Comparing Vacuum and Extreme Ultraviolet Radiation for Postionization of Laser Desorbed Neutrals from Bacterial Biofilms and Organic Fullerenes

Gerald L. Gasper\textsuperscript{a}, Lynelle K. Takahashi\textsuperscript{b,c}, Jia Zhou\textsuperscript{c}, Musahid Ahmed\textsuperscript{c}, Jerry F. Moore\textsuperscript{d}, and Luke Hanley\textsuperscript{a,\ast}

\textsuperscript{a}Department of Chemistry, University of Illinois at Chicago, 845 W. Taylor Street (m/c 111), Chicago, IL 60607, USA
\textsuperscript{b}Department of Chemistry, University of California, Berkeley, Room 419 Latimer Hall, Berkeley, CA 94720-1460, USA
\textsuperscript{c}Chemical Sciences Division, Lawrence Berkeley National Laboratory, 1 Cyclotron Road, Berkeley, CA 94720, USA
\textsuperscript{d}MassThink LLC, 500 East Ogden Avenue Suite 200, Naperville, IL 60563
\textsuperscript{\ast}Corresponding author: Tel.: 312.996.0945; fax: 312.996.0431. E-mail: lhanley@uic.edu

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Abstract
Vacuum and extreme ultraviolet radiation from 8 - 24 eV generated at a synchrotron was used to postionize laser desorbed neutrals of antibiotic-treated biofilms and a modified fullerene using laser desorption postionization mass spectrometry (LDPI-MS). Results show detection of the parent ion, various fragments, and extracellular material from biofilms using LDPI-MS with both vacuum and extreme ultraviolet photons. Parent ions were observed for both cases, but extreme ultraviolet photons (16 - 24 eV) induced more fragmentation than vacuum ultraviolet (8 - 14 eV) photons.
1. Introduction

Vacuum ultraviolet (VUV) single photon ionization (SPI) of laser or ion desorbed neutrals has demonstrated sensitive detection and the capability for mass spectrometric imaging of atomic and molecular analytes [1-4]. Laser desorption VUV postionization mass spectrometry (LDPI-MS) of biofilms showed improved sensitivity above 8 eV photon energy with optimal signal to noise at 10.5 eV, given constant photon flux [5]. By contrast, multiple photon ionization of fullerenes and organic compounds showed increased spectral complexity [6,7] with the absorption of many photons imparting excess energy and causing an inability to mitigate channels of dissociation and ionization [1]. Prior researchers argue that extreme ultraviolet (EUV) radiation near 26 eV may reduce fragmentation for single photon ionization of van der Waals and H-bonded clusters [8,9,10,11] with spectra appearing similar to those obtained at 10.5 eV [12]. Yet, considerable molecular fragmentation of small molecules was present in these investigations [8,9,10]. The role of protonated clusters in these experiments to enhance ionization, reduce fragmentation, or facilitate cooling is unclear [11]. LDPI-MS data was collected here to compare postionization by VUV and EUV radiation. Two different thin film samples were examined: a pure fullerene derivative and a bacterial biofilm with dissolved antibiotic rifampicin.

2. Experimental

Studies were performed with a modified commercial secondary ion mass spectrometer (TOF.SIMS 5, ION TOF Inc., Münster, Germany) coupled to a 349 nm Nd:YLF desorption laser (Explorer, Newport Corp, 5 ns pulse length, 300 & 25 µm spot, 2 - 100 MW/cm²) and tunable, monochromatized 8 - 24 eV synchrotron radiation at the Chemical Dynamics Beamline of the Advanced Light Source (Lawrence Berkeley National Laboratory, Berkeley, CA). Mass spectrum were averaged from 80,000 to 200,000 laser shots. The experimental apparatus and parameters were similar to those described previously [3,5,13]. [6,6] diphenyl C₆₂ bis(butyric acid methyl ester) (C₆₂, mol. wt. 1100 Da, Sigma-Aldrich) was deposited on coated glass microscope slides of indium tin oxide (ITO, Sigma-Aldrich). Biofilms were prepared from bacterial stock solutions of Staphylococcus epidermidis (ATCC 35984, Manassas, VA, USA) [14]. Drip flow biofilms were grown on ITO slides by methods previously described [5,15]. Each sample was scanned for direct ionization by laser desorption (LD, no VUV) and synchrotron
background (SB, no LD). EUV spectra were not normalized to the beam flux to account for differences in synchrotron fill and beamline transmission at different wavelengths and hence an absolute comparison to the VUV data cannot be performed.

3. Results & Discussion

LDPI-MS of neat [6.6] diphenyl C$_{62}$ bis(butyric acid methyl ester), referred to here as C$_{62}$, are displayed in Figure 1, using both VUV and EUV single photon ionization. The parent ion peak (M$^+$) at m/z 1100 was clearly observed at all six photon energies in the VUV and EUV. Three known fragments included m/z 190, 720, and 910, which correspond to the loss of one or two of the 190 Da side chains to the 720 Da C$_{60}$ core, as well as the core itself (chemical structure shown in Figure 2). Fragmentation of the ligand becomes more apparent at higher photon energies as the abundance of peaks from m/z 57 to 175 increase with respect to m/z 190. Prominent peaks at m/z 648 and 660 were observed at all photon energies, including at 7.87 and 8.5 eV (data not shown), indicating they are probably impurities of C$_{62}$. Mass resolution was ~1100 at m/z 720 (C$_{60}$) in the VUV and limited by the temporal energy distribution of ion extraction through a continuous source. Neither direct ionization by laser desorption (LD, no VUV) and nor synchrotron background signal (SB, no LD) were observed.

Parent to fragment ion ratios (M/F) for m/z 190, 720, and 910 for the data from Figure 1 and another data set are shown in Figure 2. All M/F ratios decreased as the photon energy increased. From 10.5 to 24 eV, all three M/F values reduce to ~33%. The relative abundance of m/z 910 was about 3 times less than either m/z 190 or 720 at most photon energies.

Rifampicin was doped at a concentration of 120 µM into a bacterial biofilm and measured by EUV LDPI-MS. Figure 3 shows a comparison of the result to similar spectra recorded at VUV (8.0 to 12.5 eV) photon energies [5]. The rifampicin parent ion at m/z 823 was barely observable above the noise in the EUV spectra, while it was readily observed in prior VUV spectra. Fragment ions at m/z 84, 284, 510, 524, 641, and 655 (indicated by asterisks) were observed at all three EUV photon energies, as previously identified by VUV LDPI-MS. Mass resolution was between 200 and 400 at major rifampicin fragment peaks.

Peaks observed from m/z 1000 - 1400 derived from the biofilm response to rifampicin were also observed by both EUV and VUV LDPI-MS, as previously discussed [5]. Increasing
the photon energy from 20 to 24 eV enhanced the signal to noise ratio for these peaks. Mass resolution ranged from 100 to 300 at m/z 1157 with the best resolved peaks at 24 eV.

Table 1 shows parent to fragment ion ratios (M/F) for LDPI-MS of rifampicin doped biofilm at known antibiotic fragments m/z 84, 284, and 655 at 10.5 and 20 eV photon energies. Sensitivity decreases rapidly above m/z 800 at 20 eV and is reflected in the M/F values. Table 1 shows the abundance of fragments increased 10 - 200 times from 10.5 to 20 eV. Yet, fragments m/z 84 and 284 increased and remain prominent at 20 eV. Studies by Nuutinen et al. suggest this may result due to stabilization by a piperazine group, represented in these two rifampicin fragments, and dissociation from its methyl group [16]. Similar VUV work with neat rifampicin observed the prominence of fragments containing methylated piperazine [5].

4. Conclusion

Single photon ionization with EUV radiation was effective for the two organic/biological systems examined here, but did induce more fragmentation compared with VUV radiation. However, fragmentation is sometimes an unavoidable consequence of the laser desorption process, and addition of matrix compounds or other methods for cooling and stabilizing the desorbed species prior to postionization might mitigate this effect. Further work is clearly required to explore whether the enhanced fragmentation observed here with EUV radiation occurs in other examples of laser desorption and if any such enhancement is offset by the overall increase in ionization efficiency for EUV vs. VUV radiation.

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References


FIGURES & CAPTIONS:

Figure 1: LDPI-MS with VUV (<15 eV) as well as EUV photon energies (>15 eV) photon energies of neat C\textsubscript{62} displaying parent ion (M\textsuperscript{+}). Asterisks indicate known fragments at m/z 190, 720, and 910, and circles indicate M\textsuperscript{+}. EUV spectra were scaled by 10\texttimes. Synchrotron background (SB) was recorded at 20 eV.

Figure 2: Parent to fragment ion ratios (M/F) for known C\textsubscript{62} fragments at m/z 190 (C\textsubscript{12}H\textsubscript{24}O\textsubscript{2}), 720 (C\textsubscript{60}), and 910 (C\textsubscript{60} + 190).
Figure 3: LDPI-MS with EUV radiation (16, 20, and 24 eV) of 120 µM concentration of rifampicin doped biofilm. Synchrotron background (SB) was recorded at 24 eV. Fragment ions are denoted by asterisks.

Table 1: Values of parent to fragment ion ratios (M/F) for LDPI-MS at 10.5 and 20 eV of rifampicin doped biofilm using fragments m/z 84, 284, and 655.

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<th>M/F</th>
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<tr>
<td></td>
<td>$\bar{X}$</td>
<td>$\sigma$</td>
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<td>M/84</td>
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<tr>
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<td>0.18</td>
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