Measurement of MeV Ion Track Structure in an Organic Solid

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Radial velocities and effective ejection radii of hydrocarbon secondary ions sputtered electronically from polymer films have been experimentally determined. Using these data, the radial profiles of the energy and momentum density in an individual MeV ion track on a picosecond time scale are directly mapped out. [S0031-9007(96)00742-9]

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Ion tracks, formed by the interaction of energetic charged particles with matter, have played an important role in the development of modern physics. More recently, MeV-GeV ion track production has been used to modify materials in a controlled manner [1,2], being a potential tool to engineer material properties in the nanometer to micrometer scale. MeV-GeV ions are finding recently wider use in radiation therapy for, e.g., treatment of different forms of cancer. In spite of this, the understanding of the evolution of energy deposited around the path of an ion has not progressed markedly since early works in the sixties and seventies [3–5].

Ions with velocities higher than the Bohr velocity \(v_B = 0.22 \text{ cm/ns}\) deposit their energy in electronic excitation and ionizations in a cylindrically symmetric region around the path of the impacting ion, forming the track core. The incident ions cause ejection of energetic secondary electrons that transport part of this energy out of the core [3,4]. The mean deposited energy density, \(e(r)\), after the ion passage is approximated by a \(e(r) \propto r^{-2}\) dependence, where \(r\) is the radial distance from the ion path. Such a dependence is obtained from theory employing the continuous slowing down approximation [4,6] for the secondary electrons, Monte Carlo simulations [5–7], as well as experiments performed in gases [8,9]. Several energy-transport mechanisms [10,11] have been advanced to describe the subsequent energy propagation and dissipation. However, no direct experimental determination of the initial or evolved energy distribution exists for targets in the condensed phase. In this work a first estimate of \(e(r)\), the radial profile of the mean deposited energy density, is made for a solid material.

When fast ions interact with dielectric materials, a trail of radiation damage in the bulk (damage tracks) [3] as well as material erosion of the surface (electronic sputtering) are observed [12,13]. Whereas the characteristics of latent damage tracks have been determined by various physical methods \textit{ex situ}, long after the impacting event [14], the study of the sputtered material can in principle provide “real time,” \textit{in situ} information on the complex physical and chemical processes occurring in ion tracks [15–17]. Here we demonstrate the use of high resolution time-of-flight mass spectrometry of the ionic ejecta [13,17] to reconstruct the track of an MeV ion in an organic polymer on the picosecond time scale.

Radial velocity distributions of hydrocarbon secondary ions, ejected from poly(vinylidene fluoride), PVDF \([\text{[CH}_2\text{CF}_2]_n\]\), and other polymer targets have been correlated to their chemical compositions [18]. This is exemplified in Fig. 1, where the mean radial velocities, \(\langle v_r \rangle\), and the FWHM \((\sigma(v_r^2))\) of the velocity distributions of hydrocarbon ions ejected from a pristine PVDF film are plotted as a function of the ion mass. These moments of the radial velocity distributions are seen to vary systematically with the degree of hydrogenation of the secondary ions. Such an effect demonstrates that there is a correlation between the secondary ion chemical composition and the ion formation and ejection processes. The composition of the ejected positive ions is directly related to their chemical and thermal history in the track. For example, ionic species, e.g., \(\text{C}_n^+\), formed by a strong depletion of H atoms from the macromolecular chains have the highest \(\langle v_r^2 \rangle\) values (Fig. 1), suggesting these ions originate from the “hotter” (and inner) regions of the track. In contrast, saturated (e.g., \(\text{C}_n\text{H}_{2n}^+\)) species show the lowest \(\langle v_r^2 \rangle\) values. Thus the dependence of \(\langle v_r^2 \rangle\) on ion mass in Fig. 1 actually reflects the radial profile of the deposited energy. These data can, in principle, be used to determine the radial profile of the energy density, as recently argued [18], but determining this would require a spatial model for the ejection process.

Alternatively, having an independent determination of the location of ejection of the ions, the results from Fig. 1 can be converted into radial profiles of the energy density in the track of a single ion. A direct measurement on the nanometer scale of the region in the track from which a particular ion originates is, of course, difficult to realize experimentally. Here we show that an experimental estimate of the mean ejection radius for the secondary ions can be inferred from the “disappearance cross section” [14,19,20]. In the context of sputtering, the disappearance cross section is derived from the decrease in the yield of
This decrease typically follows a Poisson distribution, leading to Eq. (1). Because of the relationship between the production of a fragment and damage, the quantity $\sigma_i$ can be associated with an effective ejection area. That is, within the area $\sigma_i$ the energy density is too high for an ion of a particular size to be formed and ejected intact. At some larger distance, the energy density will be too low for such ions to be formed and ejected. Thus ejection can be considered to occur from a ring with inner radius, roughly $(\sigma_i/\pi)^{1/2}$, and some outer radius $(\sigma_i/\pi)^{1/2} + \Delta r$. Assuming that $\Delta r \ll (\sigma_i/\pi)^{1/2}$, the effective ejection radius $\eta$ is approximately $(\sigma_i/\pi)^{1/2}$. This interpretation of the measured damage cross section is seen below to be consistent with the data on $\langle v_x^2 \rangle$ and $\langle v_y \rangle$.

The disappearance cross sections for positive secondary ions ejected from PVDF targets, using charge-equilibrated 55 MeV $^{127}$I as primary ions, were obtained in the following experiment [21]. A $1 \times 1 \text{mm}^2$sample area was bombarded at $45^\circ$ angle of incidence with a beam current less than 5 nA/cm$^2$ to the desired fluence. Secondary ion mass analysis of the irradiated target was done in situ after each irradiation by reducing the beam intensity to $= 2000 \text{ ions/s}$. The ejected secondary ions were analyzed in a reflectron time-of-flight mass spectrometer [22]. The ion flight time registration was performed in an event-by-event mode by employing a time-to-digital converter. Typically $4 \times 10^8$ primary ion impacts were accumulated for each mass spectrum. The process was repeated by increasing the fluence (from $10^9$ to $1.3 \times 10^{13} \text{ ions/cm}^2$), bombarding one and the same target spot. The vacuum in the experimental chamber was always better than $5 \times 10^{-9} \text{ torr}$.

The value of $\sigma_i$ for a specific ion is obtained by fitting Eq. (1) to the data points for ion intensity, $Y_i(\phi)$, as a function of fluence $\phi$. The $\sigma_i$ values obtained for different hydrocarbon ions, $C_nH_m^+$, in the range up to $M/Z = 100$, where $M$ and $Z$ are, respectively, the ion mass and charge, are shown in Fig. 2. The $\sigma_i$ values are seen to exhibit a regular behavior with the ion mass, meaning that $\sigma_i$ is correlated with the chemical composition of the ions, as were $\langle v_x^2 \rangle$ and FWHM. The

![FIG. 1. (a) Mean radial velocities $\langle v_x \rangle$ for the $C_nH_p^+$ ion series ejected from PVDF by charge-equilibrated 72.3 MeV $^{127}$I ions. (b) FWHM ($= 0.5M[(v_z^2) - (v_x^2)]$) of the $v_x$ distributions for the same ions (adapted from Ref. [18]).](image)

![FIG. 2. Disappearance cross sections $\sigma_i$ for hydrocarbon secondary ions ejected from PVDF targets by 55 MeV $^{127}$I, as a function of mass $M$ (see text for further details).](image)
cross sections, or equivalently the effective ejection radii, increase with increasing number of hydrogen atoms in the ion species for the same C-atom series. This is sensible since the heat of formation of these ions depends on their chemical composition and the degree of fragmentation within the track region depends on the transient local energy density in the track. Therefore, the observed dependence of $\sigma_i$ on ion mass indicates the origin of ion ejection. Ions with a lower number of hydrogen atoms originate from the hotter areas of the ion track, thus having both smaller effective ejection radii and higher $\langle u_r^2 \rangle$. In comparison, ions with higher hydrogen content are formed further away from the track center. Therefore, the three sets of data (the $\langle u_y \rangle$, $\langle u_z^2 \rangle$ and $\sigma$) are self-consistent if we use the interpretation above for the damage cross section as an effective ejection radius. A slight increase in $\sigma$ with increasing mass (i.e., size) of the species is also discerned for the pure carbon cluster ions, but the hydrogen content is more sensitive to the local energy density.

The cross sections derived from secondary ion yields also determine the eventual macroscopic transformations of the properties of the targets, such as mechanical and electrical properties, coupled to the carbonization of the polymer under long-term irradiation [21]. Thus the cross sections were estimated excluding the high fluence points. Not surprisingly, the cross section values of $C_nH_mF$ secondary ions, also included in the plots, follow the same trend of the pure hydrocarbon ion series. We note that the velocity distributions were obtained using 72.3 MeV $^{127}$I ions, with a slightly higher stopping power ($\approx 10\%$), and velocity ($\approx 15\%$), compared to 55 MeV $^{127}$I ions. This results in a higher mean deposited energy density in the track for 55 MeV $^{127}$I ions. The effect on the velocity distributions should be small, as $\langle u_x^2 \rangle$ and $\langle u_y \rangle$ increase slowly with $dE/dx$, the stopping power [24]. However, previous data on the damage cross section obtained for polymer films indicate an increase in the cross sections of up to a factor of 2 for 55 MeV I bombardments compared to 78 MeV I ions [25]. To account for these differences the radii in Fig. 3 were divided by a factor of $\sqrt{2}$.

The experimental points in Fig. 3(a) are well described by a power law decrease. The best fit to the data is $\langle u_x^2 \rangle \propto r^{-1.2}$, but powers between $r^{-1}$ and $r^{-2}$ also fit considering the scattering of the data and the uncertainties introduced by the corrections to $r$.

In Fig. 3(b), the mean velocity (related to the impulse transferred to the ions) is displayed as a function of the effective radius. The following trends can be distinguished. At small $r$ (up to $\approx 25$ Å), ions have a positive $\langle u_y \rangle$ (corresponding to ejection backwards along the track; see inset of Fig. 1), as well as higher $\langle u_z^2 \rangle$. At large $r$ ($>35$ Å), ions tend to have similar mean velocities, but away from the line of incidence of the MeV ion. A relatively steep transition region between 25 and 35 Å is also seen which is roughly consistent with the simple pressure pulse model for electronic sputtering [26]. In Fig. 3(b) the dashed lines mark approximately these three regions. It is seen from Figs. 3(a) and 3(b) that the near-surface energy density in the track produces fragmentation and ejection processes that are closely intertwined. The local energy density triggers the fast “hot” chemistry responsible for bond rearrangement in the track [25]. However, the dependence of the mean velocity of the ions with the place of emission indicates ejection occurs in response to the volume force arising from the energy density gradient in the track [26]. It provides the momentum transfer critical for removing newly formed species from the surface sufficiently rapid for survival. We note that the direction of the momentum transferred to the target species, and thus the variation of $\langle u_y \rangle$ with $r$, is coupled to the geometry of the MeV ion impact. For example, in an impact at normal incidence it
is expected that $\langle v_y \rangle = 0$ for the same ions [27], distinct from the dependence found here for 45° impacts.

In conclusion, we have derived directly from experiments the radial variation of quantities related to the mean energy density ($\langle x v_y^2 \rangle$) and the surface momentum transfer ($\langle x v_x \rangle$) for the track of a polymer film using secondary ion ejection. This was done in a self-consistent way using the damage cross section for the radial scale. This quantitative result can be used as a test for modeling track formation. The following qualitative picture of the track structure is obtained: Species emerging closer to the core formation. The following qualitative picture of the track can be used as a test for modeling track ing the damage cross section for the radial scale. This work was supported by the Swedish Natural Science Research Council (NFR), the Wallenberg Foundation, and the Ångström and Cluster Consortia. M. Inokuti and C. T. Reimann are acknowledged for helpful comments.

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[23] The total energy is related to the kinetic energy ($\langle x v_y^2 \rangle$) plus the chemical energy of rearrangement and fragmentation reactions. The Coulomb potential energy "stored" in the electric field of the localized ionic species in the track core may also contribute to the $\langle x v_y^2 \rangle$ value by, e.g., accelerating the positive ions during ejection.


