Skimming-Trajectory Effect for Energy Losses of Medium Energy He Ions Passing along Major Crystal Axes of KI(001) and RbI(001)

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Abstract

The energy loss of an ion passing along a major crystal axis before and/or after undergoing a large-angle collision is strongly enhanced, because its path is close to a lattice-site atom located at the crystal axis, the so-called skimming effect. One may find this effect in high-resolution medium-energy ion scattering (MEIS) spectra from sub-mono layers of single crystals. In the present study, the MEIS spectra were observed for medium energy He$^+$ ions backscattered from KI(001) and RbI(001) crystals at various scattering geometries. The observed surface peaks were decomposed into scattering components from top, second, and third layer atoms considering the hitting probabilities, which were derived from Monte Carlo simulations of ion trajectories, taking account of the enhanced and correlated thermal vibrations. The results obtained here demonstrate that the energy loss of the ions which move skimming-trajectories is expressed reasonably well by the impact parameter dependent energy loss calculated using the non-perturbative coupled-channel method.

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I. INTRODUCTION

In the ion scattering spectrometry, the energy loss of ions allows for depth-profiling constituent elements of thin films. The energy loss or stopping power values have been compiled in a data-base semi-empirically by Ziegler and coworkers[1-3]. These data, however, can be applied only to polycrystalline and amorphous targets and also to random penetration in crystalline solids. On the other hand, the ions channeling along a crystal axis or plane undergo a smaller mean energy loss than that for the ions passing along a random direction, because the channeled ions penetrate through a region of a low electron density[4-6]. In contrast, the ions incident along a crystal axis and scattered to a large angle or scattered to a blocking direction undergo a large energy loss because they penetrate electron shells close to target nuclei. Recently, Hentz et al.[7] reported trajectory-dependent energy losses of medium energy protons incident on Cu(111) crystals. They performed Monte Carlo simulations of ion trajectories as well as coupled-channel calculations of impact parameter dependent energy loss and demonstrated that the energy spectrum observed for 100 keV H$^+$ ions is reproduced well by the simulations.

In the present study, we measured the energy spectra for 70 and 80 keV He$^+$ ions incident on KI(001) and RbI(001) for various scattering geometries. The low Debye temperatures of 130 K for KI and 115 K for RbI are quite convenient to see clearly the scattering component from the second layer atoms. The spectra were analyzed in a layer-by-layer fashion due to an excellent energy resolution ($\Delta E/E = 1.0 \times 10^{-3}$; full width at a half maximum) of a toroidal electrostatic analyzer (ESA)[8]. In order to decompose uniquely an observed surface peak into each scattering component from the top-, 2nd-, and 3rd-layer atoms, we performed Monte Carlo simulations of He ion trajectories considering enhanced and correlated thermal vibrations. The difference between the emerging energies ($\Delta E_{1,2}$) for the ions scattered from the top- and 2nd-layer atoms corresponds to the ion path between 1st and 2nd layer only. As both lines are observed for the same backscattering atom in the same backscattering direction, the kinematic (quasi-elastic) energy reduction as well as the enhanced inner-shell excitation is identical for both backscattering events. The electronic surface position (effective jellium edge) is always in front of the surface defined by the target nuclear positions and this distance is similar to the interaction range that determines electronic excitations during the collision with a backscattering atom in the 2nd layer. Thus, for the ion path corresponding to $\Delta E_{1,2}$, we need to consider only the incident and emerging angles of the ions and the nuclear interlayer distance. The resulting energy-loss difference corresponds to two processes, (i) local energy loss incurred during passing close to a lattice-site atom (skimming effect) and (ii) average
energy loss proportional to the path-length. The latter corresponds to the random stopping power. The results obtained experimentally are compared with theoretical predictions based on the coupled channel method[9,10] that does not rely on a perturbative treatment of the interaction dynamics.

II. EXPERIMENTAL

Medium energy He\(^+\) ions were provided by a duo-plasma ion source coupled with a direct current power supply with a voltage ripple better than \(5\times10^{-4}\). After acceleration, a switching magnet selected the ion species and energy. The ion beam was then collimated to a size of \(0.18\times2.0\text{ mm}^2\) in the horizontal and vertical directions, respectively before impinging on a target surface. The energies of scattered He\(^+\) ions were selectively analyzed by the toroidal electrostatic analyzer with a central deflection radius of 150 mm. The entrance slit size was \(0.2\times4.0\text{ mm}^2\) in the horizontal and vertical directions, respectively and the narrow slit opening in the horizontal plane is responsible for the energy selection. The deflected He\(^+\) ions arrived at a three-stage micro-channel plate combined with a position sensitive detector (20 mm in diameter) of a semiconductor (Si) type named PIAS (Hamamatsu Photonics) whose position resolution was 40 μm. The large inter-electrode distance of 16 mm covers a wide energy range (~8 % of a pass energy) for deflected He\(^+\) ions at a constant voltage applied on the toroidal electrodes. The condition mentioned above gives rise to the excellent energy resolution of \(1.0\times10^{-3}\) without degrading the statistics.

The KI(001) and RbI(001) crystals take a NaCl-type structure with lattice constant of 7.066 and 7.342 Å, respectively. A crystal substrate with a size of \(14\times14\times1\text{ mm}^3\) was obtained by cleaving the bulk crystal in the atmosphere and then immediately introduced into a scattering chamber evacuated to an ultrahigh vacuum (UHV: ~\(2\times10^{-10}\) Torr). Before the MEIS measurement, infrared radiation heated the substrate to ~150°C for 5 min to eliminate a contamination[11]. A clean surface without C and O adsorption was confirmed by Auger electron spectroscopy.

The sample surface was covered with Al foils except for an ion-irradiation area to be charge-up free and positively biased at 90 V to suppress secondary electrons emission. In spite of that, in many cases, the slope of the front edges of observed MEIS spectra was smeared out significantly in comparison with the expected system energy resolution, probably due to a slight charge-up. In order to avoid radiation damage, we shifted the irradiated area by ~0.2 mm in the horizontal direction after accumulating a beam current of \(1\ \mu\text{C}\).
III. MONTE CARLO SIMULATIONS OF ION TRAJECTORIES

In order to deduce accurately the emerging-energy difference between the scattering components from the top- and 2nd-layer atoms, the observed surface peak should be decomposed uniquely. For this purpose, it is essential to see the hitting probability for each layer atoms as well as the line shape for each scattering component. Concerning the latter, we employed the exponentially modified Gaussian (EMG) line shape[12,13] and the Lindhard-Scharff formula[14] to calculate the energy straggling, whose reliabilities were confirmed experimentally in advance[13,15-17]. The hitting probability for each layer atoms is calculated by Monte Carlo (MC) simulations of ion trajectories, which are explained in detail below.

Figures 1 illustrates the scattering geometries taken in the present MEIS spectrum observation. The surfaces are slightly relaxed (KI: −1.6 %, Rbl: −1.1 %) and rumpled (KI: 1.8 %, Rbl: 2.2 %)[11,18,19] but such reconstructions were neglected in the present MC simulations, which give the hitting probabilities (close encounter probabilities: $P_{CL}$) for the 2nd- and 3rd-layer atoms normalized by that for the top-layer atoms. In this Monte Carlo simulations, we assumed root-mean-square (rms) one-dimensional (1D) bulk thermal vibration amplitudes ($u_{bulk}$) of 0.20 and 0.205 Å, respectively for I and Rb of Rbl(001) and 0.190 and 0.195 Å, respectively for I and K of KI(001) determined experimentally[11,18,19]. In addition, also assumed are the correlations calculated from molecular dynamics using the Catlow potentials[20] and the thermal vibration amplitude of the top-layer atoms in the surface normal direction enhanced by $\sqrt{2}$ compared with the $u_{bulk}$. The normalized hitting probability for the $n$th-layer atoms located in a crystal string ($z$-axis) is calculated assuming a single-row approximation[21] by

$$P_{CL}(n) = \int ... \phi(\vec{r}_1, \vec{r}_2, ..., \vec{r}_{n-1}, \vec{r}_0 + \Delta_n) d^2\vec{r}_1 d^2\vec{r}_2 ... d^2\vec{r}_{n-1} d^2\vec{r}_0$$

$$= \int ... \phi(\xi_1, \xi_2, ..., \xi_{n-1}, \xi_0 + \Delta_n) d\xi_1 d\xi_2 d\xi_3 ... d\xi_{n-1} d\xi_0$$

$$= Q \int ... \frac{\phi(\xi_1, ..., \xi_{n-1}) \cdot q(\xi_1, ..., \xi_{n-1})}{Q} d\xi_1 d\xi_2 ... d\xi_{n-1}$$

$$\approx \frac{Q}{N} \sum_{j=1}^{N} \frac{\phi(\xi_1^{(j)}, ..., \xi_{n-1}^{(j)})}{q(\xi_1^{(j)}, ..., \xi_{n-1}^{(j)})}$$

$$\xi_i = \frac{x_i}{\sigma_{x(n)}}, \quad \xi_2 = \frac{y_1}{\sigma_{y(n)}}, \quad ..., \quad \xi_{n-1} = \frac{x_{n-1}}{\sigma_{x(n-1)}}, \quad \xi_{2n-2} = \frac{y_{n-1}}{\sigma_{y(n-1)}}, \quad \xi_{2n-1} = \frac{X_n}{\sigma_{x(n)}}, \quad \xi_{2n} = \frac{Y_n}{\sigma_{y(n)}},$$

where $\sigma_{x(n)}$ is an rms 1D thermal vibration amplitude of the $n$th-layer atoms in $x$-direction and ($\vec{r}_1, \vec{r}_2, ..., \vec{r}_{n-1}$) is the correlated $2(n-1)$-dimensional position vector. Here, the 2$n$-variate
normal probability distribution function is expressed by \(2n\)-dimensional correlation matrix \(S\).

\[
\varphi(\xi_1, \xi_2, \ldots, \xi_{2n}) = \frac{1}{\sqrt{(2\pi)^{2n}|S|}} \exp\left[-\frac{1}{2} \sum_{i=1}^{2n} \sum_{k=1}^{2n} (S^{-1})_{ik} \xi_i \xi_k \right]
\]

\[
q(\xi_1, \xi_2, \ldots, \xi_{2(n-1)}) = \frac{1}{A} \frac{1}{\sqrt{(2\pi)^{2(n-1)}|S|}} \exp\left[-\frac{1}{2} \sum_{i=1}^{2(n-1)} \sum_{k=1}^{2(n-1)} (S^{-1})_{ik} \xi_i \xi_k \right]
\]

where \(\bar{R}_0\) is a random starting position, \(A\) an incident area, \(\bar{R}_n : (X_n, Y_n) = \bar{R}_0 + \bar{A}_n\) the incident position in the \(n\)th-layer, \(N\) the number of incident ions, and \(S\) is the correlation matrix given by \(S_{ij} = \langle \xi_i \cdot \xi_j \rangle\) (\(\langle \rangle\) means a time average). The \(\bar{\xi} = (\bar{\xi}_1, \bar{\xi}_2, \ldots, \bar{\xi}_{2(n-1)})\) is correlated \(2(n-1)\)-variate normal distribution expressed by

\[
Q = \int \cdots \int q(\xi_1, \xi_2, \ldots, \xi_{2(n-1)}) d\xi_1 d\xi_2 \ldots d\xi_{2(n-1)} = 1
\]

FIG. 1. (Color on line) Scattering geometries taken in the MEIS measurements. Small and large circles denote cation and anion, respectively.

\(\bar{\xi} = T^{-1}\eta\), where \(\eta \equiv (\eta_1, \eta_2, \ldots, \eta_{2(n-1)})\) is the normalized position vector generated by assembling the normal distribution of the \(2(n-1)\)-univariate independent random variables and the transformation matrix \(T\) is given by \(\bar{T} T = S^{-1}\)\([22]\). A nuclear encounter would have taken place if the atom residing in the \(n\)-th layer were located exactly at \(\bar{r}_n = \bar{R}_n (= \bar{R}_0 + \bar{A}_n)\). Note
that only the coordinates in the (x, y) planes are relevant to the problem. We also assumed that
the motion in the x-direction is independent of that in the y-direction. The basic treatment to
calculate the hitting probabilities considering the correlations is given in the literature[21].
Thus the surface peak observed was deconvoluted uniquely into each scattering component
from the top-, 2nd-, and 3rd-layer atoms, considering the hitting probabilities (assumed to be
highly accurate) calculated from the MC simulations of He ions trajectories described above.

IV. RESULTS AND DISCUSSION

Figures 2 shows a typical MEIS spectrum (full circles) observed for 80 keV He\(^+\) ions
incident along the [00\overline{1}] axis of RbI(001) and scattered to 100° (random direction: 2° off
from [100]-azimuth; see Fig. 1: trajectory (8)). The surface peaks are decomposed into three
scattering components from the top-, 2nd-, and 3rd layer I and Rb atoms (solid curves). Note
that the shadowing effect was weakened owing to large root-mean-square thermal vibration
amplitudes expected from the low Debye temperature of 115 K. Here, we employed the EMG
function as the line shape[12,13]. The best-fit was obtained by assuming the energy
differences (\(\Delta E_{I-2}\)) of 520±20 and 540±30 eV between the scattering components from the
top- and 2nd-layer I and Rb atoms, respectively. In order to fit the leading edge, it was also
assumed that the energy straggling took a value of 0.35 \(\Omega_{B}^2\) (\(\Omega_{B}\): Bohr straggling)[14,15],
which is significantly larger than that given by the Lindhard-Scharff formula. This is probably
due to the slight charge-up, as mentioned before. The hitting probabilities (\(P_{CL}\)) deduced here
for the 2nd and 3rd-layer I and Rb atoms indicate significant correlations of +0.35 and +0.10,
respectively between the first and second nearest neighbor atoms in the [001]-string for the
motion perpendicular to this string. Here, the positive correlation represents an attractive
motion caused by acoustic phonon modes, which lowers the close encounter probability
because of a stronger shadowing effect. Note that acoustic phonon modes are generally
dominant rather than optical modes because the number of created phonons is proportional to
the density of state multiplied by the Planck distribution function. Figure 3 shows the MEIS
spectrum observed for 80 keV He\(^+\) ions incident along the [10\overline{1}] axis of RbI(001) and
backscattered to [101] direction (see Fig. 1: trajectory (1)). A best-fitted MEIS spectrum was
obtained assuming the energy difference of \(\Delta E_{1-2} = 390\pm10\) eV and a significant correlation
of +0.15 between the first nearest neighbor I atoms in the [101]-string. The smaller correlation
for the [101]-string compared with that for the [001]-string is attributed to the larger
inter-atomic distance by \(\sqrt{2}\). The correlations estimated here are consistent with those
calculated from molecular dynamics simulations reported previously[11,18,19]. This
evidences the validity of the present deconvolution procedure.

FIG. 2. MEIS spectrum (full circles) observed for 80 keV He$^+$ ions incident along the [00$ar{1}$]-axis of Rbl(001) and scattered to 100° about 2° off from [100]-azimuth (random emergence). Thick (brown) and thin (blue) solid curves denote best-fitted spectrum and scattering components from top-, 2nd- and 3rd layer atoms, respectively. Best-fitted spectrum was obtained by assuming energy difference $\Delta E_{1-2} = 520$ eV for I and 540 eV for Rb and hitting probabilities of 0.58 and 0.30 for the 2nd and 3rd-layer I atoms and 0.51 and 0.27 for Rb atoms in [001]-crystal axis, which correspond to correlations of +0.35 and +0.10, respectively between 1st and 2nd nearest neighbor atoms.

From the energy differences ($\Delta E_{1-2}$) derived here for 80 keV He$^+$ ions scattered from Rbl for various kinds of scattering geometries, as indicated in Table I, we can deduce the local atomic energy loss (skimming effect) and random energy loss ($S^R_{\text{tot}}$: eV/Å) proportional to the path-length. For simplicity, as the first approximation, it is assumed here that the local energy loss is subjected to the He ions during passage through a major crystalline axis, while the ions penetrate along random directions undergo a random energy loss. Later we constrain the region of the local energy loss within a Wigner-Seitz cell to match the coupled channel calculations. The energy difference ($\Delta E_{1-2}$) observed for the [10$ar{1}$]-incidence and backscattered by I atoms to the [101]-direction, the local energy loss subjected by skimming through the top-layer I atoms $\Delta E(I)$ is determined to be 195±10 eV (just the half of the
energy difference $\Delta E_{i-2} = 390$ eV). This local energy loss value would correspond to 195 eV/Å = 37.6 eV/Å, which is much larger than the Ziegler’s random stopping power of 12 eV/Å for 80 keV He$^+$ ions passing through RbI. The local energy loss subjected by passing close to a Rb atom $\Delta E(\text{Rb})$ is then derived to be 190±15 eV and the random energy loss ($S_{\text{random}}^{\text{Rb}}$) of 15.8 eV/Å is obtained from the energy difference ($\Delta E_{i-2}$) observed for the [10$\bar{1}$]-incidence and backscattered from Rb and I atoms to the [102]-direction (see Table I and also Fig. 1: trajectory (2)). The random energy loss values are also deduced to be 15.6 and 16.3 eV/Å, respectively from the energy differences ($\Delta E_{i-2}$) observed for the [00$\bar{1}$]-incidence and scattered to 100° using the above local energy loss values (see Fig. 1: trajectory (8)). Combined with the energy differences derived for the two scattering geometries; [00$\bar{1}$]-incidence and scattered to 100° (trajectory (8)) and [11$\bar{1}$]-incidence and scattered to [112] direction (trajectory (6)), we obtain the random energy loss of 16.5 eV/Å. The local energy loss for Rb is also derived to be $\Delta E(\text{Rb}) = 206\pm4$ eV from the $\Delta E_{i-2}$ observed for the [10$\bar{2}$]-incidence and scattered to [201] direction (see Fig. 1: trajectory (4)). However, in this scattering geometry the path length along the [201]-axis (skimming path) is too long (8.21 Å). This probably gives an apparently larger local energy loss value. Therefore, it is reasonable to constrain the region of the atomic (local) energy loss within a critical volume, for example, a Wigner-Seitz cell. Indeed, this treatment matches the coupled channel approach in which the excitation and ionization probabilities are calculated along the incident ion path. Here, we assume that the volume of the Wigner-Seitz cell is equal to the volume of a sphere, $4\pi R^3/3$ and thus we obtain the radius of $R = 2.277$ for RbI and 2.192 Å for KI.

The path-length for random energy loss is corrected by subtracting $2R\times$(number of skimming atoms) from the total path-length, as indicated in Table I. In the case of the [10$\bar{1}$]-incidence and backscattered from I to the [101]-direction (see Fig. 1: trajectory (1)), we neglected the random energy loss. This should lower slightly the local energy loss determined previously for He ions skimming through an I atom, while slightly increase the random energy loss. Note that the inter-atomic distance in the [101]-string is slightly longer than the diameter of the Wigner-Seitz sphere and thus correction is small enough. At present we adopt the averaged random energy loss value of 16 eV/Å in spite of some uncertainty roughly estimated to be ±1 eV/Å. The corrected local energy loss values for I and Rb are shown in Table I. Thus we obtain the average local energy losses of $\Delta E(I) = 187 \pm 20$ and $\Delta E(\text{Rb}) = 174 \pm 20$ eV for 80 keV He$^+$ ions skimming through an I and Rb atom of RbI(001), respectively.

Similar analysis was also carried out for 70 keV He$^+$ ions incident on KI(001). Figures 4 and 5 show the MEIS spectra observed for 70 keV He$^+$ ions incident along the [10$\bar{1}$]-axis and
FIG. 3. MEIS spectrum (full circles) observed for 80 keV He\(^+\) ions incident along [1\(\bar{1}\)0]-axis of RbI(001) and scattered from I to [101]-direction. Notation is quite the same as that in FIG. 2. Best-fitted spectrum was obtained by assuming energy difference \(\Delta E_{1-2} = 390\) eV for scattering component from I and hitting probabilities of 0.35 and 0.06, respectively for the 2nd and 3rd-layer I atoms in [101]-string, which correspond to correlation of +0.14 between 1st nearest neighbor I atoms.

scattered from I atoms to the [101]-direction and incident along the [11\(\bar{1}\)]-axis and scattered from K atoms to random direction (see Fig. 1: trajectory (7')), respectively. For the [10\(\bar{1}\)]-incidence, the observed MEIS spectrum was best-fitted assuming the energy difference (\(\Delta E_{1-2}\)) of 350\(\pm\)10 eV and the hitting probabilities of 0.35 and 0.03 for the 2nd- and 3rd-layer I atoms, indicating no correlation, while for the [11\(\bar{1}\)]-incidence the \(\Delta E_{1-2}\) value of 305\(\pm\)15 eV and the hitting probabilities of 0.519 and 0.238 for the 2nd- and 3rd-layer K atoms, corresponding to correlation of +0.10 between the neighboring K and I atoms in the [111]-string. We also found a significant correlation of +0.20 between the neighboring I atoms in the [10\(\bar{1}\)]-string ([10\(\bar{1}\)]-incidence and random emergence; not shown here). No correlation found for the [10\(\bar{1}\)]-[101] double alignment geometry is possibly due to a smaller correlation in the [101]-string rather than that in the [001]-string or slight misalignment. The correlations derived here are consistent with those calculated from molecular dynamics analysis, indicating the validity of the convolution procedure[11,18,19] also for the KI(001) target.

From the energy difference \(\Delta E_{1-2} = 350\pm10\) eV observed for the [10\(\bar{1}\)]-incidence and [101]-emergence, the local energy loss for 70 keV He\(^+\) ions skimming through an I atom is determined to be 175\pm10 eV significantly smaller than that (195\pm10 eV) for 80 keV He\(^+\) ions.
FIG. 4. MEIS spectrum (full circles) observed for 70 keV He$^+$ ions incident along [10$ar{1}$]-axis of KI(001) and scattered from I atoms to [101]-direction. Thick (brown) and thin (blue) solid curves denote best-fitted spectrum and scattering components from top-, 2nd- and 3rd layer atoms, respectively. The best-fitted spectrum was obtained by assuming energy difference $\Delta E_{1-2} = 350$ eV for scattering component from I and hitting probabilities of 0.35 and 0.03, respectively for the 2nd and 3rd-layer I atoms in the [101]-string, which correspond to no correlations.

FIG. 5. MEIS spectrum observed with 70 keV He$^+$ ions incident along [11$ar{1}$]-axis of KI(001) and scattered from K to 65.3° with respect to surface normal (random direction). Notation is quite the same as that in FIG. 4. Best-fitted spectrum was obtained by assuming energy difference $\Delta E_{1-2} = 305$ eV and hitting probabilities of 0.52 and 0.24, respectively for the 2nd and 3rd-layer K atoms in the [111]-string, which correspond to correlation of +0.10 between the neighboring K and I atoms.
This local energy loss of 175 eV corresponds to 35 eV/Å, which is much larger than the Ziegler’s random energy loss of 10.5 eV/Å. Then we obtained the random energy loss value, $S_{e}^{KI}$ of 13.8 eV/Å from the energy difference ($\Delta E_{I-2}$) observed for the [10\overline{1}] incidence and random emergence (see Table I). As mentioned before for RbI(001), we must correct the path length for random energy loss. Here, we adopt the random energy loss value of 14 eV/Å, slightly larger than the above value. The corrected path lengths and the local energy loss values estimated assuming the random energy loss of 14 eV/Å are indicated in Table I. The local energy loss value for an I atom averaged for three different geometries is 165.5 eV with a slight dispersion of ±1 eV. The energy difference ($\Delta E_{I-2}$) observed for the [11\overline{1}] -incidence and scattered to 65.26° with respect to surface normal (random) gives the local energy loss of $\Delta E(K) = 120$ eV (see Fig. 1: trajectory (7)). The local energy loss of 257.2 eV for He ions skimming through a K and I atom estimated from the data observed for the [10\overline{1}] -incidence and [201] -emergence (see Fig. 1: trajectory (3)) is smaller about 10% than that expected from the other scattering geometries (120 + 165.5 = 285.5 eV), indicating roughly the uncertainty of ±(10 - 20) eV.

We calculated the impact-parameter ($b$) dependent energy loss by the coupled channel method[9]. Shortly, the time-dependent Schrödinger equation was solved for one active target electron in the framework of the independent particle model. The projectile-electron
interaction was described by a single-zeta potential for He$^+$ projectiles. Excitation and ionization probabilities were numerically calculated shell-by-shell allowing the contributions of transitions for about 500 discrete and continuum (wave packet) states with orbital quantum numbers up to $l = 8$ and energies up to $2m_e v^2$ ($m_e$: electron mass and $v$: ion velocity). The average energy transfers for 70 and 80 keV He ions to each electron from different sub-shells of I, K, and Rb atoms are indicated in Table II. Here, we assumed only near central collisions ($b$ close to 0) and a frozen ionic state of He$^+$.

The local energy losses determined from the MEIS observation are compared with the theoretical values calculated from the coupled channel method in Table III. The local energy losses for 80 keV He$^+$ ions skimming through an Rb and I atom determined from the present MEIS spectrum analysis agree well with the theoretical predictions. Surprisingly, the local energy loss for 70 keV He$^+$ ions passing close to a $^{19}$K atom calculated from the coupled channel method is significantly larger than that for the He$^+$ ions penetrating close to an $^{53}$I atom despite the smaller number of total electrons. Note, however, that the number of electrons easily removed according to the criterion, $2m_e v^2 > I_b$ ($2m_e v^2$ is the maximum energy transfer in a free projectile-electron collision and $I_b$ is the shell binding energy) is larger for $^{19}$K. Thus, the observed local energy loss values are basically consistent with the coupled channel calculations, although the $\Delta E(K)$ value derived experimentally is smaller than the calculated one by almost 30%. This deviation may come from the assumed independent particle model that does not consider dynamical screening and increased binding energy in multiple ionizations, which may be more important for the case of slow He projectiles compared to protons. Other uncertainties of the present calculations such as the use of a target-centered basis set (not suitable to describe capture events) and frozen projectile charge-state lower than the corresponding equilibrium value may also be responsible for the observed disagreement.

As mentioned above, the energy losses due to the skimming effect $<\Delta E>/2R$ (35 eV/Å for RbI and 30 eV/Å for KI) are much larger than the Ziegler’s stopping power values of 12 eV/Å for RbI and 10.5 eV/Å for KI[2]. Concerning the random energy loss $S_e$, the roughly estimated values of $S_e^{RbI} = 16$ and $S_e^{KI} = 14$ eV/Å, respectively for 80 and 70 keV He$^+$ passing through RbI and KI, respectively are also considerably larger than the Ziegler’s stopping powers. There is, of course, an energy uncertainty coming from incoming and outgoing energy difference. However, the outgoing energy is above 90% of the incoming energy for the present scattering geometries. Therefore, this energy uncertainty contributing to random energy loss is 5% at most. In the case of the local energy loss, the contribution to the
energy loss uncertainty is less than 10%, comparable with the errors estimated from the uncertainties mainly originating from the deconvolution procedure. Note that the Ziegler’s random energy loss formula acts as a useful guide but does not always give reliable data in the low and medium energy regimes.

Table II. Shell-by-shell energy-loss values calculated from the coupled-channel method for 70 keV He\(^+\) ions impact on KI(001) and 80 keV He\(^+\) ions incident on Rbl(001). The impact parameter is assumed to be 0 (head-on-collision) and electronic shells with binding energies exceeding 600 eV have been excluded. The numbers in parentheses indicate the electron occupancies.

| Shell | Energy Loss (eV/electron) | Shell | Energy Loss (eV/electron) |
|-------|---------------------------|-------|---------------------------|
| 2s (2) | 4.0                       | 4s (2) | 3.0                       |
| 2p0 (2) | 1.6                      | 4p0 (2) | 4.2                      |
| 2p1g (2) | 0.0                      | 4p1g (2) | 1.7                      |
| 2p1u (2) | 0.0                      | 4p1u (2) | 0.9                      |
| 3s (2) | 21.9                      | 4d0 (2) | 15.6                     |
| 3p0 (2) | 30.1                      | 4d1g (2) | 4.7                      |
| 3p1g (2) | 11.4                      | 4d1u (2) | 3.9                      |
| 3p1u (2) | 11.1                      | 4d2g (2) | 2.6                      |
| 4s (1) | 9.1                       | 4d2u (2) | 2.8                      |
|       |                           | 5s (2) | 19.3                     |
|       |                           | 5p0 (2) | 18.9                     |
|       |                           | 5p1g (2) | 7.3                      |
|       |                           | 5p1u (1) | 7.4                      |
| total (eV) | 169.6                   | total (eV) | 158.8                   |

| Shell | Energy Loss (eV/electron) | Shell | Energy Loss (eV/electron) |
|-------|---------------------------|-------|---------------------------|
| 3s (2) | 0.7                       | 4s (2) | 3.7                       |
| 3p0 (2) | 6.5                       | 4p0 (2) | 4.1                       |
| 3p1g (2) | 1.5                       | 4p1g (2) | 1.9                       |
| 3p1u (2) | 1.5                       | 4p1u (2) | 1.1                       |
| 3d0 (2) | 10.5                      | 4d0 (2) | 17.8                      |
| 3d1g (2) | 1.2                       | 4d1g (2) | 5.5                       |
| 3d1u (2) | 0.6                       | 4d1u (2) | 5.0                       |
| 3d2g (2) | 0.5                       | 4d2g (2) | 3.0                       |
| 3d2u (2) | 0.4                       | 4d2u (2) | 3.5                       |
| 4s (2) | 20.5                      | 5s (2) | 18.9                      |
| 4p0 (2) | 26.4                      | 5p0 (2) | 19.7                      |
| 4p1g (2) | 9.3                       | 5p1g (2) | 7.5                       |
| 4p1u (2) | 9.2                       | 5p1u (1) | 7.6                       |
| 5s (1) | 8.3                       |       |                           |
| total (eV) | 185.8                   | total (eV) | 187.0                   |
V. CONCLUSION

The energy loss of ions passing close to a lattice site atom before and after a large angle collision is strongly enhanced, due to the so-called skimming effect. This phenomenon is quite different from the energy losses of ions penetrating in a random or channeling direction, because the ion impinging and/or emerging along a major crystal axis before or after a large angle collision takes a path close to a lattice site atom located in the crystal axis and thus tends to excite the inner shell electrons. The excellent energy resolution of the toroidal ESA makes it possible to resolve energetically each scattering component from the top- and 2nd-layer atoms. The resulting energy difference between the two scattering components originates from two parts, (i) local energy loss subjected by skimming through a lattice site atom and from (ii) usual random energy loss proportional to a path-length. We measured the MEIS spectra using 70 and 80 keV He\(^+\) ions for KI(001) and RbI(001) substrates under various kinds of scattering geometries and decomposed the surface peaks into basically three scattering components from the top-, 2nd-, and 3rd-layer atoms considering the hitting probabilities, which were calculated from the MC simulations of He ion trajectories taking account of enhanced and correlated thermal vibrations. The local energy losses were determined to be \(\Delta E(\text{I}) = 187 \pm 20\) eV for 80 keV He\(^+\) incident on RbI(001) and \(\Delta E(\text{Rb}) = 174 \pm 20\) eV for 80 keV He\(^+\) incident on KI(001). The above local energy losses \(<\Delta E>/2R\) are much larger than the Ziegler’s random stopping powers more than 3 times. Simultaneously, the random energy losses for 80 and 70 keV He\(^+\) passing through RbI and KI, respectively are estimated to be 16±1 and 14±1 eV/Å, which are also considerably larger than the Ziegler’s energy loss values of 12.0 and 10.5 eV/Å. This may be due to either additional excitation channels at the very surfaces or inaccuracies of the Ziegler’s stopping procedure at low and medium energies. The local energy losses determined experimentally for skimming through an Rb and I atom of RbI(001) and for passing close to an I atom of KI(001) are in good agreement with the calculated ones by the coupled channel method, while for a K atom of KI(001) the observed local energy loss deviate significantly from the couple channel calculations almost 30 %. This deviation may be attributed to use of the independent particle model, where the effect of dynamical screening is not taken into account, which may be important for slow heavy projectiles. Despite that, the local energy losses obtained by the MEIS spectrum analysis are basically consistent with the impact-parameter dependent energy losses given by the coupled channel calculations.

Such a strongly enhanced energy loss works effectively to resolve each scattering component from the top-, 2nd-, and 3rd-layer atoms in MEIS spectra. However, we must note
that surface peaks should be deconvoluted carefully considering the hitting probabilities for each layer atoms, which can be calculated from MC simulations of ion trajectories. It is emphasized that such a strongly enhanced electronic energy loss is a key issue to analyze exactly the MEIS spectra from subsurface ordered structures.

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