CORE LEVEL EXCITATION OF ATOMIC AND MOLECULAR CLUSTERS


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ABSTRACT

Core level excitation of weakly bonded atomic and molecular species is reported. Excitations into unoccupied orbitals with valence or Rydberg character as well as into the corresponding K- and L-continua are compared with the isolated and condensed species. Core level excitation of clusters is also accompanied with extensive fragmentation of singly and doubly charged clusters. Coincidence techniques such as photoelectron-photoion-coincidences (PEPICO) and photoion-photoion-coincidences (PIPICO) are used to study the photochemical fragmentation pathways of core excited clusters. Kinetic energy releases related to charge separation of cluster dications (Coulomb explosion) are derived from PIPICO spectra. The fragmentation energetics is discussed in relation to cluster dication threshold energies.

INTRODUCTION

The field of core excitation processes of clusters is relatively unexplored. Most studies that have been performed in the energy regime of core level excitation (E>150 eV) have concentrated on isolated atoms or molecules in the gas phase and condensed species, i.e. solids or adsorbates. Valuable information is obtained from these studies, such as energy positions and relative intensities of resonant processes above or below the corresponding ionization energies. It has been found that the shapes of inner shell absorption spectra of isolated atoms differ from those of the corresponding condensed species. In contrast to valence transitions of core excited molecular species (1s→π*) show typically the same energy positions in the gas and condensed phase. However, intensity changes of discrete resonances relative to the corresponding ionization continua are observed. Recently, we have reported progress in the field of core level excitation of isolated clusters in the gas phase. It has been shown that in the case of argon clusters the cation yield spectra of mass selected clusters differ considerably from those of the isolated cation in the L3/L2 excitation regime (240-300 eV). Cluster specific resonances are observed, which are more similar to the absorption spectrum of the solid rather than to the atom. Broad features are observed in the corresponding L1/L2 continuum, which are assigned as single scattering (EXAFS) processes in clusters. More recent results in this field show that also above the L1 edge similar EXAFS structure is observed (320-500 eV), which gives further confidence in the given assignments. We have also studied fragmentation pathways of core excited argon cluster dications. With the photoion-photoion-coincidence (PIPICO) technique it was shown that doubly charged clusters undergo asymmetric fission, leading to series of products, such as monomer cations, dimer cations, and trimer cations in coincidence with cluster cations. The kinetic energy releases are significantly lower than expected from a simple electrostatic picture which is in agreement with findings for collision induced fragmentation carbon dioxide cluster dications, where internal excitation has been assumed to be responsible for the low kinetic energy releases.

In this paper we report recent progress in the field of core level excitation in atomic and molecular clusters in the energy regimes of element K- and L-edges: argon 2p (240-280 eV), carbon 1s (270-330 eV), and oxygen 1s (520-570 eV). Besides photoion yield curves of mass selected cluster cations, which provide spectroscopic information, we have also studied fragmentation reactions of core excited cluster monocations and dications. Threshold energies of unstable argon cluster dications are measured, which are used to interpret the energetics of core excited dication decay.

EXPERIMENTAL

The experiments were performed at the electron storage ring BESSY. The high energy TGM-II beam line allows the dispersion of soft x rays between 150 eV and 600 eV. The energy resolution (∆E) is obtained from the full width at half maximum signal of the argon (L1, 4s) ion signal at 244.390 eV. The experiments were performed typically with an energy resolving power (E/∆E) of 300-500.

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The TGM-7 beam line is used for threshold experiments of direct cluster double ionization. This monochromator disperses synchrotron radiation between 7 eV and 150 eV. The experiments are performed with a 950 l/mm grating with an optimized dispersion range between 20-120 eV. Higher order light and stray light was filtered by an aluminum foil.

The experimental setup shown in Fig. 1 is similar to that reported earlier. A time of flight mass spectrometer (TOF-MS) is used for cation detection. The design of the mass spectrometer involves conventional Wiley/McLaren space focussing. Under typical experimental conditions a mass resolution (M/ΔM) of 125 is obtained for Ar⁺ (m/z=40). The spectrometer is operated with moderate acceleration fields in the ionization region (100 to 200 V/cm).

Opposite to the TOF-MS a threshold photoelectron spectrometer is mounted. This device can be replaced by a total yield detector.

With this setup we have measured mass spectra at resonant excitation energies, total cation and electron yield curves, photoionization efficiency curves of mass selected cations, threshold photoelectron (TPES) spectra, photoelectron-photoion-coincidence (PEPICO) spectra, and PIPICO spectra. Mass spectra were recorded by the use of pulsed extraction of cations into the mass spectrometer, whereas PEPICO and PIPICO spectra were obtained with static voltages. Further experimental details are given in ref. 7.

**Fig. 1. Experimental setup**

The stagnation conditions for the supersonic expansion are similar to those reported earlier. We have extended the stagnation temperature range by cooling the gas inlet line and the nozzle down to 90 K with liquid nitrogen. From scaling laws and correlations to average cluster sizes we expect for T₀ =300 K an average cluster size (N) of 2 (p₀= 2 bar) to 10 (p₀= 5 bar), depending on the expansion conditions. Kinetic energy releases (KER) for charge separation in cluster dications are calculated according to the formulas given in ref. 9. The samples were of commercial quality and were used without further purification.
RESULTS AND DISCUSSION

I. ATOMIC CLUSTERS

Fig. 2 shows the photoion-photoion-coincidence (PIPCICO) spectrum of argon clusters recorded at 248 eV. This energy corresponds to 3d (3/2) exciton state in clusters and the solid. Three series of charge separation product channels are identified: \( \text{Ar}^+/\text{Ar}_n^+ \), \( \text{Ar}_2^+/\text{Ar}_n^+ \) and \( \text{Ar}_3^+/\text{Ar}_n^+ \). These series of charge separation show that asymmetric fission of argon cluster dications is dominant. The signal at \( \Delta t=0 \) is shown to be due to symmetric charge separation processes. However, forthcoming photoelectron-photoion-photoion-coincidence experiments will unequivocally show the fraction of symmetric versus asymmetric charge separation in core excited argon clusters.

The series of coincidences involving cation pairs of dimers and clusters is stronger than the other ones. This is consistent with the high stability of the argon dimer cation. Its formation is a dominant process in single ionization of argon clusters. Recent experimental results from mass resolved molecular beam scattering show that small neutral clusters form preferentially monomer and dimer cations after electron impact ionization. This is in agreement with photofragmentation studies of mass selected cluster cations, where for species smaller than \( \text{Ar}_{15}^+ \) the major fragments \( \text{Ar}_2^+ \) and \( \text{Ar}_3^+ \) are detected. From photoelectron spectra of argon cluster beams and DIM calculations the chromophors \( \text{Ar}_3^+ \) (for small clusters) and \( \text{Ar}_{13}^+ \) (for large clusters) are proposed.

Asymmetric fission of molecular cluster dications has been studied with electron impact ionization techniques. It is reported that more than 90% of the mass is retained in the observed fragments, whereas stable doubly charged argon clusters undergo a loss of one or two single atoms, where no Coulomb explosion is observed.

Fig. 2. (a) Photoion-photoion-coincidence (PIPCICO) spectrum of argon clusters at 248 eV excitation energy, (b) shows a detailed view of the \( \text{Ar}^+/\text{Ar}_2^+ \) signal. The energy scale refers to the two body dissociation \( \text{Ar}_3^+ \rightarrow \text{Ar}^+/\text{Ar}_2^+ \). Experimental conditions: \( p_0=4.7 \text{ bar}, T_0=215 \text{ K} \).
(3.9(1) Å), the solid (3.75 Å), and matrices (3.7 Å).

The time separation of both maxima of the PIPICO signal corresponds to a KER of 0.7(2) eV. It is remarkable that the shape of the PIPICO signal remains nearly unchanged if the average cluster size is decreased.

A likely explanation for the low KER's can be found by considering a complex charge separation mechanism in cluster dications: According to the double ionization model for solids a two step process leads to double ionization, where first step is the photoionization of one atom within the cluster followed by subsequent electron impact ionization of a neighboring atom. Immediate Coulomb explosion will result in a high KER, which accounts for the widest part of the PIPICO signal. As a competing process charge separation within the clusters prior to Coulomb explosion is likely. Some of the total energy in the cluster dication may be consumed by evaporation of neutrals or can be stored as internal energy of the singly charged fragments. Similar results are reported by Gotts and Stace for collision induced fragmentation of carbon dioxide cluster dications. The difference between the predicted KER's for Coulomb repulsion and the experimental results are interpreted in terms of simultaneous internal excitation.

In an oversimplified picture of purely electrostatic charge separation the low KER can be understood in terms of medium sized cluster dications, such as Ar$_7^{2+}$ to Ar$_8^{2+}$, where a charge separation distance of 20.5 Å appears to be in a reasonable range, if non-compact cluster structures are assumed to exist prior to charge separation.

Related to the KER's in clusters is the question of energy position of the dissociating dication state. The asymptote of the Ar$_7^{2+}$/Ar$_8^{2+}$ product channel is 30 eV as calculated from thermodynamical reference data. With the experimental KER of 3.8 eV one obtains 33.8 eV as a lower limit for the dissociating dication state of Ar$_5^{2+}$

Experimental efforts have been made to measure thresholds of unstable cluster dications in order to correlate the above estimated energy of dissociative dication states with experimental data. Since small cluster dications are unstable we have used the PIPICO-technique to investigate both the thresholds of the Ar$_7^{2+}$/Ar$_8^{2+}$ and Ar$_5^{2+}$/Ar$_6^{2+}$ PIPICO signals as a function of average cluster size and photon energy. It is observed that the thresholds of both product channels depend on the average cluster size. The results are listed in Tab. I:

**Table I: Threshold energies for double ionization in atomic, clustered and solid argon. The average cluster sizes are estimated to the references listed in ref. 5. The error limits for cluster threshold determination are 0.3 eV.**

<table>
<thead>
<tr>
<th>species</th>
<th>threshold energy / eV</th>
<th>reference</th>
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<tbody>
<tr>
<td>atom</td>
<td>43.4</td>
<td>15a</td>
</tr>
<tr>
<td>cluster</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\bar{N}=3$</td>
<td>36.0</td>
<td>this work</td>
</tr>
<tr>
<td>$\bar{N}=4$</td>
<td>35.5</td>
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<tr>
<td>$\bar{N}=6$</td>
<td>35.0</td>
<td>this work</td>
</tr>
<tr>
<td>$\bar{N}=10$</td>
<td>33.5</td>
<td>this work</td>
</tr>
<tr>
<td>$\bar{N}=20$</td>
<td>32.8</td>
<td>this work</td>
</tr>
<tr>
<td>Ar$_{101}$</td>
<td>32</td>
<td>15b</td>
</tr>
<tr>
<td>solid</td>
<td>28.6</td>
<td>15c</td>
</tr>
</tbody>
</table>

The data in Tab. I show that the double ionization thresholds of the solid and the atom differ by nearly 15 eV. The value for solid argon (28.6 eV) is obtained from recent photoelectron-photoelectron-coincidence measurements, which is in good agreement with the double ionization energy of the solid. From electron impact mass spectrometry a threshold for the stable cluster dication Ar$_{101}^{2+}$ is reported. The thresholds of cluster dications undergoing Coulomb explosion show values that fall between the stable cluster dication (Ar$_{101}^{2+}$) and the atomic dication. The estimated lower limit of the Ar$_5^{2+}$ dissociative state (33.8 eV) agrees well with the appearance energies of small cluster dications ($\bar{N}<6$).

**II. MOLECULAR CLUSTERS**

Fig. 3 shows photoionization efficiency curves of CO and its dimer in the energy regime of the carbon K-edge as well as the C$^+$ and dimer spectra at the oxygen K-edge. All spectra are dominated by the C (1s)$\rightarrow$$\pi^*$ transition (287.4 eV), and O (1s)$\rightarrow$$\pi^*$ transition (534.11 eV), respectively.
In contrast to the cluster size dependent energy positions of Rydberg states below the argon 2p edges, we do not observe energy shifts for this valence transition in clusters. This is in agreement with photon stimulated desorption of solid carbon monoxide, where no shift relative to the gas phase spectrum is observed. The most interesting difference in spectral shape is the decreased intensity of both 1s→π* bands relative to the K-continuum intensities and changes in spectral shape in both Rydberg regimes. The x-ray absorption spectrum of CO shows under high resolution conditions evidence for several Rydberg series below the carbon K-ionization continuum. In the dimer spectrum we find an onset of intensity at 292 eV without any fine structure, a shoulder at 293 eV, and an increase in intensity up to 305 eV, which corresponds to the energy of the C(1s)→σ* transition. The spectral shape of the dimer cation mimicks that of cations desorbing from the solid (photon stimulated desorption (PSD)). The ion yield curve of CO shows also weak contributions of cluster ion fragmentation above 292 eV. In gas phase CO one finds that there is no parent cation intensity above the carbon K-ionization energy, since double and multiple ionization dominate this energy regime. This shows that cluster fragmentation leads to CO as a final product of dissociative ionization processes. This is also consistent with the PEPICO spectrum of CO clusters recorded at 305 eV, where a considerable CO signal is observed (cf. Fig. 4a). Another interesting difference of the cluster cation yield is a comparison with coordinated CO in transition metal carbonyls. It is found that quenching of Rydberg states is evident as observed for van der Waals clusters of CO. Due to metal 3d level backbonding the intensity of the C(1s)→π* transition is decreased in organometallics.

Similar differences in spectral shape are observed for oxygen K-electron excitation, where a considerably stronger K-continuum is observed relative to the O (1s)→π* resonance. This is in good agreement with PSD of cations from the solid. Relative changes in intensity of π* resonances vs. K-continuum intensities have been rationalized for the solids in terms of a delayed onset behavior. These phenomena are consistent with reneutralization or screening effects by the solid. It is interesting to note that the spectral shape of the C fragment shows a considerably stronger O (1s)→π* resonance compared to the dimer cation. This is consistent with the spectra shown in ref. 17, indicating that C is a product of unclustered CO fragmentation.
The photochemical behavior of carbon monoxide clusters are studied with the PEPICO and PIPICO techniques. The PEPICO spectrum of CO clusters is shown in Fig. 4a. The dominant peak in the spectrum is the dimer cation, which is one of the preferred fragmentation products of CO cluster dications, as deduced from the width of the PEPICO signal. Besides the dimer signal we find a relative strong monomer intensity as well as singly or multiply charged atomic fragments. The latter ones show unchanged intensity upon clustering, indicating that they are not involved in cluster dissociation pathways.

The PIPICO spectrum of CO clusters shows besides the well-known molecular charge separation channels (C⁺O⁻) several coincidences corresponding to cluster dication decays. Most products involve asymmetric charge separation into entire molecular and cluster units. The widest portion of the CO⁺/(CO)₂⁺ PIPICO channel corresponds to a KER of 2.6(3) eV for a two body dissociation, whereas the peak maximum gives 0.9(2) eV. In terms of charge separation distances in an electrostatic (Coulomb explosion) picture this corresponds to 5.5 Å and 17 Å, respectively. The lower value is twice the molecular diameter, which is maximum charge separation distance in a dimer. The higher value corresponds to a low KER which is probably due to internal excitation of the cluster fragments, as discussed for argon clusters.

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QUESTION

BAER - I am surprised that you obtained similar bond distances for the Ar$_2$ clusters as for the Ar$_n$ clusters from your NEXAFS data. What about multiple scattering?

RUHL - The cluster EXAFS analysis shows that an Ar-Ar distance of 3.9 ± 0.1 Å is obtained. This value is slightly higher than values obtained for the solid (3.75 Å) or matrices (3.7 Å). An obvious reason for the slightly increased Ar-Ar distance in small argon clusters (with N = 10) can be found in the spherical (icosahedral) structure of argon clusters. In contrast to this, the solid is known to exist in a f.c.c. lattice. Further EXAFS experiments in the energy regime of Ar(1s) excitation are needed to confirm the results obtained at the Ar(2p) - edge (250 eV). The advantage of Ar(1s) - EXAFS will be, that the energy space is not limited as in the case of Ar(2p) - EXAFS, where the Ar(2s)-edge disturbs the 2p EXAFS above 330 eV. Another advantage of Ar(1s)-EXAFS will be that only one threshold energy has to be considered in the EXAFS analysis. Our EXAFS analysis has shown, that the oscillations that are observed in photoion yield curves of argon clusters above the 2p-and 2s-edges can be described with a single scattering mode, whereas the resonance centered at approximately 256 eV is probably due to multiple scattering.

DUIJARDIN - Experimental question about the PIPOCO curves you showed. They have a non zero background. So it seems that the false coincidences have not been substituted. Could this change some of the results?

RUHL - The false coincidence background is extremely weak, so that is was not necessary to subtract any background. Therefore, we do not think the results are affected by false (or random) coincidences.

MENZEL - Two questions related to the comparison of clusters and solids:

1. In solid Argon, surface and bulk 2p resonances can be distinguished by using the detection of different ion signals such as Ar$^+$ and Ar$_2^+$ (see e.g. D. Menzel, Appl. Phys., 1990 and work cited there) or by detecting decay electrons at different exit angles (G. Rocker et al., to be published in Phys. Rev. B). Can a similar distinction be made in the case of clusters?
2. Again in solid Argon, evidence for a bi-exciton excitable by electrons (P. Feulner et al., PRL June 1991) or even by simple photons (P. Feulner et al., to be published) has been found. Is there any evidence that such an excitation also exists in Ar clusters?

RUHL -

1. Different cation channels will not give a distinction between different processes. However if the beam expansion conditions are changed, the NEXAIFS structure at the Ar(2p)-edges changes its shape independent on the cation channel. We have analyzed the different components of spectral features in the energy regime of the 4s Rydberg/exciton state (244-246eV) using a total electron detection method (E. Ruhl et al., to be published). When the average cluster size is changed we observe how the individual contributions of small clusters (surface excitons), medium sized clusters, and large clusters (bulk excitons) change their relative intensity.
2. We have found no evidence of these processes in clusters so far.

WONG - Was the Fourier transform of the Ar EXAIFS phase corrected to give the Ar-Ar distance you quoted? If so, which (whose) phase shift value did you use?

RUHL - The EXAFS analysis was done by the use of Mc Kale parameters. Further details can be found in: E. Ruhl et al., Chem. phys. lett. 178, 558 (1991).