Thermal quenching of electronic shells and channel competition in cluster fission

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Experimental and theoretical studies of fission of doubly-charged Li, Na, and K clusters in the low fissility regime reveal the strong influence of electronic shell effects on the fission products. The electronic entropy controls the quenching of the shell effects and the competition between magic-fragment channels, leading to a transition from favored channels of higher mass symmetry to the asymmetric channel involving the trimer cation at elevated temperatures.

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Explorations of factors controlling the stability and the collapse or fragmentation of finite systems range from early studies of hydrodynamical instabilities and droplet formation [1] to studies of fission and fragmentation of atomic nuclei [2], and of atomic and molecular free clusters and supported nanostructures [3,4]. Here we focus on the charge instability and fission of doubly charged cationic metal clusters, that is $M^+_N \rightarrow M^+_P + M^{-}_N - P$. Such instabilities in macroscopic fluid systems have been studied by Rayleigh [1], and adaptations of Rayleigh’s model (RM) for the description and analysis of nuclear [2] and cluster fission [5] are referred to as the Liquid Drop Model (LDM). In the LDM, the clusters are viewed as charged classical liquid drops whose shape and dynamics are controlled by the competition between the repulsive Coulomb ($E_{\text{Coulomb}}$) and the binding surface ($E_{\text{surface}}$) energies [6].

Various fission regimes may be classified by the fissionability parameter $X = E^{\text{sph}}_{\text{Coulomb}} / 2E_{\text{surface}}^{\text{sph}}$, where the superscript “sph” indicates that the equilibrium shapes of the LDM are spherical. For $X \geq 1$, spontaneous barrierless fission is predicted and observed [7,8]; this is also the regime studied originally by Rayleigh who predicted a mass-symmetric fission mode (see also Ref. [7]). In the $X < 1$ regime, which is the focus of our paper, the fissioning system must overcome a barrier. For atomic clusters in this regime, tunneling is suppressed, unlike the case of nuclei, and fission requires thermal activation. From $X_{\text{cr}} = 1$, the RM critical size for barrierless fission is $N^*_{\text{cr}} = z^2e^2 / (16\pi r_s^2\sigma)$, where $r_s$ is the Wigner-Seitz radius and $\sigma$ is the surface-energy; for doubly charged ($z = 2$) Li, Na, and K clusters, one has $N^*_{\text{cr}} \approx 10$. Thus, with the exception of the smallest ones, fissioning clusters must be hot [9] in order to overcome the fission barrier.

For the low fissility $X < 1$ regime, the LDM predicts the dominance of a strongly mass asymmetric fission process, as deduced from the finite-temperature [6,10] $Q'_F = F^+_P + F^{-}_N-P - F^2_N$ values, that express the free-energy balance for the possible $(P,N-P)$ fission channels. Such asymmetric fission has been observed in experiments for Na$_{2N}^+$ with $N \approx 24$ (i.e., near the appearance size, see below), where the preferred fission channel involved the “magic” Na$_3^+$ fragment [9]. The prevalence of this channel, compared to other asymmetric ones in the neighborhood of the LDM minimum, originates from quantum mechanical shell effects [11].

In this paper we present experimental and theoretical results, pertaining to cluster fission for $X << 1$, which cannot be described by the classical RM, demonstrating instead the dominance of quantum-size effects, i.e., electronic shell contributions [6]. In particular, we study alkali-metal (Li, Na, and K) clusters with $N = 24, 26, 28$, and 30. These systems correspond to the lowest observable fissility regime with $N >> N_{\text{cr}}^{\text{RM}}$; e.g., for $N = 30$ one has $X \approx 0.25$ for the doubly charged alkali clusters. Moreover, this range of sizes is particularly suitable for explorations of shell effects, since it includes cases where the clusters may fission into doubly “magic” fragments, i.e., $M_{21}^{2+} \rightarrow M_{21}^{-} + M_2^{+}$ and $M_{30}^{-} \rightarrow M_{21}^{-} + M_{9}^{+}$.

We find that, in addition to $M_{21}^{-}$, the more mass symmetric magic fission channels (involving the magic fragments $M_9^+$ and $M_{21}^+$) do appear and compete with the channel involving the trimer (even becoming the preferred channels in several instances). Furthermore, from comparisons between the experiments and calculations based on the finite-temperature shell-correction method (FT-SCM) [12], we find that the channel distributions are greatly influenced by thermal quenching of the shell effects due to the increase (compared to $T = 0$) of the electronic entropy; this quenching results from thermal promotion of electrons to unoccupied single-particle levels which smears out the shell structure. The more mass-symmetric fission channels are favored at lower temperatures, whereas the asymmetric ones (e.g., involving the $M_9^+$ fragment) dominate as $T$ increases. These trends reflect the more pronounced (quenching) effect of the electronic entropy on the shell-stabilization of the larger magic sizes (e.g., $M_9^+$ and $M_{21}^+$).

In the experiment [13], a distribution of relatively large
neutral clusters is formed by a gas aggregation source. The clusters are ionized, photo-excited and warmed by a 15 ns laser pulse (Nd-YAG laser at $h\nu = 3.50$ eV). A rapid sequential evaporation follows the warming of the clusters for about 1 µs (i.e., the residence time in the ionizing region). This results in a charged cluster distribution shifted down to smaller sizes, called an “evaporative ensemble” (EE) [13]. The cluster temperature associated with the EE is determined [13] by the cluster dissociation energy and by the experimental time window, i.e., the residence time in the ionizing/heating region. Under our experimental conditions, the temperatures of the EE’s of lithium, sodium and potassium clusters are $700\pm100$ K, $400\pm100$ K and $300\pm100$ K, respectively.

The EE of ionized clusters enters a tandem time-of-flight (TOF) mass spectrometer. Parents of interest are mass selected in the first TOF region. Because they are at a given temperature, subsequent unimolecular dissociation of these parents takes place and the resulting fragments are mass analyzed by the second TOF device. Enhanced optimization of the electrostatic potentials in the current experiments has led to significant increases in both the mass and energy resolutions. In particular, the latter allows us to resolve the kinetic energy released during the fission process (see the splitting of the peak of some of the fission fragments in Fig. 1).

For cluster sizes in the vicinity of the appearance size (see below), the unimolecular dissociation of doubly charged clusters portrays the competition between the fission (1a) and evaporation (1b) processes:

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\begin{align*}
M_N^{2+} &\rightarrow M_P^{+} + M_{N-P}^{+} \quad (1a) \\
&\rightarrow M_{P'}^{+} + M_{N-P'}^{+}, \quad P' = 1 \text{ or } 2.
\end{align*}
$$

We focus here on the fission decay channels. The explored size domain is bounded by two values, $N_-$ and $N_+$. The lower value $N_-$ is determined by the appearance size $N_{\text{app}}$ ($N_{\text{app}} = 24$ for Li$_N^{2+}$ and Na$_N^{2+}$ clusters, and 19 for K$_N^{2+}$ clusters [4]), below which fission is the dominant dissociation process [see Eq. (1a)]. Below this size no doubly charged clusters are present in the mass spectra, since their fast fission process prevents their observation during the experimental time window. $N_+$ is determined by the loss of the signal corresponding to the fission process, resulting from the increase of the inner part of the fission barrier with the size of the cluster relative to the essentially constant monomer evaporation energy. Fig. 1 summarizes the results obtained for the three alkali metals in the same size range.

Although we have recorded the fragmentation spectra for all the sizes $24 \leq N \leq 30$, only results for clusters with an even number of atoms are shown (due to pairing they exhibit simpler features). In our experimental setup, only the heavy fragments $M_P^+$ with $P > N/2$ are detected. The main features are as follows: For the lowest parent size ($N = 24$) and for all three elements, only one fission channel is present, i.e., the doubly magic $M_P^+ + M_{21}^+$. Increasing the parent size, however, results in differentiation between the three elements regarding the observed fission channels. A larger parent size (i.e., $N \geq 26$) opens the channels of heavier fragments and this trend is more pronounced for potassium than for lithium. In particular, the trimer appears to be the dominant product in the Li fission, while $M_{21}^+$ is the dominant one for K, with Na exhibiting an intermediate behavior. That is, the asymmetry of the fission process is reduced for the K clusters as compared to the Na and Li ones.

We have performed FT-SCM calculations of the $Q_T$-values for the various fission channels for all three alkali-metal sequences at the corresponding experimental temperatures. Since for each species the Coulomb repulsion (QR) at the scission point (measured by the energy release) is approximately constant for all fission channels in the size-range considered here ($X \ll 1$), the effective inner part of the fission barrier can be approximated by adding the calculated $Q_T$-values to the QR term. Thus the ordering of the fission channels can be directly in-

![FIG. 1. Fission channels recorded in TOF measurements for Li$_N^{2+}$, Na$_N^{2+}$, and K$_N^{2+}$ clusters, with $N = 24$, 26, 28, and 30. In each case, the peak on the right (marked by an arrow) corresponds to the parent cluster and the ones on the left of it correspond to the larger fission fragments. Fragment labelings are determined from the voltage setup. Peak heights are in arbitrary units. The splitting of the TOF ion-fragment peaks are due to the kinetic energy released during fission, which is of the order of 1 eV (see, e.g., the case $N = 26$).](image)
ferred from the $Q^T$-values.

The theoretical zero-temperature $Q$-values provide an inadequate description of the experimental data. For example, the $Q$-values for $Li^{2+}_{2N}$ (plotted in Fig. 2) cannot explain the observed trend (see Fig. 1, left column) of strong asymmetric fission, with the $Li^+_{2N}$ being the preferred channel for all parents $N = 24, 26, 28,$ and $30$ [14]. In particular, the $T = 0$ theoretically deduced preferred fission channels for $N = 26$ and $N = 30$ are $(5,21)$ and $(9,21)$, respectively, in contradiction with the measurements (see Fig. 1). In contrast, the theoretical $Q^T$-values (Fig. 3) agree well with the experiment. Specifically, for Lithium the calculations at $T = 700$ K show that $Li^+_{2N}$ is indeed the preferred channel for all parents; only for $Li^{2+}_{30}$, the $Li^{2+}_{2N}$ channel becomes competitive, remaining however less abundant than the one involving the trimer.

We reiterate that in Fig. 3 there is a clear tendency toward opening the channels involving the heavier magic fragments ($P = 9$ and $P = 21$), both when going up vertically from the lighter to the heavier parents (from $N = 24$ to $N = 30$), as well as when going horizontally from left to right (i.e., from Li to K, with Na representing an intermediate case).

The physics underlying this agreement between the experiment and the FT-SCM results can be revealed by an examination of the influence of the electronic entropy on the various fission channels ($P, N − P$) as a function of $P$. For brevity, we limit our discussion to the case of the $K^{2+}_{30}$ parent; the conclusions, however, extend to all three species and the other parent sizes. Figure 4 displays the FT-SCM $Q^T$-values for $K^{2+}_{30}$ at five temperatures (200 K to 1000 K). At the lowest temperature, the $(9,21)$ channel is the preferred one; however, an increase in $T$ is accompanied by quenching of the shell effects; i.e., at high temperature (1000 K) the shell effects have almost vanished, and the $Q^T$ curve exhibits a smooth shape similar to that of the LDM (although with a steeper variation).

Most important for the competition between the various magic fragments, however, is the fact that the electronic-entropy quenching of the shell effects is non-uniform, with the higher occupied shells of the heavier fragments being influenced the most as $T$ increases, since the energy spacing between the major electronic shells, $\hbar \omega_0(N) = 49 eV \cdot a_0^2/(\gamma_{2N} N^{1/3})$, decreases with larger $N$. Consequently, a higher $T$ favors the trimer ($M^{3+}_{2N}$) channel over those involving the magic $M^{2+}_{N}$ and $M^{2+}_{2N}$ fragments. This happens even in the case of $N = 30$ where the competing $(9,21)$ channel is doubly magic. Note that, unlike the $T = 700$ K experimental temperature of Li, the temperatures of 300 K for Potassium and 400 K for Sodium are low enough so that the heavier-fragment channels retain their $T = 0$ original prevalence for all three parent sizes $N = 26, 28,$ and $30$. 

FIG. 2. Theoretical SCM zero-temperature energetics ($Q$-values) for the processes $Li^{2+}_{2N} \rightarrow Li^+_{2N} + Li^+_{P}$. $N = 24, 26, 28,$ and $30$. The dashed lines correspond to LDM calculations. In each case, the horizontal axis corresponds to various SCM channels $2 \leq P \leq N − 2$.

FIG. 3. Finite-temperature SCM results ($Q^T$-values) for the fission process $M^{2+}_{N} \rightarrow M^{3+}_{N−P}$ for $Li^{2+}_{N}$, $Na^{2+}_{N}$, and $K^{2+}_{N}$ with $N = 24, 26, 28,$ and $30$. The corresponding temperatures (in degrees Kelvin), taken from the experiments, for each species are shown at the top. The dashed lines correspond to LDM calculations. In each case, the horizontal axis corresponds to various SCM channels $2 \leq P \leq N − 2$. The length of the vertical bar in each frame corresponds to 0.4 eV.
Further insight can be gained by examining the ratio \( f_{sh}(N) = \Delta F_{sh}(N, \tau)/\Delta F_{sh}(N, 0) \) of the free-energy shell-correction term at \( \tau = 2\pi^2 k_B T/h\omega_0(N) \) over the shell correction term at \( \tau = 0 \). For a schematic single-particle spectrum consisting of equally spaced major shells with the same degeneracy, it can be shown in the case of closed shells (see p. 608 of Ref. [2] and Ref. [10]) that \( f_{sh}(N) \approx -\tau \exp(-\tau) \) for \( \tau \geq 1 \). Thus, because of the large prefactor \( 2\pi^2 \), the electronic-entropy effects manifest themselves even at the rather low temperatures of the cluster experiments. In Fig. 4 (bottom frame on the right), we compare the \( f_{sh}'s \) for \( K_3^+ \) and \( K_3^- \) calculated with the realistic single-particle spectrum of our FT-SCM. Note that, as a function of \( T \), the realistic spectrum suppresses the shell effect of the heavier-than-the-trimer magic fragments at a faster rate than the estimate above.

The controlling influence of the electronic entropy on the stabilization of electronic shells, discussed previously [6,10] in studies of ground-state properties of clusters, has been shown here to govern the fission patterns and their temperature dependence at the low-fissility regime [15]. This provides the impetus for further temperature-controlled studies of charge instabilities in clusters.

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Fig. 4. Theoretical FT-SCM temperature dependence of the \( Q^0 \)-values for the process \( K_f^{\pm} \rightarrow K_3^+ + K_3^{--} \). The dashed lines correspond to LDM calculations. In each case, the horizontal axis corresponds to various SCM channels \( 2 \leq P \leq 28 \). In the bottom right frame, we display \( f_{sh} \) vs. \( \tau \) (see text) for the \( K_3^+ \) and \( K_3^- \) fragments.

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[12] The FT-SCM [10] is an extension of the zero-temperature SCM [C. Yannouleas and U. Landman, Phys. Rev. B 51, 1902 (1995); *ibid.* 48, 8376 (1993)] and it incorporates triaxial deformations, single-particle electronic entropy, and thermal averaging over the shape fluctuations. In the SCM approach, the total free energy of a finite system of interacting delocalized electrons is divided into two contributions: a part that varies smoothly with the system size and thus coincides with the LDM, and an oscillatory term accounting for the quantal shell effects. In keeping with the experimental analysis [13], the LDM curvature coefficients were taken to be zero, while the surface coefficients for Li, Na, and K were taken as 1.30 eV, 0.85 eV, and 0.70 eV, respectively. The temperature dependence of these coefficients was neglected. For the rest of coefficients of the FT-SCM, see the first reference in this footnote for Na and K, and C. Yannouleas and U. Landman, J. Chem. Phys. 107, 1032 (1997) for Li.
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