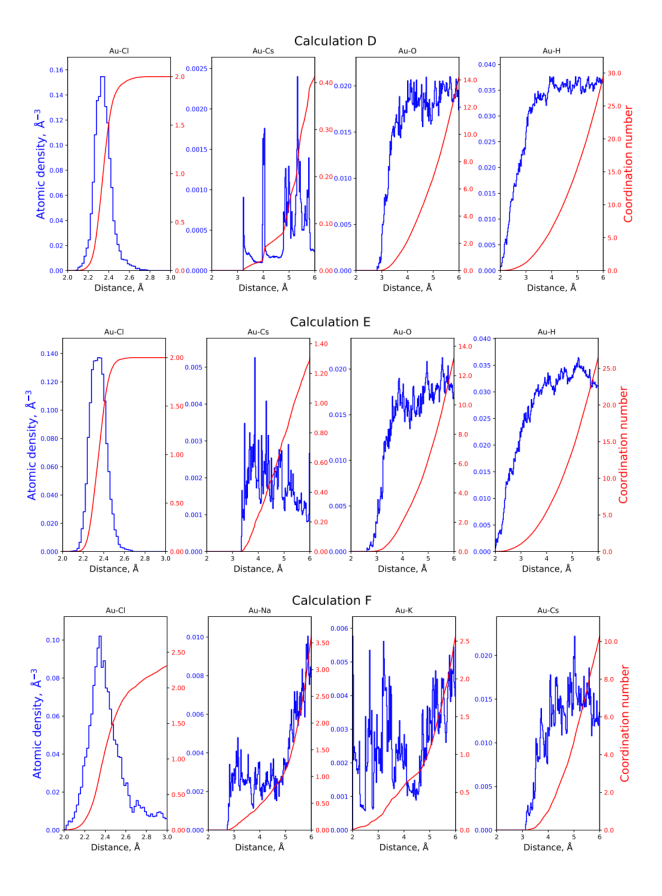


Fig. 5 – continued

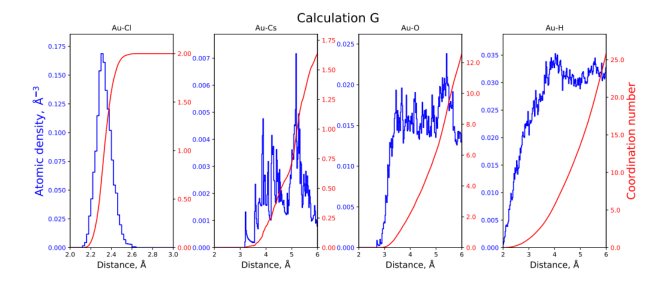


Fig. 5 – continued

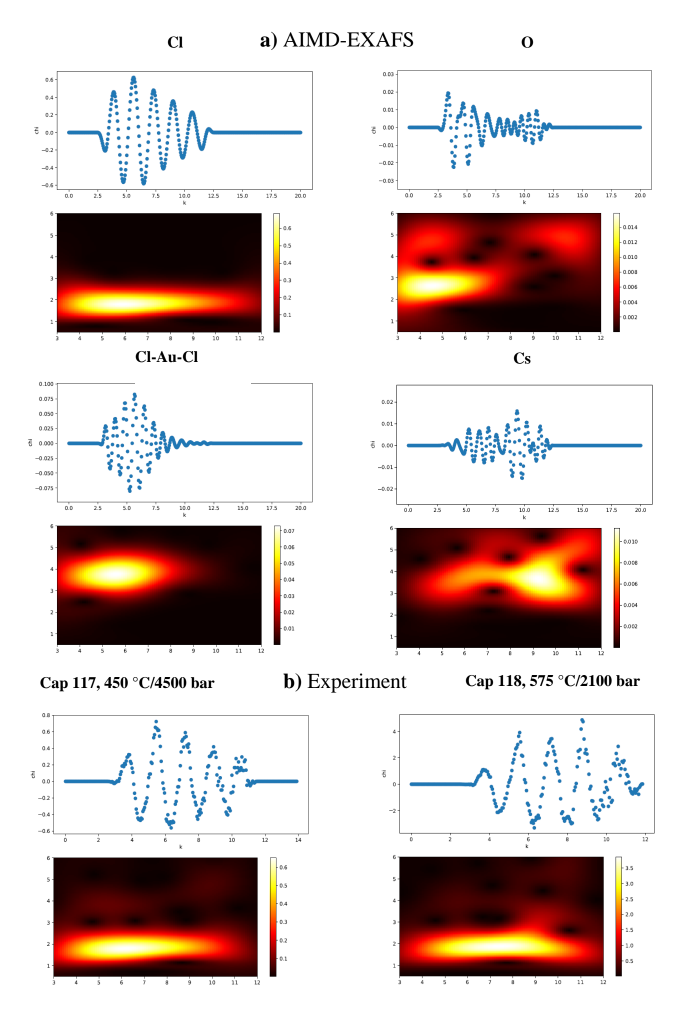


Fig. 6

Au L₃ XANES spectra

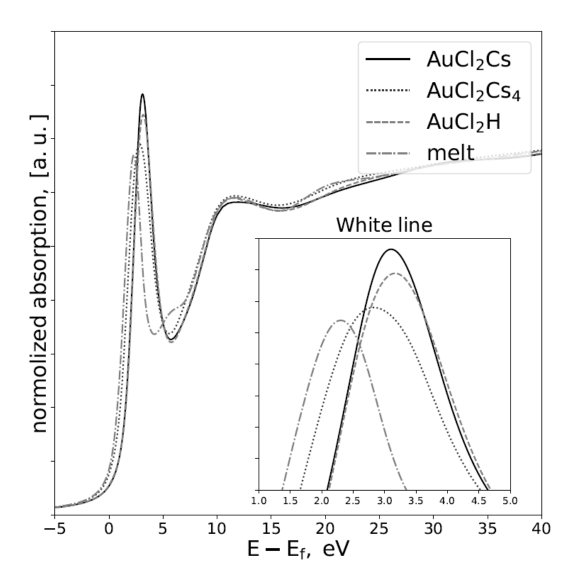


Fig. 7

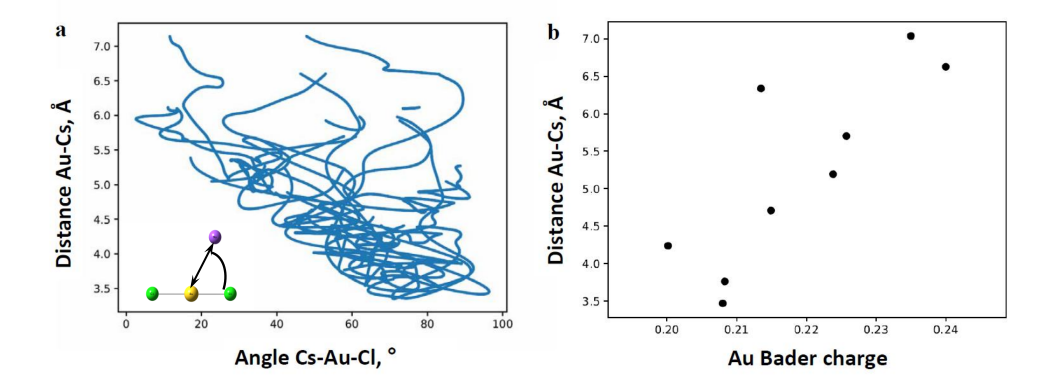


Fig. 8

Electronic Annex

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