Supplementary Materials

Photodegradation of Riboflavin under Alkaline Conditions: What can Gas-Phase Photolysis Tell Us About What Happens in Solution?

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Supporting Information:

S1. Photodepletion laser power dependence measurements

S2. Electron detachment yield vs. photodepletion yield interpretation

S3. Higher-energy collisional dissociation of [RF-H]⁻

S4. Solution-phase photofragment production curves

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S1. Photodepletion laser power dependence measurements

Laser power measurements were conducted on $[RF-H]^-$ at several of its respective absorption maxima when electrosprayed in deionized H₂O. The plot displayed in Figure S1 shows the parent ion photodepletion intensities (Int_{OFF} – Int_{ON}) at the selected photon energies. Following standard protocols, the data has been plotted and fit to a power function.[1–3] The resultant slope is thereby proportional to the number of absorbed photons implicated in the experiment(s).



Figure S1. Power dependence measurements for [RF–H]⁻ at five absorption maxima of (**a**) 2.5 eV (496 nm), (**b**) 3.0 eV (414 nm), (**c**) 3.5 eV (354 nm), (**d**) 4.7 eV (264 nm), and (**e**) 5.0 eV (246 nm).

Multiphoton events *via* instantaneous absorption of multiple photons in the Franck-Condon region are negligible as the laser beam is only softly focused through the ion-trap region. The slopes of all photon energies are less than 1.0, confirming that photodepletion $[RF-H]^-$ at 0.1 mJ are evidently not multiphoton in nature.

S2. Electron detachment yield vs. photodepletion yield interpretation

In Figure S2 where we present ED* spectra, we overlay such data with the photodepletion yield (PD*) for ease of comparison; PD* is the normalized photodepletion ion count (Eq. 6), which provides the most straight-forward comparison to ED* (Eq. 5). Previous work on the vertical detachment energies (VDEs) of flavins have shown predicted VDEs of *ca*. 4.0 and 3.8 eV for deprotonated alloxazine and *ca*. 4.6 and 4.7 eV for deprotonated structures of lumichrome [4] with deprotonation on the ribose side chain likely giving a VDE close to that of an alkoxide [5].



Figure S2. (a) Electron detachment yield (ED*; blue) *vs.* photodepletion yield (PD*; red) of [RF–H]⁻. The solid lines are a five-point adjacent average of the data points.

S3. Higher-energy collisional dissociation of [RF-H]⁻



Figure S3. Parent ion dissociation curves $[RF-H]^-$ (*m/z* 375) and the extent of the minor thermal fragments produced between 0% and 60% HCD energy, as electrosprayed in EtOH. The curved lines are a five-point adjacent average of such data points and are provided as a viewing guide, to emphasize the profile for each individual fragment.

S4. Solution-phase photofragment production curves



Figure S4. (a) Relative ion intensity plot highlighting the solution-phase photofragment ions of aqueous RF produced over a 15 minute interval of irradiation at 365 nm when delivered *via* ESI-MS in the negative ion mode. (b-c) Linked inset plot highlighting the relative ion intensities of the minor photofragments observed upon the solution-phase irradiation of aqueous RF. The curved lines included with the data points are a three-point adjacent average of such data points and are extracted from an average of 3 repeat runs.

S5. Design of custom-made on-line syringe cell



Figure S5. Various cross-sectional views of the custom-made on-line syringe photolysis cell.

S6. Control solution-phase studies



Figure S6. Static UV-visible absorption spectra monitoring the absorption spectrum of riboflavin, prepared in aqueous solution with trace amounts of NH_3 (0.4% v/v), between 0–240 min. The overall stability of the spectra depicted support the idea that hydrolysis does not occur on this time scale.



Figure S7. Static UV-visible absorption spectra monitoring the changes in the absorption spectrum of riboflavin, prepared in aqueous solution with trace amounts of NH_3 (0.4% v/v), between 0–20 min upon irradiation (365 nm; 50% light int.).



Figure S8. Static UV-visible absorption spectra monitoring the changes in the absorption spectrum of riboflavin, prepared in aqueous solution, between 0–20 min upon irradiation (365 nm; 50% light int.).



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Figure S9. Positive ion mode electrospray ionization (ESI) mass spectrum of protonated riboflavin $([RF+H]^+)$ at m/z 377. Note that the ions at noted at m/z 399 and 415 represent its $[RF+Na]^+$ and $[RF+K]^+$ systems, respectively.



Figure S10. Representative total ion chromatogram and extracted ion chromatogram for ions at m/z 751, 617, 375, 255, and 241. Note that the ion intensities of the m/z 751, 617, 375, and 255 ions barely fluctuate relative to one another across the 0–30 min time period of irradiation. Notably the production of the major photofragment ion in the solution phase at m/z 241 (as identified previously in Figures 8 and 9 of the main text) is not induced when the syringe is covered, and where UV-A light is not transmitted through to the solution.



Figure S11. Relative ion intensity plot highlighting the solution-phase ions of aqueous $[RF-H]^-$ produced over a 30 minute interval of irradiation at 365 nm (3.4 eV) when delivered via ESI-MS in the native ion mode when the syringe is covered with a black cloth, preventing UV-A light to be transmitted into the solution for irradiation. The curved lines included with the data points are a three-point adjacent average of such data points and are extracted from an average of 3 repeat runs.



Figure S12. Plot of the data obtained *via* use of a thermocouple (Six Channel Handheld Temperature Data Logger; Omega RDXL6SD, Serial No. 003308) employed to monitor the temperature within the photolysis cell at two separate locations within the on-line photolysis cell.

References

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