Ultrafast Dynamics of a Nucleobase Analogue Illuminated by a Short Intense X-ray Free Electron Laser Pulse

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Understanding x-ray radiation damage is a crucial issue for both medical applications of x rays and x-ray free-electron-laser (XFEL) science aimed at molecular imaging. Decrypting the charge and fragmentation dynamics of nucleobases, the smallest units of a macro-biomolecule, contributes to a bottom-up understanding of the damage via cascades of phenomena following x-ray exposure. We investigate experimentally and by numerical simulations the ultrafast radiation damage induced on a nucleobase analogue (5-iodouracil) by an ultrashort (10 fs) high-intensity radiation pulse generated by XFEL at SPring-8 Angstrom Compact free electron Laser (SACLA). The present study elucidates a plausible underlying radiosensitizing mechanism of 5-iodouracil. This mechanism is independent of the exact composition of 5-iodouracil and thus relevant to other such radiosensitizers. Furthermore, we found that despite a rapid increase of the net molecular charge in the presence of iodine, and of the ultrafast release of hydrogen, the other atoms are almost frozen within the 10-fs duration of the exposure. This validates single-shot molecular imaging as a consistent approach, provided the radiation pulse used is brief enough.

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I. INTRODUCTION

During the last decade, new research avenues have been opened for both spectroscopic and structural investigations of matter, through the advent of the x-ray free-electron-laser (XFEL) [1–3]. XFELs deliver coherent x-ray pulses, combining unprecedented intensities and extremely short pulse durations, as short as a few femtoseconds only. For selected applications, these qualities offer important advantages over conventional x-ray sources. Time-resolved spectroscopy on the time scale of femtoseconds to picoseconds allows us to monitor, for the first time, atoms and electrons in action [4–14]. For structural analysis, the extreme intensity focused into a couple of μm2 transforms single-shot diffraction imaging of biomolecules and nanosize objects from a remote goal into a tangible reality [15–19].

Connecting these two realms of XFEL-based research is the fundamental issue of radiation damage. X-ray...
absorption often leads to the breakage of chemical bonds, geometrical changes, and distortions of electron distributions that manifest the deterioration of the original object. In XFEL-based experiments for a single-molecule or a nanosize object, the amount of energy deposited in the target by a single pulse is sufficiently high to cause its total destruction; from a structural imaging point of view, the question is whether the time scale of these changes is slow enough for the collected data to capture essential high-quality structural information about the original object [20]. Our present study addresses this question.

The phenomenon of radiation damage is also interesting by itself because it is a consequence of the complex interplay between fast electron dynamics and slower nuclear dynamics, following the rapid creation of highly charged, highly energetic states. X-ray radiation creates charges by core ionization, which is followed by internal electronic relaxations, i.e., Auger processes that may occur in cascades. Core ionization is thus predominantly the first step of radiation damage in biomolecules from living tissue subjected to medical or unintentional x-ray exposure [21]. The cascading processes of creating low-energy electrons and ionic radicals, which play a crucial role in the damage, are also of current fundamental interest [22,23]. XFEL pulses, which are the subject of the present work, are known to cause, even for isolated atoms, major changes in electronic structure via multiphoton absorption followed by Auger processes [24–27]. At the other end of the scale, in condensed matter and clusters of $10^3–10^5$ atoms, the intense XFEL pulses produce a dense nanoplasma, in which the dynamics of electrons and ions are strongly correlated with each other [28,29].

The coupled electron and ion dynamics strongly depend on the size of the system, laser fluence, and photon energy. Studying the response of a single molecule composed of a small number of atoms to the XFEL pulse is perhaps the most rewarding, as various levels of theory and experimental methods inapplicable to larger systems may be applied. This was illustrated by Erk et al., who reported ionization and fragmentation of methyleneselenol (CH$_3$SeH) molecules by intense ($>10^{17}$ W/cm$^2$) 5-fs long x-ray pulses of about 2 keV energy, using coincident ion momentum spectroscopy [30]. They reported a molecular expansion of about 50% within the first $5–10$ fs. Motomura et al. reported a similar study on iodomethane (CH$_3$I) molecules using XFEL pulses of 5.5 keV at SACLAXFEL in Japan [31]. The latter study concluded that the C-I bond length increased by only 10% during the first 10 fs.

The target of the present study is a more complex molecule, a 5-halouracil, which is an analogue of uracil, one of the four canonical nucleobases of ribonucleic acid (RNA). While 5-flourouracil (FU) was identified several decades ago as an effective anticancer drug [32], iodine substitution makes the uracil molecule much more vulnerable to x-ray damage owing to iodine’s large absorption cross section for hard x rays. Indeed, a possible use of 5-iodouracil (IU) as a radiosensitizer in tumor therapy has been discussed [33] and the radiation damage involving 5-IU has been studied extensively by simulations [34,35]. To our knowledge, however, there has been no experimental report on the destruction of 5-IU exposed to x rays.

The aim of this work is to trace back the intricate details of the early stages of the Coulomb explosion of 5-IU, as a biomolecule of practical relevance, utilizing XFEL pulses as the ionizing radiation and advanced multiparticle coincidence detection techniques as an analysis tool. The experiment is interpreted with an improved version of our modeling approach elaborated for the first time with iodomethane [31]. The present results provide the basis for the understanding of the molecular mechanisms of the radiosensitizing capacity of 5-IU or any similar radiosensitizing molecules, as well as the basis for single-shot bio-imaging with XFEL.

II. RESULTS

Here, we report the results of the ion momentum imaging measurements for 5-IU using the intense x-ray pulses at the SACLAXFEL facility. The fragment ions resulting from the destruction of the sample, by multiphoton x-ray absorption, were detected by a multicoincidence recoil ion momentum spectrometer [36,37] [see Fig. 1(a)], which provides the mass-to-charge ratios of all detected ions (via their flight time toward the detector), as well as the full reconstruction of their three-dimensional momenta. Let us first review the fragmentation pattern of 5-IU. Figure 1(b) shows the ion time-of-flight (TOF) spectrum of 5-IU, where it can be seen that (i) atomic fragments dominate the spectrum, (ii) doubly charged ions such as C$^2+$, O$^2+$, N$^2+$, as well as multiply charged iodine ions $I^{q+}$ ($q = 1–4$), are formed, and (iii) singly charged polyatomic fragments, CC+$, CN+$, CO+, CNH+, COH+, are also present, albeit with quite small abundances. The x-ray energy of 5.5 keV used in this experiment can ionize electrons from and above the L shell of iodine with a much higher cross section than those from the K shells of light elements, making iodine the dominant x-ray absorber in 5-IU.

Regarding x-ray absorption, iodine behaves very similarly to its neighboring element xenon, where a prior XFEL experiment saw the creation of charge states up to $+26$ [27]. In the case of the smallest iodine-containing organic molecule, CH$_3$I, iodine ions up to $q = +15$ and carbon ions up to $q = +4$ were observed at similar XFEL conditions [31]. The significantly lower charge states of iodine and carbon atomic fragments from the larger molecule 5-IU suggest a very efficient redistribution [38] of the charges from the iodine to the entire molecule, over many more constituent atoms.

The dynamics of the absorption, charge redistribution, and early-stage fragmentation leaves an imprint on the final
momentum vectors of the fragment ions. Figure 2 represents this information in a compact form, as the kinetic energy distributions (KEDs) of the fragment ions derived from their observed momenta. Each dissociating 5-IU molecule results in several charged fragments; these were detected in coincidence with each other, allowing us to distinguish processes that produce, for example, C$^+$ together with the I$^+$, or C$^+$ combined with I$^+$, or C$^+$ together with the I$^{4+}$ ions.

In the kinetic energy analysis, the momentum correlation of the coincident ions is not revealed, but let us consider the angles formed between the momenta of the atomic fragments in the Coulomb explosion. The cosine of the angle, between the momenta of the two ionic fragments A and B, is

$$\cos(\theta) = \frac{\vec{p}_A \cdot \vec{p}_B}{|\vec{p}_A||\vec{p}_B|}. \quad (1)$$

We denote the distributions of $\cos(\theta)$ between the momenta of various ejected ion pairs as $\mathrm{SP}_2(A, B)$, and they are shown in the left column of Fig. 3. The observed maxima in the distributions $\mathrm{SP}_2(I^{4+}, H^+)$, $\mathrm{SP}_2(I^{4+}, O^+)$, and $\mathrm{SP}_2(I^{4+}, N^+)$ correspond well to the angles in the equilibrium geometry of the parent 5-IU molecule, and they are insensitive to the charge state of the coincident iodine. In contrast, $\mathrm{SP}_2(I^+, C^+)$ has a maximum at around 60°, which tends to disappear when the iodine charge state increases and the angle between iodine and carbon ions becomes larger.

During the Coulomb explosion of 5-IU, almost always more than two ions are created. Up to now, we have studied them in pairs. Investigating triple coincidences allows us to extract even more detailed information on momentum correlations. In order to see how well the momenta of three ions A, B, and C, lie on a molecular plane, a triple product was calculated from the observed momenta, giving the cosine of the polar angle:

$$\cos(\phi) = \frac{(\vec{p}_A \times \vec{p}_B) \cdot \vec{p}_C}{|\vec{p}_A \times \vec{p}_B||\vec{p}_C|}.$$ \quad (2)

The experimental distributions, denoted as $\mathrm{SP}_3(A, B, C)$, are shown in the left column of Fig. 4. Note that the distribution function $\mathrm{SP}_3$ for three randomly oriented and uncorrelated vectors is flat against $\cos(\phi)$, so the curves presented directly reflect the deviation from isotropic emission. Coulomb explosion that occurs preferentially in the plane then results in the distributions of $\mathrm{SP}_3(A, B, C)$ peaked at $\cos(\phi) = 0$. In Fig. 4, we chose the combinations $\mathrm{SP}(I^{4+}, H^+, x)$, where x stands for the H$^+$, O$^+$, N$^+$, or C$^+$. $I^{4+}$, H$^+$, N$^+$, and O$^+$ tend to be ejected preferentially in the plane, but this is not necessarily the case for C$^+$.

### III. DISCUSSIONS

In order to interpret the experimental results and quantify the underlying dynamics, we carried out classical MD simulations of the Coulomb explosion within a parametric model of charge buildup and redistribution. The model follows what was developed for the analysis of the CH$_3$I molecule [31], keeping up with its original purpose of being easily extendable to various systems. The model appropriately considers the effects of multiple photon absorption within the XFEL pulse duration (about 10 fs) by the deep atomic shells of iodine. Concomitant to this absorption, but also partly subsequent to it, Auger cascades are the dominant mechanism of charge multiplication at an exponentially decreasing rate. We modeled these charge-creation processes statistically by assuming an increase of the total charge of the molecule ($Q^{\text{tot}}$) with time $t$ (measured from the onset of the XFEL pulse) as $Q^{\text{tot}}[t] = Q^{\text{max}}[1 - \exp(-t/\tau)]$, where $Q^{\text{max}}$ is the final charge. Charge redistribution from iodine to the uracil ring (C$_4$H$_3$N$_2$O$_2$) is described by the rate equation $dQ^{\text{ring}}[t]/dt = RQ^{\text{ring}}[t]$, with a constant charge redistribution rate $R$. Here, $Q^{\text{ring}}[t]$ and $Q^{\text{tot}}[t]$ represent the charge of the uracil ring and iodine at $t$, respectively. This model describes the initial charge creation and the subsequent charge redistribution over the molecule as a result of, e.g.,
molecular Auger decay that occurs during the later stages of the Auger cascades. The two empirical constants $\tau$ and $R$ can be adjusted to best describe the experimental results.

As an important development of our earlier model, we now include molecular vibrations as dependent on the internal temperature $T$. First, molecules with randomly selected atomic velocities were prepared. Then, ion trajectories in the Coulomb explosion were obtained numerically using Coulomb repulsion forces between ions, which are regarded as point charges with the charge development depicted by the above model. The effect of covalent bonding in the 5-IU molecule was neglected. Thus, the di- and polyatomic fragments were not taken into account in the simulation. As shown in Fig. 1(b), complete fragmentation into atomic ions is the dominant process. A large number (1000 runs for each $Q_{\text{max}}$) of trajectories were simulated for each set of parameter combinations of $\tau$, $R$, and $T$, and for a given charge $Q_{\text{max}}$, differing by the random set of initial atomic velocities. Thousands of different sets of initial conditions afforded simulated statistical distributions, which can be directly comparable with the experimental ones, as seen in Figs. 2–4.

As seen in Fig. 2, the experimentally observed broadening of KEDs is well reproduced by the MD simulations (red solid lines). The MD simulations confirm that the broad widths of KED mainly come from the different charge distributions on atoms. The results of the MD simulations for KEDs without consideration of the effect of charge buildup and charge redistribution are shown in Fig. 2 by blue dotted lines. One can see that these simplified simulations fairly reproduce the experimental KEDs of O$^+$ and N$^+$, although they provide discrepancies for H$^+$ and C$^+$, where they overestimate the kinetic energies. In the case of H$^+$, this discrepancy can be directly attributed to the neglect of charge buildup time; in reality, the very mobile protons escape before the full charge buildup is reached.

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FIG. 2. Kinetic energy distribution (KED) of fragment ions. (a) KED of H$^+$ emitted from a XFEL irradiated 5-iodouracil and detected in coincidence with iodine ions. (b) KED of O$^+$, (c) KED of N$^+$, and (d) KED of C$^+$. Symbols are the experimental data, and lines are the result of MD simulations at $T = 300$ K. Dotted blue lines and thin solid green lines correspond to charge buildup times $\tau = 0$ and 10 fs, respectively, assuming that the charge redistribution is instantaneous. The thick red lines represent a modeling with $\tau = 10$ fs and $R = 0.5$ fs$^{-1}$.
[31]. Indeed, taking into account a realistic charge buildup time constant of 10 fs improves the agreement dramatically (thin green solid lines in Fig. 2), especially for coincidences of H\(^+\) with the low-charge states of iodine. Thus, the KEDs of H\(^+\) are particularly sensitive indicators of early molecular and charge dynamics.

Finally, the charge redistribution rate constant \( R \) of 0.5 fs\(^{-1} \) was included in the most realistic model (thick red solid lines). For the heavier ions, the simulated KEDs are less sensitive to the model parameters. These ions travel only over short distances during the charge buildup, and thus most of their kinetic energy is accumulated after the

![Graphs showing distributions of normalized scalar product of momentum vectors, \( \text{SP}_2(A, B) \), plotted as a function of \( \cos(\theta) \).](image)

FIG. 3. Distributions of normalized scalar product of momentum vectors, \( \text{SP}_2(A, B) \), plotted as a function of \( \cos(\theta) \). (a) \( \text{SP}_2(A, B) \) for the (I\(^+\)-H\(^+\)) pair, (b) \( \text{SP}_2(A, B) \) for (I\(^q=1\)-N\(^+\)), (c) \( \text{SP}_2(A, B) \) for (I\(^q=1\)-O\(^+\)), and (d) \( \text{SP}_2(A, B) \) for (I\(^q=1\)-C\(^+\)). Gray shadow areas indicate the angles of atoms in the neutral parent molecule. (e)--(h) \( \text{SP}_2(A, B) \) calculated with temperature, charge buildup, and charge redistribution taken into account, using the parameters \( T = 300 \) K, \( \tau = 10 \) fs, and \( R = 0.5 \) fs\(^{-1} \).
final charge is reached, in contrast to the H⁺ case. Thus, it is
difficult to extract information on charge dynamics from
the KEDs of ions heavier than H⁺. The remaining discrep-
cancies in the KEDs may come from neglecting the
molecular bonds, such as initial carbon departure as dimers.
Also, the smooth charge buildup model is best suited for
highly charged particles and can lead to discrepancies in the
case of very low charge states [31]. The discrepancy between
the measured and simulated KEDs is most significant for the
C⁺ case. It may be worth noting that three of the carbon
atoms are close to the iodine atom, forming part of the uracil
ring. Such geometrical factors neglected in the present model
may also account for the discrepancy.

Previously, Motomura et al. demonstrated that the
momentum correlations are well suited for the reconstruction
of early molecular dynamics [31]. Simulated SP₂(A, B) and
SP₃(A, B, C) are displayed in the right-hand side of Figs. 3
and 4, respectively. We found good agreement between the
experiment and MD results with T = 300 K, τ = 10 fs, and
R = 0.5 fs⁻¹ (see Ref. [39] for many more parameter
combinations). These values are close to the ones for
CH₃I (τ = 9 fs and R = 0.37 fs⁻¹ [31]), suggesting that
the model employed is indeed of general applicability. It may
be worth noting also that these values of R are comparable to
the Auger decay rates of individual decay stages in Xe, as
calculated by Son and Santra [40]. The details of parameter
dependence of SP₂(A, B) and SP₃(A, B, C) are presented in
Ref. [39], but it is worth pointing out here that the internal
temperature also strongly affects the correlated motion of the
particles in the molecular Coulomb explosion.

Our results from both experiment and modeling show
that there is a clear difference in the behavior of carbon ions
compared to oxygen, nitrogen, and also hydrogen ions. All
the latter exhibit a good angular correlation with the
iodine’s momentum that is consistent with the initial
geometry of 5-IU (Fig. 3); they also show a marked
tendency of planar emission together with I⁺⁺ and H⁺
(Fig. 4). Hydrogen and oxygen ions, being attached to the
ring, experience mostly radial forces. Also, the nitrogen
ions experience outward, mostly planar, forces after the
definite departure of protons. Carbon ions, however, are
surrounded by elements of equal or larger mass, and the
forces on some of the carbon ions, such as C₁ and C₂, are
not necessarily outwards. Such a typical example is shown
in Fig. S7 of Ref. [39]. This may explain a peculiar
behavior in the observed momentum correlations in which
the momentum of the carbon ion is involved.

Since we validated the model by comparison with the
experimental observables, we can now use it to give
insights into motions of the individual atoms in real space
that may deteriorate x-ray scattering signals necessary for
the structure retrieval of the original object with XFEL. The
simulated evolution of interatomic distances using
realistic charge dynamics is displayed in Fig. 5. The results
are an average of over 1000 runs for each Q²max, also
weighted by the charge state distribution of parent molecu-
lar ions, estimated from our CH₃I experiment [31]. MD
results reveal that after 10 fs (the XFEL pulse duration), the
H-C and H-N distances increase by 50%–100%, whereas
the increases of the O-C, C-N, and C-C bonds are of only a
few percent; it takes more than 10 fs for heavy atoms,
except for H, to move 10 pm. The smallest change takes place in the C-I distance and is due to the large mass of iodine and the fact that the Coulomb forces acting on C$_1$ tend to cancel out at the early stages of the explosion. These changes are much smaller than what would be expected from instantaneous charge creation (Ref. [39], Fig. S7).

### IV. CONCLUSION

We carried out recoil-ion momentum imaging measurements for 5-iodouracil (IU) using intense XFEL pulses at SACLA. The parametric, classical Coulomb explosion model, which includes both charge and nuclear dynamics, succeeded in reproducing the experimental observations. We found that both the charge buildup and redistribution take place within the XFEL pulse duration of 10 fs, at the early stages of the Coulomb explosion. Good agreement between experiment and MD results is found when selecting an internal temperature, charge buildup time, and charge redistribution rate of 300 K, 10 fs, and 0.5 fs$^{-1}$, respectively. The present values of charge buildup time $\tau$ and charge redistribution rate $R$ are very close to those we obtained for CH$_3$I ($\tau = 9$ fs and $R = 0.37$ fs$^{-1}$ [31]). The time dependence of the interatomic distances, deduced from the MD simulations, suggests that within 10 fs, the displacements are large for the hydrogens and are considerably smaller for the other heavier atoms. This result, obtained on a realistic, biologically relevant molecule, gives two important indications, which, being independent of the exact composition of the studied nucleobase, are both of very general relevance to other systems of this kind.

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### APPENDIX: MATERIALS AND METHODS

1. **Experimental methods**

The experiment was carried out at the experimental hutch 3 (EH3) of beam line 3 (BL3) of SACLA [41]. The XFEL beam is focused by the Kirkpatrick-Baez (KB)
mirror system to a focal size of 1 μm (FWHM) in diameter [42].

The photon energy was set at 5.5 keV and the photon bandwidth was about 60 eV (FWHM). The repetition rate of the XFEL pulses was 10 Hz. The pulse length was not measured but was estimated to be about 10 fs (FWHM) [43]. XFEL pulse energies were measured by the beam-position monitor [44] located upstream of the beam line. The monitor was calibrated by a calorimeter [45] so that output signals from the monitor could be transformed into the absolute value of the pulse energy, which was 235 μJ on average. The relative x-ray pulse energy passing through the interaction point was also measured shot-to-shot by a p-intrinsic-n (PIN) photodiode. The shot-to-shot pulse energy fluctuation was about 7% (14% FWHM). The peak fluence was 26 μJ/μm² on average. The absolute value of the peak fluence was calibrated just after the experiment using Ar [27].

The 5-iodouracil (>99.0%) was purchased from Tokyo Chemical Industry Co., Ltd. and used without further purification. The solid sample was heated up to 190°C in the reservoir of a modified electromagnetic solenoid valve [46]. The resulting sample vapor was seeded by helium gas and introduced to the focal point of the XFEL pulses in the ultrahigh-vacuum reaction chamber as a pulsed supersonic gas jet. Any change of sample due to heating was not confirmed by 1H NMR measurements.

The ions were detected by a multicoincidence recoil ion momentum spectrometer [36,37] to measure three-dimensional momenta of each fragment ion. The molecular beam was crossed with the focused XFEL beam at the reaction point, and the emitted ions were projected by electrostatic fields onto an 80-mm-diameter microchannel plate (MCP) detector, in front of a delay-line anode. The three-dimensional momentum vectors were determined using both the TOF and the detector-hit position for each reaction point, and the emitted ions were projected by the ion spectrometer [36,37] to measure three-dimensional momentum vectors. The molecular fluence was negligible for the coincidence analysis.

The observed particle hit rate was kept at less than 0.1 events per FEL shot during experiments. False coincidence was negligible for the coincidence analysis.

2. MD simulations

Classical molecular dynamics simulations were carried out, where we evaluated the Coulomb repulsive force between ions during ion trajectory calculations, while the effects of covalent bonds in the 5-IU molecule were neglected. We traced the trajectories of the emitted ions to up to 10 ps with a time step of 0.2 fs. The ensemble of initial positions and velocities of atoms in the parent molecules was prepared by B3LYP/3-21G(d) quantum-chemical calculations using the GAMESS package [47]. Thermal vibration effects were taken into account using Nose-Hoover thermostat methods implemented in the GAMESS package.

For each trajectory of the MD simulations, the final charge state of parent 5-IU molecules, Q\textsuperscript{max}, was chosen according to the experimental charge distribution of the CH$_3$I molecules [31] that was measured under the same XFEL conditions. We also determined the final charge of each atom in such a way that the charges are randomly distributed over the atoms in the parent molecule. In addition, we assumed that the iodine ions take at least +1 charge. The time evolution of the atomic charges was governed by the equations given in Sec. III. We carried out numerous MD simulations by scanning the parameters \( r \), \( R \), and \( T \), as presented in Ref. [39].


