The all-seeing eye of resonant Auger electron spectroscopy: a study on aqueous KCl

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Abstract

X-ray absorption and Auger electron spectroscopies are powerful tools to probe the electronic structure and immediate surroundings of ions in a solution. In this work we use a combination of these methods to study the electronic structure and decay in aqueous KCl at the K-edges of K⁺ and Cl⁻. Although the two ions are isoelectronic, their Auger electron spectra as a function of the photon energy exhibit notably different

features. To explain these differences, we carried out *ab initio* calculations of both the core excited states and the final Auger states of K^+ , Cl^- and their microsolvated clusters. Our calculations show that the energy order of the 3d and 4p orbitals is inverted in K^+ with respect to Cl^- . The reverse orbital order in the two ions is reflected in both the ordering of the core excited states and in the final states populated in the resonant Auger decay. The energetic proximity of the 3d and 4p virtual states in the bare K^+ ion leads to their mixing in the presence of the solvent, and to the population of the dipole forbidden $1s \rightarrow 3d$ state upon K-shell excitation in an aqueous solution. The resonant Auger decay of this state results in a separate feature in the Auger electron spectrum of K^+ which is absent in the spectrum of Cl^- .

Introduction

X-ray absorption and Auger electron spectroscopies are powerful tools to study the electronic structure and the nearest environment of atoms and molecules in gas, liquid and solid phase. Understanding how atoms or molecules respond to irradiation with x-rays gives insight into the structure of solutions (Ref.¹ and references therein), and the mechanism of radiation damage ²⁻⁴. Upon absorption of an x-ray photon, core excited or core ionized states of a specific atom are populated depending on the photon energy. The relaxation of these highly energetic states involves an ultrafast cascade of intraatomic processes, such as radiative and Auger decays. Furthermore, if the initially excited or ionized species is embedded in an environment, interatomic processes such as charge and energy transfer ⁴⁻⁸ are possible.

The course of a decay cascade depends on the character of the initially populated states. This has been well understood in atoms and molecules in gas phase⁹⁻¹⁴. In the case of a core ionized state, the Auger decay process, designated as normal Auger decay (see Fig. 1), leads to the population of doubly ionized final states localized on the initially ionized unit ⁹⁻¹². Auger processes in rare gas clusters have been also investigated (refs). The normal Auger decay process in clusters proceeds similarly to that in atoms or molecules. However,

in the case of a core excited state, the resonant Auger process competes with the process of delocalization of the excited electron in clusters. If the initially core excited electron delocalizes within the lifetime of the core hole, then normal instead of resonant Auger decay is observed ¹⁵.

In a solution, the electronic decay processes initiated by x-ray photoabsorption are different compared to those in rare gas clusters due to the shorter distances and stronger interatomic interactions. In particular, the solvent molecules have two effects – first, they strongly affect the excited ¹⁶ or ionized states of the ion and second, they can participate in the decay processes, leading to the population of charge-separated final states, and ionization of the surrounding environment ^{4-6,8}. Moreover, the process of delocalization of the initially excited electron also occurs in aqueous solutions ^{17,18}. In the case of pure water, the rate of delocalization of the O1s excited electron takes place on a femtosecond to sub-femtosecond time scale depending on the photon energy, thus being commensurate with the lifetime of the O1s core hole which is 6 fs ¹⁷. The O K-edge is located in the soft x-ray range of photon energies. Going higher in photon energy, in the tender and hard x-ray regimes, the lifetimes of the core ionized or core excited states become even shorter, on the order of 1 fs. And thus, it is even more imperative to reveal whether the delocalization of the core excited electron occurs within the lifetime of the core hole.

The aim of this work is to elucidate the nature of the states populated upon x-ray irradiation of solvated ions in the tender x-ray regime, and furthermore, to understand whether the process of delocalization influences the resonant Auger decay. To this end, we used Auger electron spectroscopy together with x-ray absorption spectroscopy in the tender x-ray regime to study aqueous potassium chloride at the K-edges of both K⁺ and Cl⁻. In particular, we demonstrate experimentally that at photon energies below the K-edges of the two ions, localized core excited states are populated. These states undergo resonant Auger decay within less than 1 fs. In both ions, there is a competition between resonant Auger decay and delocalization of the excited electron. Using the core-hole clock method we show

that in K⁺ delocalization at the pre-edge is weak, whereas in the case of Cl⁻, due to the energetic proximity of the core excited state to the K-edge, the rate of delocalization is of the same order as that of the resonant Auger process. Moreover, we observe that although the K⁺ and Cl⁻ ions are isoelectronic, they have different fingerprints in the resonant Auger spectra. With the aid of high-level *ab initio* calculations of the initial and final states of the resonant Auger process of both the bare ions and their microsolvated clusters, we demonstrate that these differences result from different electronic structures of the two ions, thus confirming that the combination of XAS and AES techniques is a sensitive probe of the electronic structure of solutions.

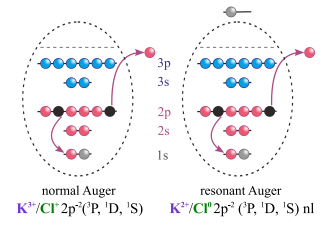


Figure 1: Schematic representation of the normal and resonant Auger processes of the isoelectronic K⁺ and Cl⁻ ions. The charges of the final Auger states in the two ions are also shown.

Methods

Experimental

For the present experiment we used the newly operational microjet setup that was specifically designed for the HAXPES station of the GALAXIES beamline ^{19,20}. A differentially-pumped tube in which the microjet head is inserted, is mounted on a 3-axes manipulator in front of the spectrometer lens. Two holes of 2 mm diameter allow the photons to go in and out. At the

end of the tube and in front of the lens, a 500 μ m diameter hole skimmer allows the electrons created at the interaction point to go in the direction of the spectrometer. The microjet head is mostly composed of a 30 μ m diameter vertical glass capillary facing a temperaturecontrolled catcher in CuBe having a 300 µm hole, and a camera. Piezo motors allow their precise alignment relative to each other and to the photon beam. The catcher is placed at a distance of about 5 mm from the capillary and is permanently pumped in order to extract the liquid. For the present experiment, a 0.5M KCl aqueous solution is injected in the capillary by a HPLC pump with a constant flux of 1.6 ml/min. The alignment of the setup is performed on a salted aqueous solution by measuring the O1s XPS peak intensity and by optimizing the liquid phase vs gas phase ratio. The pressure in the main chamber is kept below the 10^{-5} mbar range whereas it is kept at about 10^{-4} mbar in the differentially-pumped tube when the HPLC pump is ON. Our equipment is an updated version of the equipment used in the reference. 21 The aqueous potassium chloride solution was prepared by mixing >99% KCl salt with deionized water. Filtering and degazing procedures were systematically performed before injecting the solution. The spectrometer resolution of about 0.6 eV was achieved with the 500 eV pass energy and 0.5 mm slits. The photon energy resolution achieved at 2.8 keV and 3.6 keV was about 0.3 eV and 0.4 eV, respectively

Ab initio calculations

The theoretical X-ray absorption spectra were computed for the hexa-coordinated clusters of both ions, $K^+(H_2O)_6$ and $Cl^-(H_2O)_6$, which can be considered as representatives of the complete first solvation shell of the two ions $^{22-24}$. The two structures were optimized at the DFT level of theory using the B3LYP functional and the 6-311++G(2d,2p) basis set 25,26 . The geometry optimization was performed with the Gaussian 09 package 27 . In order to obtain more realistic structures corresponding to the bulk solution, we carried out constrained geometry optimization starting with the equilibrium gas-phase geometries 28 (REF for Cl-) belonging to the D₃ point group and increasing the angle θ between the K-O/Cl-O bond

and the C_3 axis to 55° in the case of K⁺ and 50° in the case of Cl⁻. This angle was chosen such that the O-K-O and O-Cl-O angles are around the maxima of the angular distributions obtained from quantum mechanics / molecular mechanics dynamical simulations in Ref.²⁴. Moreover, we fixed the K-O and Cl-O distances to 2.840 Å and 3.140 Å, respectively, such that they correspond to the distances from other theoretical and experimental works ^{22–24,29,30}. The hexa-coordinated structures are presented in Fig. 4.

The energies and transition moments of the core excited states of the microsolvated clusters were computed with the algebraic diagrammatic construction method for the polarization propagator³¹ within the core-valence separation approximation^{32–34} (CVS-ADC(2)x) as implemented in the Q-Chem package^{35–38}. In the case of Cl⁻ the 6-311++G(3df,3pd) basis set^{25,39} (excluding the f functions) was used for all atoms, whereas in the case of K⁺ we used the 6-311+G(2d,p) basis set^{25,26} on all atoms, and an additional set of 2s, 2p and 2d diffuse functions was added on K. In our calculations the core space comprises the 1s orbital of K⁺ or Cl⁻, whereas the remaining occupied orbitals are included in the valence space. For the calculations of the XAS spectra we used the C₂ point group in the case of K⁺(H₂O)₆ and Cl⁻(H₂O)₆. To account for the lifetime broadening due to the Auger decay of the core excited states, we convolved the theoretical spectra with a Lorentzian of FWHM 0.74 eV and 0.62 eV in the case of K⁺ and Cl⁻, respectively⁴⁰. We analyzed the core excited states by expanding the singly occupied natural orbitals (SONOs) ψ_i of the microsolvated clusters in the basis of SONOs of the bare K⁺ or Cl⁻ ion χ_{nl}

$$\psi_i = \sum_{nl} a^i_{nl} \chi_{nl} \tag{1}$$

where n and l stand for the principal and orbital quantum numbers as described in Ref. ¹⁶. The expansion coefficients a_{nl}^i show the degree of delocalization of the excited electron and the mixing of the core excited states in the crystal field created by the surrounding water molecules (see Fig. 4).

The final states following KLL resonant Auger decay of K⁺ and Cl⁻ were computed at the Configuration Interaction Singles (CIS) level using the Graphical Unitary Group Approach (GUGA) as implemented in the GAMESS-US package ^{41–43}. In order to account for the relaxation effects upon core ionization, we used a restricted open-shell Hartree-Fock reference wave function with a hole in the 2s orbital of both K⁺ and Cl⁻. We used the 6-311++G(2d,2p) basis set ²⁶ augmented with 2s, 2p, 2d diffuse functions on K⁺, and the cc-pVTZ basis set augmented with 6s, 6p, 6d diffuse Kaufmann-Baumeister-Jungen continuum-like functions ⁴⁴ on Cl⁻. The active space comprises the 2s and 2p orbitals of K/Cl with occupancy fixed to 6 and all virtual orbitals with occupancy fixed to 1. The remaining doubly occupied orbitals were frozen in the calculation. ⁴⁵

Results and discussion

Normal Auger decay

The $KL_{2,3}L_{2,3}$ normal Auger decay following K-shell ionization of aqueous K^+ and Cl^- can be written as follows

$$\gamma + \mathrm{K}_{\mathrm{aq}}^{+} \to \mathrm{K}_{\mathrm{aq}}^{2+}(1s^{-1}) \to \mathrm{K}_{\mathrm{aq}}^{3+}(2p^{-2}) + e_{\mathrm{Auger}}^{-}$$

 $\gamma + \mathrm{Cl}_{\mathrm{aq}}^{-} \to \mathrm{Cl}_{\mathrm{aq}}^{0}(1s^{-1}) \to \mathrm{Cl}_{\mathrm{aq}}^{+}(2p^{-2}) + e_{\mathrm{Auger}}^{-}$

It results in the population of $2p^{-2}(^{3}P, ^{1}D, ^{1}S)$ final states. The ^{3}P final states are expected to have very little intensity since at the first order they are forbidden from parity conservation rules. In addition, in our photon energy range of interest, the tail of the ^{1}D state described in the next paragraph, and the ^{1}D $2p^{-2}V'$ dispersive state (see next section) overlap in the same kinetic energy region making it impossible to properly determine the ^{3}P state position. In the case of Cl_{aq}^{-} , the lines corresponding to the Cl^{+} $(2p^{-2})$ ^{1}S and ^{1}D states are located at $2373.2\,\mathrm{eV}$ and $2382.1\,\mathrm{eV}$ kinetic energy (see Fig. 3). For K_{aq}^{+} the maxima of the ^{1}S and ^{1}D

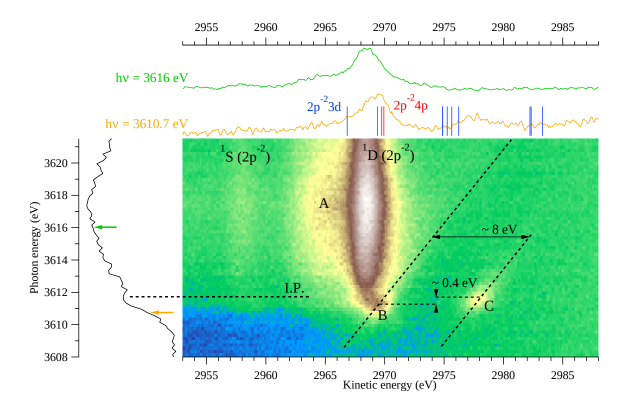


Figure 2: 2D map showing the kinetic energy of the electrons emitted in $KL_{2,3}L_{2,3}$ Auger decay vs the photon energy in the vicinity of the K-edge of aqueous K^+ . The black curve on the left represents the experimental partial electron yield spectrum of K^+ obtained after integrating over the kinetic energies of the Auger electrons in the energy range presented on the figure. The upper panel shows two spectra at photon energies 3610.7 eV, and 3616 eV below and above the ionization potential at 3611.9 eV, respectively. The bars in the pre-edge cut correspond to the final $2p^{-2}$ 3d (blue) and $2p^{-2}$ 4p(red) doublet resonant Auger states of K^+ computed at the CIS level (see Sec. for details). The features A, B and C are discussed in the text.

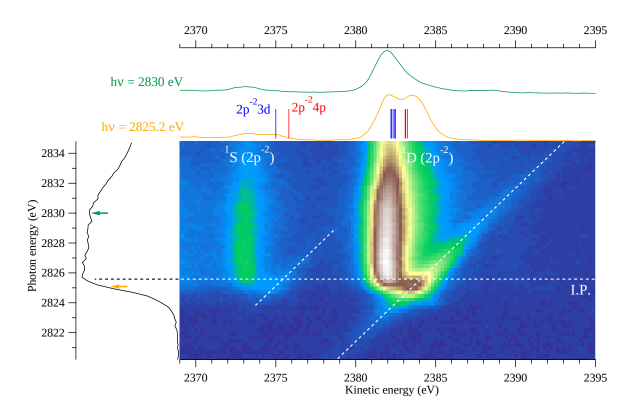


Figure 3: 2D map showing the kinetic energy of the electrons emitted in $KL_{2,3}L_{2,3}$ Auger decay vs the photon energy in the vicinity of the K-edge of aqueous Cl^- . The black curve on the left represents the experimental partial electron yield spectrum of Cl^- obtained after integrating over the kinetic energies of the Auger electrons in the energy range presented on the figure. The upper panel shows two spectra at photon energies $2825.2\,\mathrm{eV}$, and $2830.0\,\mathrm{eV}$ below and above the ionization potential at $2825.4\,\mathrm{eV}$, respectively. The bars in the pre-edge cut correspond to the final $2p^{-2}$ 3d (blue) and $2p^{-2}4p(\mathrm{red})$ doublet resonant Auger states of Cl^- computed at the CIS level (see Sec. for details).

KL_{2,3}L_{2,3} Auger lines are located at 2958 eV and 2968.4 eV, respectively (see Fig. 2).

The KL_{2,3}L_{2,3} Auger lines do not disperse with photon energy except close to threshold due to the interaction between the photoelectron and Auger electron, i.e. the so-called post-collision interaction (PCI). As a result of this interaction, first, the peaks in the Auger spectrum become asymmetric with a shoulder at high kinetic energies, and second, they are shifted to higher kinetic energies close to threshold ^{46,47}. Consequently, one can attribute the high kinetic energy shoulder of the ¹D and ¹S peaks on Figs. 2 and 3 as resulting from PCI effect.

Further, we compare the positions of the normal KLL Auger lines of both Cl_{aq}^- and K_{aq}^+ close to threshold with those recorded far from threshold, at photon energies $h\nu=5\,\mathrm{keV}$ (see Ref. 8 for details). In this case, the maxima of the 1D and 1S states were found at 2381.1 eV and 2372.3 eV for Cl_{aq}^- , and 2967.4 eV and 2957 eV kinetic energy for K_{aq}^+ , respectively. The lines observed at photon energies far from threshold and close to it appear to be shifted by $\sim 1\,\mathrm{eV}$. The magnitude of the shift is the same for both ions suggesting that it does not depend on the initial charge of the ion, and it is almost constant in the photon energy range presented here. A possible explanation of the shift observed in our experiment is given in Ref., 48 which focuses on the Auger decay of large Kr clusters on and just above the $3d_{5/2}$ ionization threshold. The observed $4d^4(^1D)$ Auger peak was found to be shifted by 0.7 eV to higher kinetic energies compared to the position of the peak far above threshold. Moreover, the shift did not vary with the photon energy close to threshold. Consequently, it was proposed that this feature originates from a process of internal ionization, i.e. excitation of the photoelectron into the conduction band. Further investigations are, however, planned in the case of liquid samples.

Finally, the normal Auger ¹D main line of K⁺ differs from that of Cl⁻ by the presence of a large shoulder (A on Fig. 2) on the low kinetic energy side at about 2965 eV kinetic energy. This shoulder is attributed to electron transfer from the water solvent molecules ⁸ to the 3d orbital of K⁺. In the case of Cl⁻, there was no experimental evidence of such intense

localized electron transfer processes.

Resonant Auger decay

The $KL_{2,3}L_{2,3}$ Auger decay following resonant K-shell excitation of solvated K^+ and Cl^- can be written as follows

$$\gamma + \mathrm{K}_{\mathrm{aq}}^+ \to \mathrm{K}_{\mathrm{aq}}^{+*}(1s^{-1}V) \to \mathrm{K}_{\mathrm{aq}}^{2+}(2p^{-2}V') + e_{\mathrm{Auger}}^-$$

 $\gamma + \mathrm{Cl}_{\mathrm{aq}}^- \to \mathrm{Cl}_{\mathrm{aq}}^{-*}(1s^{-1}V) \to \mathrm{Cl}_{\mathrm{aq}}^0(2p^{-2}V') + e_{\mathrm{Auger}}^-$

where V and V' denote the virtual orbitals in the excited and singly ionized excited states, i. e. in the presence of the $1s^{-1}$ and $2p^{-2}$ core holes. The character of these states will be discussed below.

The pre-edge regions of the x-ray absorption spectra of K⁺ and Cl⁻ shown to the left on Figs. 2 and 3 do not exhibit any high intensity features due to the lifetime broadening and energetic proximity of the core excited states to the ionization threshold. Consequently, solely from these spectra, one cannot conclude whether there are core excited states in the pre-edge structure, which can undergo resonant Auger decay. However, one can determine their excitation energies from the maxima of the resonant Auger features. Thus, for Cl⁻, the lowest core excited state is located at 2825.2 eV, which agrees very well with the position of the Cl⁻ 1s \rightarrow 4p excitation determined from Cl K-edge XAS experiments in MgCl₂.6H₂O and of SrCl₂/SrCl₂.6H₂O⁴⁹ and MCl₄ compounds⁵⁰. In the case of K⁺, there are two dispersive features with maxima at photon energies of 3611.2 eV (B) and 3611.6 eV (C), respectively (Fig. 2). The positions of these two core excited states are close to the energy of the 1s \rightarrow 4p excitation in bare K⁺, 3610.7 eV⁵¹.

The resonant Auger features produced in the decay of these core excited states appear to be quite different for Cl⁻ and K⁺. In the spectrum of Cl⁻ shown in Fig. 3 there are two dispersive features on the high kinetic energy side of the main ¹S and ¹D lines, i.e. at

2825.2 eV photon energy and 2374.6 and 2383.4 eV kinetic energy. In the case of K⁺, the ¹S dispersive line cannot be clearly identified due to the presence of strong background. The dispersive feature close to the ¹D main peak is observed (feature B) with a maximum located at $h\nu = 3611.2$ eV and 2969.2 eV kinetic energy. An additional feature appears as a separate island away from the main lines on the 2D map of K⁺. It is located at $h\nu = 3611.6$ eV and 2978.1 eV kinetic energy (feature C), thus it is separated by approximately 400 meV photon energy and 8.3 eV kinetic energy from the feature B.

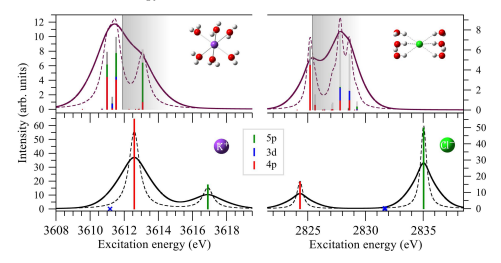


Figure 4: XAS spectra of the lowest K-shell transitions in the bare K^+ (lower left panel) and Cl^- (lower right panel) ions and their 6-coordinated clusters (upper left panel, $K^+(H_2O)_6$, and upper right panel, $Cl^-(H_2O)_6$). The theoretical stick spectra were convolved with a Lorentzian profile of FWHM 0.74 eV for K^+ and 0.62 eV for Cl^- (dashed line) and a Voigt profile to account for the lifetime broadening and the experimental resolution. The stick spectrum corresponds to the projections $|a_{nl}^i|^2$ of the SONOs corresponding to the core excited states of the 6-coordinated clusters on the basis of SONOs corresponding to the $1s \rightarrow 3d$, $1s \rightarrow 4p$, and $1s \rightarrow 5p$ states in the bare K^+ and Cl^- ions (Eq. 1). The theoretical XAS spectra of both K^+ and Cl^- were shifted to higher photon energies such that the excitation energies lowest core excited states correspond to the experimentally determined energies – $3610.7 \, eV$ in the case of K^+ , and $2825.2 \, eV$ in the case of Cl^- . The experimental ionization thresholds are depicted as grey boxes starting at photon energies of $3611.9 \, eV$ (K^+_{aq}) and $2825.4 \, eV$ (Cl^-_{aq}).

In order to rationalize the pre-edge region of the experimental XAS spectra and the differences in the AES spectra of K^{+}_{aq} and Cl^{-}_{aq} , we computed the lowest core excited states of the bare K^{+} and Cl^{-} ions and their hexa-coordinated clusters. The theoretical XAS spectra are presented in Fig. 4. In the bare ions (lowermost panels on Fig. 4), the

lowest energy peak corresponds to the dipole allowed $1s \to 4p$ state. The next dipole allowed state, $1s \to 5p$, is located 4.3 eV and 10.8 eV higher in the cases of K⁺ and Cl⁻, respectively. Together with the two dipole allowed transitions, we show the dipole forbidden $1s \to 3d$ states of the bare ions as blue crosses at photon energies 3611.17 eV in the case of K⁺ and 2831.68 eV in the case of Cl⁻, respectively. It is noteworthy that the positions of the $1s \to 4p$ and $1s \to 3d$ states are inverted in K⁺ and Cl⁻. In the case of Cl⁻ the $1s \to 4p$ excitation has lower energy and the $1s \to 3d$ excitation is close to the $1s \to 5p$ state. On the contrary, in K⁺ the $1s \to 3d$ excitation has lower energy and lies below the $1s \to 4p$ state. We note in passing that the intensity of the Cl⁻($1s \to 4p$) state is lower than that of the Cl⁻($1s \to 5p$) state contrary to what is observed in K⁺. This difference can be explained with the lower electron density of the 4p compared to the 5p electron in the region close to the core hole which thus results in the lower oscillator strength of the $1s \to 4p$ compared to the $1s \to 5p$ transition in Cl⁻ (see Fig. 2).

The water molecules in the first solvation shell have several effects on the core excited states. First, upon addition of water molecules, the degeneracy of the $1s \rightarrow 4p$ state is lifted and the intensity of the resulting states in the cluster drops. Moreover, the character of these states changes – they are no longer of pure atomic character but they rather interact with states of the neighboring water molecules (shown as grey bars) and with other closely lying states of the bare ion, such as the dipole allowed $1s \rightarrow 5p$ and dipole forbidden $1s \rightarrow 3d$ state. Thus, the latter also acquire intensity in the cluster due to mixing with the dipole allowed states in the ligand field of the solvent. A similar effect was observed in the XAS spectra of microsolvated clusters of Na^+ and Mg^{2+16} .

Further we assume that only the lowest peak in the theoretical XAS spectra is populated in the experiment for two reasons. First, due to the lifetime broadening, it spreads over approximately 2eV which coincides with the width of the pre-edge structure in the experimental XAS spectra. Second, the splitting between the first core excited state and the ionization threshold in the experiment is 1.2 eV for K⁺ and 0.2 eV for Cl⁻, and thus it is

smaller than the splitting between the first and second peak in the theoretical spectra (1.5 eV for K^+ and ~ 3 eV for Cl^- , Fig. 4). In the 6-coordinated cluster (Fig. 4 upper left panel), which represents the complete first solvation shell around K^+ , the lowest peak in the spectrum contains three states. The lowest and highest lying states are split by approximately 0.5 eV and they have mixed 4p and 5p character. The low intensity state in between these two states has a predominantly $1s \rightarrow 3d$ character. Since the dispersive feature B appears at lower excitation energies compared to the feature C, we assume that it is produced in the resonant Auger decay of the lowest core excited states of K^+ , which are predominantly of $1s \rightarrow 4p$ character. Moreover, we can attribute the feature C to the resonant Auger decay of the low intensity dipole forbidden $1s \rightarrow 3d$ state. Thus, we explain both the energy splitting of ~ 400 meV photon energy of the two features, and the fact that island C has lower intensity than feature B.

In the hexa-coordinated cluster of Cl^- , the solvent molecules have little influence on the position and character of the first peak. Since there are no other ionic states close the the $1s \rightarrow 4p$ state in the ion, this state preserves its character in the cluster and interacts with states of the nearest water molecules. We attribute the two dispersive features on the 2D map of Cl^- and associated with the 1S and 1D terms, to the resonant Auger decay of these core-excited states involving mostly the 4p orbitals of chloride.

To fully characterize the dispersive features on the experimental 2D maps, we also computed the lowest $K^{2+}(2p^{-2}nl)$ and $Cl^0(2p^{-2}nl)$ states of the bare ions corresponding to the lowest final spectator resonant Auger states. They are shown as bars in the upper panels on the experimental resonant Auger spectra on Figs. 2 and 3. In both cases of K^+ and Cl^- , we shifted the lowest $2p^{-2}4p$ states such that they coincide with the maxima of the dispersive features on the high kinetic energy part of the 1D main line. Out of all final states we show only the doublets. Since the initial core excited states are populated by photon excitation, only doublet states are efficiently populated in the resonant Auger decay. The $2p^{-2}nl$ states of K^+ and Cl^- are substantially different and they reflect the fact that the 3d unoccupied

orbitals of K^+ are lower than the 4p orbitals, which is the opposite of what is observed in Cl^- .

As mentioned above, we attribute the island B to the decay of the lowest lying core excited state of the hexa-hydrated K^+ cluster, which is of predominantly $1s \rightarrow 4p$ character. Supposing that this state undergoes mostly pure spectator resonant Auger decay, which is the case of the $1s \rightarrow 4p$ state in the isoelectronic Ar atom⁵², then the lowest states of $2p^{-2}4p$ character located between 2969 and 2970 eV are populated. As can be seen from the Auger electron spectrum at $h\nu = 3610.7 \,\mathrm{eV}$ (upper panel of Fig. 2), the lowest $2\mathrm{p}^{-2}4\mathrm{p}$ states of $\mathrm{K^{+}}$ at electron energy of $\sim\!2970\,\mathrm{eV}$ are separated by $\sim\!5$ and $\sim\!13\,\mathrm{eV}$ from the two groups of 2p⁻²3d states located at higher electron kinetic energies. Thus, the group of 2p⁻²3d states at ~ 2975 eV almost coincides with the position of island C. Consequently, we attribute this dispersive feature as originating from the resonant Auger decay of the dipole forbidden $1s \rightarrow 3d$ state of the 6-coordinated K⁺ cluster to the group of $2p^{-2}3d$ states lying around 2975 eV. The splitting between the 2p⁻²4p and 2p⁻²3d states in our calculation is smaller than the splitting between the islands B and C. This difference may be due to the fact that we do not account for the effect of the solvent molecules in our calculation In the figure we need to show the calculations on K+(H2O)6, and we need to interpret differently the next sentence. There is a group of 2p⁻²3d states at kinetic energies of 2982 eV. Since no additional experimental features are observed, we conclude that these states are not populated.

Another argument supporting the attribution of the island C as being of $2p^{-2}$ 3d character can be found by considering the electron transfer from the water molecules (W) discussed in reference.⁸ As a result of this process, an additional feature A is observed on the low-kinetic energy side of the ¹D main peak in K⁺. This feature has the configuration $K^{2+}2p^{-2}3d$ W⁻¹. ... Another argument playing in favor of the attribution of the island C as being of $2p^{-2}$ 3d character comes from the attribution of the A area as given in reference⁸ and from the energy splitting between A and C. The A area originates from a charge transfer process between water (W) and core-excited potassium and has the configuration $K^{2+}2p^{-2}3d$ W⁻¹.

The ionization potential of water in the liquid phase is about 11 eV which fits well with the observed A-C splitting. Based only on these two simple energetic arguments we can attribute the C island to the a K²⁺ 2p⁻² 3d configuration and note that neither 2p⁻² 3d nor charge transfer states are present in the Cl⁻ case.

In the $Cl^0(2p^{-2}nl)$ spectrum there are two groups of states split by about 7 eV (see upper panel of Fig. 3). The lower kinetic energy group corresponds to the $2p^{-2}(^1S)4p$ states, whereas the higher kinetic energy group corresponds to the $2p^{-2}(^1D)4p$ states. The splitting between the two groups is in good agreement with the experimental splitting between the dispersive features on the high kinetic energy sides of the 1S and 1D main peaks. Consequently, we attribute these dispersive features as resulting from the resonant Auger decay of the $1s \rightarrow 4p$ core excited state of Cl^- to the $2p^{-2}(^1S)4p$ and $2p^{-2}(^1D)4p$ final states. Similar dispersive features originating from the decay of the Cl ($1s \rightarrow 4p$) state were observed on the 2D map of chloromethane CH_3Cl recorded in the vicinity of the Cl K-edge in gas phase. In this case, however, additional lower-lying core excited state are observed. They result from excitation to the LUMO of CH_3Cl , which is a linear combination of the Cl 2p and Cl 3p atomic orbitals. Since the 3p shell is fully occupied in Cl^- , such a core excited state is not observed in our experiment.

Delocalization vs resonant Auger decay

As mentioned above, the delocalization of core excited electrons in aqueous solutions is ultrafast and as such it competes with the resonant Auger decay. In order to estimate the delocalization rate of the core-excited electron at the pre-edges of K⁺ and Cl⁻, we used the core-hole clock method as in the reference.^{54,55}

In the case of Cl⁻, and contrary to K⁺, it was possible to perform the same data treatment as in Ref. ⁵² (do you want to explain this data treatment in 1 sentence. now the reader is forced to check the other paper.), i.e. for each photon energy step, all components of the 2D map shown in Fig. 3 were isolated and their intensity integrated to get a partial electron

yield as a function of the photon energy. The result is shown in the SI. The figure shows that there is a large overlap between the resonant and normal Auger contributions, due to the proximity of the resonance to the ionization potential and due to the very short lifetime of the corresponding states. At the specific photon energy corresponding to the lowest core excitation, $h\nu=2825.2\,\mathrm{eV}$ (Fig. 3, upper panel) a double-peak structure is observed in the interval of kinetic energies $2380-2385\,\mathrm{eV}$. The position of the first peak coincides with the main $^1\mathrm{D}$ line resulting from normal Auger, whereas the second peak at $2383.5\,\mathrm{eV}$ corresponds to the final resonant Auger states $2\mathrm{p}^{-2}(^1\mathrm{D})4\mathrm{p}$. By fitting the peak with two Voigt functions, we determine the ratio of the intensities of these peaks to be ~ 1 . Consequently, the delocalization time is of the same order as the Auger lifetime, i.e. $\sim 1\,\mathrm{fs}$. The fast delocalization in this case is a result of the fact that the energy splitting between the Cl $^-$ ($1\mathrm{s} \to 4\mathrm{p}$) resonance and the ionization threshold is $0.2\,\mathrm{eV}$, and thus, smaller than the lifetime broadening of $0.62\,\mathrm{eV}$.

For potassium the treatment is more complex due to the presence of multiple simultaneous processes – normal, resonant Auger decay, charge transfer (CT) from solvent. To extract the intensity of each component from the 2D map shown in Fig. 2, one needs the spectral fingerprints of each process to be separated. However, as can be seen, this is not the case especially close to threshold in the kinetic energy region $2965 - 2970 \,\mathrm{eV}$. For instance at $3610.7 \,\mathrm{eV}$ photon energy on the high-kinetic-energy side of the $2\mathrm{p}^{-2}(^1\mathrm{D})$ state, there are contributions from the PCI tail and from the dispersive $2\mathrm{p}^{-2}(^1\mathrm{D})4\mathrm{p}$ state related to the B island. On the low-kinetic-energy side, the charge transfer process leads to a very large structure that unfortunately cannot be easily simulated by a known profile. However, the 1s core-hole lifetime is shorter for potassium than for chloride (0.9 vs. 1 fs) and moreover, the core-excited state appears $1.2 \,\mathrm{eV}$ below the ionization threshold whereas it is only $0.2 \,\mathrm{eV}$ for chloride. Therefore, one can expect a much less efficient delocalization process compared to $\mathrm{Cl}^-_{\mathrm{aq}}$.

Conclusion

Using a combination of x-ray absorption and Auger electron spectroscopy in the tender x-ray regime, in this work we studied the electronic structure of aqueous solution of KCl at the K-edges of both K and Cl. The Auger electron spectra of both ions as a function of photon energy exhibit features of normal as well as resonant Auger processes. To interpret the resonant Auger features in the experimental spectrum, we performed ab initio calculations on microsolvated clusters of K⁺ and Cl⁻. Our calculations show that the energy ordering of the 3d and 4p virtual orbitals of Cl⁻ is inverted compared to K⁺, and also that the energy splitting between the bright $1s \rightarrow 4p$ and dark $1s \rightarrow 3d$ core excited states is larger in the chlorine case. Thus, the energetic proximity of the 3d and 4p orbitals in the bare K⁺ ion results in the dipole forbidden $1s \rightarrow 3d$ state acquiring intensity in a solution as a result of mixing with the dipole allowed $1s \rightarrow 4p$ excitation. The spectator Auger decay of this state produces an additional dispersive feature which is manifest as a separate island in the Auger electron spectrum at high kinetic energies. In the case of Cl⁻ the two core excited states do not interact, and therefore, only fingerprints of the population and Auger decay of the dipole allowed $1s \rightarrow 4p$ state are observed in the spectrum. Moreover, using the core-hole clock method we estimated the time of delocalization of the core excited electron at the pre-edge region of both ions. Whereas in K^{+}_{aq} the delocalization of the core excited electron is slower than the resonant Auger process, in the case of Cl⁻_{aq} the two processes occur on a comparable timescale.

Our work shows that the combination of x-ray absorption and resonant Auger spectroscopies is a sensitive probe of the electronic structure of solvated ions. The reported results are an important first step in the study of the electronic decay processes following photoabsorption in the tender x-ray regime, and they can have implications in revealing the mechanisms of radiation damage in biologically relevant systems.

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Supporting Information Available

• suppinfo.pdf: contains the radial density distributions of the core excited states of the bare ions.

References

- (1) Smith, J. W.; Saykally, R. J. Chem. Rev. 2017, 117, 13909–13934, PMID: 29125751.
- (2) O'Neill, P.; Stevens, D. L.; Garman, E. F. *J. Synchrotron Radiat.* **2002**, *9*, 329–332.
- (3) Carugo, O.; Carugo, K. D. *Trends Biochem. Sci.* **2005**, *30*, 213–219.
- (4) Stumpf, V.; Gokhberg, K.; Cederbaum, L. S. Nat. Chem. **2016**, 8, 237–241.
- (5) Pokapanich, W.; Bergersen, H.; Bradeanu, I. L.; Marinho, R. R. T.; Lindblad, A.; Legendre, S.; Rosso, A.; Svensson, S.; Björneholm, O.; Tchaplyguine, M.; Öhrwall, G.; Kryzhevoi, N. V.; Cederbaum, L. S. J. Am. Chem. Soc. 2009, 131, 7264–7271.
- (6) Pokapanich, W.; Kryzhevoi, N. V.; Ottosson, N.; Svensson, S.; Cederbaum, L. S.; Öhrwall, G.; Björneholm, O. J. Am. Chem. Soc. 2011, 133, 13430.

- (7) Unger, I.; Seidel, R.; Thürmer, S.; Pohl, M. N.; Aziz, E. F.; Cederbaum, L. S.; Muchová, E.; Slavíček, P.; Winter, B.; V., K. N. Nat. Chem. 2017, 9, 708.
- (8) Céolin, D.; Kryzhevoi, N. V.; Nicolas, C.; Pokapanich, W.; Choksakulporn, S.; Songsiriritthigul, P.; Saisopa, T.; Rattanachai, Y.; Utsumi, Y.; Palaudoux, J.; Öhrwall, G.; Rueff, J.-P. *Phys. Rev. Lett.* **2017**, *119*, 263003.
- (9) Stoychev, S. D.; Kuleff, A. I.; Tarantelli, F.; Cederbaum, L. S. J. Chem. Phys. 2008, 129, 074307.
- (10) Demekhin, P. V.; Scheit, S.; Stoychev, S. D.; Cederbaum, L. S. Phys. Rev. A 2008, 78, 043421.
- (11) Demekhin, P. V.; Chiang, Y.-C.; Stoychev, S. D.; Kolorenč, P.; Scheit, S.; Kuleff, A. I.; Tarantelli, F.; Cederbaum, L. S. J. Chem. Phys. 2009, 131, 104303.
- (12) Ouchi, T.; Sakai, K.; Fukuzawa, H.; Higuchi, I.; Demekhin, P. V.; Chiang, Y.-C.; Stoychev, S. D.; Kuleff, A. I.; Mazza, T.; Schöffler, M.; Nagaya, K.; Yao, M.; Tamenori, Y.; Saito, N.; Ueda, K. *Phys. Rev. A* **2011**, *83*, 053415.
- (13) Miteva, T.; Chiang, Y.-C.; Kolorenč, P.; Kuleff, A. I.; Cederbaum, L. S.; Gokhberg, K. J. Chem. Phys. 2014, 141, 164303.
- (14) Travnikova, O.; Marchenko, T.; Goldsztejn, G.; Jänkälä, K.; Sisourat, N.; Carniato, S.; Guillemin, R.; Journel, L.; Céolin, D.; Püttner, R.; Iwayama, H.; Shigemasa, E.; Piancastelli, M. N.; Simon, M. *Phys. Rev. Lett.* **2016**, *116*, 213001.
- (15) Björneholm, O.; Federmann, F.; Fössing, F.; Möller, T. *Phys. Rev. Lett.* **1995**, *74*, 3017–3020.
- (16) Miteva, T.; Wenzel, J.; Klaiman, S.; Dreuw, A.; Gokhberg, K. Phys. Chem. Chem. Phys. 2016, 18, 16671–16681.

- (17) Nordlund, D.; Ogasawara, H.; Bluhm, H.; Takahashi, O.; Odelius, M.; Nagasono, M.; Pettersson, L. G. M.; Nilsson, A. *Phys. Rev. Lett.* **2007**, *99*, 217406.
- (18) Ottosson, N.; Odelius, M.; Spångberg, D.; Pokapanich, W.; Svanqvist, M.; Öhrwall, G.; Winter, B.; Björneholm, O. J. Am. Chem. Soc. 2011, 133, 13489–13495.
- (19) Céolin, D.; Ablett, J.; Prieur, D.; Moreno, T.; Rueff, J.-P.; Marchenko, T.; Journel, L.; Guillemin, R.; Pilette, B.; Marin, T.; Simon, M. J. Electron Spectrosc. Relat. Phenom. 2013, 190, Part B, 188 192.
- (20) Rueff, J.-P.; Ablett, J. M.; Céolin, D.; Prieur, D.; Moreno, T.; Balédent, V.; Lassalle-Kaiser, B.; Rault, J. E.; Simon, M.; Shukla, A. J. Synchrotron Rad. 2015, 22, 175–179.
- (21) Faubel, M.; Schlemmer, S.; Toennies, J. P. Z. Phys. D 1988, 10, 269–277.
- (22) Ohtaki, H.; Radnai, T. Chem. Rev. 1993, 93, 1157–1204.
- (23) Soper, A. K.; Weckström, K. Biophys. Chem. 2006, 124, 180 191.
- (24) Ma, H. Int. J. Quant. Chem. **2014**, 114, 1006–1011.
- (25) Krishnan, R.; Binkley, J. S.; Seeger, R.; Pople, J. A. J. Chem. Phys. 1980, 72, 650–654.
- (26) Blaudeau, J.-P.; McGrath, M. P.; Curtiss, L. A.; Radom, L. J. Chem. Phys. 1997, 107, 5016–5021.
- (27) Frisch, M. J. et al. Gaussian 09 Revision D.01. Gaussian Inc. Wallingford CT 2009.
- (28) Lee, H. M.; Kim, J.; Lee, S.; Mhin, B. J.; Kim, K. S. *J. Chem. Phys.* **1999**, *111*, 3995–4004.
- (29) Ge, L.; Bernasconi, L.; Hunt, P. Phys. Chem. Chem. Phys. **2013**, 15, 13169–13183.
- (30) Gora, R. W.; Roszak, S.; Leszczynski, J. Chem. Phys. Lett. **2000**, 325, 7 14.
- (31) Schirmer, J. Phys. Rev. A **1982**, 26, 2395–2416.

- (32) Barth, A.; Schirmer, J. J. Phys. B At. Mol. Opt. Phys. 1985, 18, 867.
- (33) Cederbaum, L. S.; Domcke, W.; Schirmer, J. Phys. Rev. A 1980, 22, 206–222.
- (34) Barth, A.; Cederbaum, L. S. Phys. Rev. A 1981, 23, 1038–1061.
- (35) Wenzel, J.; Wormit, M.; Dreuw, A. J. Comp. Chem. 2014, 35, 1900–1915.
- (36) Wenzel, J.; Wormit, M.; Dreuw, A. J. Chem. Theory Comput. 2014, 10, 4583-4598.
- (37) Wormit, M.; Rehn, D. R.; Harbach, P. H.; Wenzel, J.; Krauter, C. M.; Epifanovsky, E.; Dreuw, A. Mol. Phys. **2014**, 112, 774–784.
- (38) Shao, Y. et al. Mol. Phys. 2015, 113, 184-215.
- (39) McLean, A. D.; Chandler, G. S. J. Chem. Phys. 1980, 72, 5639–5648.
- (40) Krause, M. O.; Oliver, J. H. J. Phys. Chem. Ref. Data 1979, 8, 329-338.
- (41) Brooks, B. R.; Laidig, W. D.; Saxe, P.; Handy, N. C.; Schaefer III, H. F. Phys. Scr. 1980, 21, 312.
- (42) Brooks, B. R.; Schaefer, H. F. J. Chem. Phys. 1979, 70, 5092–5106.
- (43) Schmidt, M. W.; Baldridge, K. K.; Boatz, J. A.; Elbert, S. T.; Gordon, M. S.; Jensen, J. H.; Koseki, S.; Matsunaga, N.; Nguyen, K. A.; Su, S.; Windus, T. L.; Dupuis, M.; Montgomery, J. A. J. Comp. Chem. 1993, 14, 1347–1363.
- (44) Kaufmann, K.; Baumeister, W.; Jungen, M. J. Phys. B At. Mol. Opt. Phys. 1989, 22, 2223.
- (45) Mosnier, J.-P.; Kennedy, E. T.; van Kampen, P.; Cubaynes, D.; Guilbaud, S.; Sisourat, N.; Puglisi, A.; Carniato, S.; Bizau, J.-M. *Phys. Rev. A* **2016**, *93*, 061401.
- (46) Russek, A.; Mehlhorn, W. J. Phys. B At. Mol. Opt. Phys. 1986, 19, 911.

- (47) Guillemin, R.; Sheinerman, S.; Püttner, R.; Marchenko, T.; Goldsztejn, G.; Journel, L.; Kushawaha, R. K.; Céolin, D.; Piancastelli, M. N.; Simon, M. Phys. Rev. A 2015, 92, 012503.
- (48) Tchaplyguine, M.; Kivimäki, A.; Peredkov, S.; Sorensen, S. L.; Öhrwall, G.; Schulz, J.; Lundwall, M.; Rander, T.; Lindblad, A.; Rosso, A.; Svensson, S.; Mårtensson, N.; Björneholm, O. J. Chem. Phys. 2007, 127, 124314.
- (49) Sugiura, C. J. Chem. Phys. 1982, 77, 681-682.
- (50) Shadle, S. E.; Hedman, B.; Hodgson, K. O.; Solomon, E. I. J. Am. Chem. Soc. 1995, 117, 2259–2272.
- (51) Hertlein, M. P.; Adaniya, H.; Amini, J.; Bressler, C.; Feinberg, B.; Kaiser, M.; Neumann, N.; Prior, M. H.; Belkacem, A. Phys. Rev. A 2006, 73, 062715.
- (52) Céolin, D.; Marchenko, T.; Guillemin, R.; Journel, L.; Kushawaha, R. K.; Carniato, S.; Huttula, S.-M.; Rueff, J. P.; Armen, G. B.; Piancastelli, M. N.; Simon, M. Phys. Rev. A 2015, 91, 022502.
- (53) Goldsztejn, G.; Marchenko, T.; Püttner, R.; Journel, L.; Guillemin, R.; Carniato, S.; Selles, P.; Travnikova, O.; Céolin, D.; Lago, A. F.; Feifel, R.; Lablanquie, P.; Piancastelli, M. N.; Penent, F.; Simon, M. Phys. Rev. Lett. 2016, 117, 133001.
- (54) Björneholm, O.; Nilsson, A.; Sandell, A.; Hernnäs, B.; Mrtensson, N. Phys. Rev. Lett. 1992, 68, 1892–1895.
- (55) Karis, O.; Nilsson, A.; Weinelt, M.; Wiell, T.; Puglia, C.; Wassdahl, N.; Mårtensson, N.; Samant, M.; Stöhr, J. Phys. Rev. Lett. 1996, 76, 1380–1383.

Graphical TOC Entry

