Measurement of the Population of Electrosprayed Deprotomers of Coumaric Acids Using UV–Vis Laser Photodissociation Spectroscopy

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ABSTRACT: The measurement of deprotonation sites in multifunctional molecules following electrospray ionization is important to better inform a wide range of spectroscopic and photophysical studies that use electrospray to prepare molecular species for study in the gas phase. We demonstrate that low-resolution UV–vis laser photodissociation spectroscopy can be applied in situ to identify the deprotomers of three coumaric acids, trans-para-coumaric acid (CMA), trans-cafeic acid (CA), and trans-ferulic acid (FA), formed via electrospray. Electronic absorption spectra of the deprotonated coumaric acids are recorded via photodepletion and photofragmentation following electrospray from solutions of ethanol and acetonitrile. By comparing the experimental spectra to wave function theory calculations, we are able to confirm the presence of phenoxide and carboxylate deprotomers upon electrospray for all three coumaric acids, when sprayed from both protic and aprotic solvents. Ratios of the phenoxide:carboxylate deprotomers are obtained by generating summed theoretical absorption spectra that reproduce the experimental spectra. We find that choice of electrospray solvent has little effect on the ratio of deprotomers obtained for deprotonated CMA and CA but has a greater impact for FA. Our results are in excellent agreement with previous work conducted on deprotonated CMA using IR spectroscopy and demonstrate that UV photodissociation spectroscopy of electrosprayed ions has potential as a diagnostic tool for identifying deprotomeric species.

1. INTRODUCTION

The influence of electrospray ionization (ESI) conditions on the location of protonation and deprotonation sites of electrosprayed ions is a topic of keen debate.\textsuperscript{1–13} Acid–base reactions are of key importance throughout chemistry and biology, so correctly identifying the structures of protomers and deprotomers can be crucial to understanding reactive processes. While ESI has been successfully employed across analytical chemistry for many years, it is increasingly being used to probe solution-phase reactions and reactive intermediates for both chemical and biochemical systems.\textsuperscript{8,11} The role of the electrospray process in determining the location of protonation and deprotonation sites is therefore of key chemical interest.

Roithova and co-workers performed what is perhaps the seminal investigation of how the electrospray process affects the gas-phase ratios of deprotonated isomers by studying the para-hydroxybenzoic acid molecule.\textsuperscript{13} NMR was used to probe the solution-phase structures and ion-mobility mass spectrometry (IM-MS) was used to identify the gas-phase isomers. They were able to show that while the carboxylate isomer is preferred in solution irrespective of the solvent, the opposite is true for the gas-phase isomers. However, the exact ratio of isomers formed in the gas phase was found to depend strongly on the ESI solvent, pH, and solution concentration.\textsuperscript{5,7,11,13} These results led Roithova and co-workers to conclude that the gas-phase populations do not accurately reflect solution-phase populations, a conclusion that has been confirmed by a number of subsequent studies. As a consequence, it is very important to have tools available to determine accurately the identity of protonation/deprotonation isomers and also to have a full understanding of how the gaseous population relates to the solution-phase population as a function of the experimental conditions. Toward this end, recent studies have investigated the possibility of applying in situ spectroscopy to identify correctly electrosprayed protomers or deprotomers.\textsuperscript{2,4,6,8,13}

In this work, we present a combined UV–vis laser photodissociation spectroscopy and quantum chemical study of a series of electrosprayed deprotonated coumaric acids (Scheme 1): namely, trans-para-coumaric acid (CMA), trans-cafeic acid (CA), and trans-ferulic acid (FA). The para-
Scheme 1. Schematic Diagram of (1) trans-para-Coumaric Acid (CMA), (2) trans-Caffeic Acid (CA), and (3) trans-Ferulic Acid (FA)

hydroxycinnaminate unit is widespread in nature, being found as the chromophore in photoactive yellow protein and as sunscreen molecules in plants. Given their biological importance, deprotonated coumaric acids have been the focus of a number of recent gas-phase studies investigating their intrinsic photophysics and photochemistry. Here, our goal is to identify which deprotomers are present as a function of electrospray solvent through applying a combination of theory and experiment. We also aim to assess the potential of low-electrospray solvent as an analytical tool for identifying gaseous deprotomers. UV laser photodissociation spectroscopy should be highly sensitive to the electronic chromophore of the coumaric acid moiety, thus allowing us to identify unambiguously which isomers were present in those previous studies, and for electrosprayed coumaric acids generally.

CMA, CA, and FA are widely used as antioxidant agents in the cosmetic industry, leading to their long-standing use in antiaging cosmetics. They also absorb strongly in the UVA (320–380 nm) and UVB (320–280 nm), making them viable sunscreen agents. Antioxidants have considerable synergistic potential as components of sunscreen formulations since, in addition to their filter in Japan. Indeed, FA is already approved as a UV filter in Japan. The potential utility of coumaric acids as sunscreens has led to a number of studies of the fundamental photophysics of CMA to explore its excited state nonradiative relaxation pathways. Laser spectroscopic studies on the deprotonated forms of the CA and FA antioxidants are sparse, with only solution-phase transient absorption studies having been conducted to date. This is despite the fact that sunscreens are exposed to varying pH environments in common usage, with seawater and salt water lakes being alkaline, for example. Recent photochemical studies on how pH environment could affect the sunscreen performance at the molecular level have demonstrated that the deprotonated forms of oxybenzone, 2-phenylbenzimidazole-5-sulfonic acid, and benzophenone-4 are all able to photogenerate free radicals species via active photodecay channels in the gas phase. It is therefore important to investigate the deprotonated forms of the antioxidants studied here in the same context.

2. METHODS

Gas-phase UV–vis photodissociation experiments were conducted in an AmaZon SL dual funnel ESI quadrupole ion-trap mass spectrometer (Bruker Daltonics Inc., Billerica, MA, USA), which was modified to allow laser-interfaced mass spectrometry (LIMS). This instrument has the advantages of a commercial mass spectrometer, coupled with the ability to record UV–vis photodissociation spectra in a routine manner. trans-para-Coumaric acid (4-hydroxycinnamic acid) and trans-caffeic acid ((2E)-3-(3,4-dihydroxyphenyl)acrylic acid) were purchased from Fluorochem Ltd. (Hadfield, Derbyshire, UK). trans-Ferulic acid ((2E)-3-(4-hydroxy-3-methoxyphenyl)-prop-2-enolic acid) was purchased from Sigma-Aldrich, Inc. (St. Louis, MO, USA). HPLC-grade EtOH and MeCN were purchased from Fisher Scientific, Inc. (Pittsburgh, PA, USA), all used as received. Solutions of CMA, CA, and FA (1 × 10⁻⁵ M) in EtOH or MeCN were introduced into the mass spectrometer by ESI using typical instrumental parameters: nebulizing gas pressure 14.0 psi; injection rate 0.33 mL h⁻¹; drying gas flow rate 10.0 L min⁻¹; and run in the negative ion mode at a capillary temperature of 160 °C to form deprotonated ions. Trace amounts of NH₃ solution (0.4%) was added to aid deprotonation.

Photodepletion (PD) experiments were conducted with an ion accumulation time of 10 ms and a fragmentation time of 100 ms, thereby ensuring that each mass-selected ion packet interacted with one laser pulse to minimize the likelihood of multiphoton events. 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. In the limit where fluorescence is negligible, the UV-excited gaseous ion will fragment upon excited state relaxation, yielding an action absorption spectrum by photo-dissociation. Photodetection (PD) of [CMA-H]⁺, [CA-H]⁺, and [FA-H]⁺ were measured as a function of the scanned wavelength, with the photofragment (PF) production also recorded simultaneously at each wavelength (see eqs 1a–1c):

\[
\text{Photodepletion (PD) intensity} = \frac{\ln (\text{Int}_{\text{OFF}} / \text{Int}_{\text{ON}})}{\lambda \times P} \quad (1a)
\]

\[
\text{Photofragmentation (PF) production intensity} = \frac{\ln (\text{Int}_{\text{PF}} / \text{Int}_{\text{OFF}})}{\lambda \times P} \quad (1b)
\]

\[
\text{Relative ion yield} = \frac{\text{Int}_{\text{PF}}}{\text{Int}_{\text{OFF}}} \quad (1c)
\]

where \(\text{Int}_{\text{off}}\) and \(\text{Int}_{\text{on}}\) are the peak parent ion intensities with laser off and on, respectively, \(\text{Int}_{\text{pf}}\) is the fragment intensity with the laser on, \(\lambda\) is the excitation wavelength (nm), \(P\) is the laser pulse energy (mJ), and \(\text{Int}_{\text{pf}}\) is the sum of the photofragment ion intensities with the laser on. Photodecomposition laser power dependence measurements are available in Section S3. PD intensities were taken from an average of three runs at each scanned wavelength. Fragment ions with \(m/z < 50\) are not detectable in our mass spectrometer since low masses fall outside the mass window of the ion trap.

Electron detachment yield (ED) spectra were calculated by assuming that any depleted ions not detected as ionic photofragments are decaying via means of electron detachment, as determined using eq 2a. This analysis assumes that both the
parent ions and photofragments are detected equally in the mass spectrometer. In the figures where we present ED* spectra (Figures 5 and 6), we overlay such data with the photodepletion yield (PD*). PD* is the normalized photodepletion ion count (eq 2b), which provides the most straightforward comparison to the electron detachment yield (eq 2a).

Electron detachment yield (ED*) = \[ \frac{\text{Int}_{\text{CAB}} - \text{Int}_{\text{CAB}}^*}{\lambda \times P} \] (2a)

Photodepletion yield (PD*) = \[ \frac{\text{Int}_{\text{CAB}} - \text{Int}_{\text{CAB}}^*}{\lambda \times P} \] (2b)

Higher-energy collisional dissociation (HCD) was employed to determine the ground state thermal fragmentation characteristics of [CMA-H]-, [CA-H]-, and [FA-H]- in EtOH and in MeCN, using an Orbitrap Fusion Tribid mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) with an ESI source, run in negative ion mode between 0 and 40% HCD (Thermo Fisher Scientiﬁc, ESI source). A number of minimum-energy deprotomer structures were identiﬁed for each deprotontated carboxylic acid, both for the isolated (gaseous) and solution-phase environments. Table 1 lists the zero-point-corrected gas-phase energies of [CMA-H]-, [CA-H]-, and [FA-H]-.

| deprotomer | gas-phase \( E_{\text{rel}} \) (kJ mol\(^{-1}\)) | VDE (eV) | VDM (D) | EtOH \( E_{\text{rel}} \) (kJ mol\(^{-1}\)) | MeCN \( E_{\text{rel}} \) (kJ mol\(^{-1}\)) |
|------------|-----------------------------------|--------|--------|------------------------------|-------------------------------|
| CMA        | carboxylate                        | 0.0    | 3.73   | 1.93                         | 0.0                           |
|            | phenoxide                         | -41.8  | 2.42   | 3.79                         | +16.3                         |
|            | phenoxyde (para)                  | -81.0  | 2.47   | 3.42                         | +0.4                          |
|            | phenoxyde (meta)                  | -66.5  | 3.02   | 5.01                         | +5.7                          |
|            | carboxylate (C\(_1\))            | 0.00   | 3.70   | 3.64                         | 0.0                           |
|            | carboxylate (C\(_2\))            | +22.2  | 3.75   | 3.94                         | +14.7                         |
|            | carboxylate (C\(_3\))            | -26.6  | 2.23   | 2.70                         | +73.4                         |
|            | phenoxyde (C\(_1\))              | -32.8  | 2.42   | 2.39                         | +30.9                         |
| FA         | carboxylate (C\(_1\))            | 0.00   | 3.70   | 3.64                         | N/A\(^{\dagger}\)             |
|            | carboxylate (C\(_2\))            | +22.2  | 3.75   | 3.94                         | +14.7                         |
|            | carboxylate (C\(_3\))            | -26.6  | 2.23   | 2.70                         | +73.4                         |
|            | phenoxyde (C\(_1\))              | -32.8  | 2.42   | 2.39                         | +30.9                         |
|            | phenoxyde (C\(_2\))              | 0.00   | 3.70   | 3.64                         | N/A\(^{\dagger}\)             |

\(^{\dagger}\)All energies are zero-point energy (ZPE) corrected. VDEs/VDMs are evaluated at the optimized gas-phase geometries. All values are evaluated at the RI-MP2/aug-cc-pVDZ level of theory. \(^{\ddagger}\)VDE = E (neutral at optimized anion geometry) – E (anion). This is included in the table for comparison with the experimental data. \(^{\dagger}\)VDM is the dipole moment of the neutral at the geometry of the optimized anion geometry. \(^{\ddagger}\)Unobtainable as a stable minimum-energy geometry.

3. RESULTS AND DISCUSSION

3.1. Ab Initio Wave Function Theory Calculations of the Deprotomers of CMA, CA, and FA. High-level ab initio wave function theory calculations at the RI-MP2/ADC(2)/aug-cc-pVDZ level were performed on the selected carboxylic acids in the gas phase and solution (EtOH and MeCN) to assign the experimental spectra. A number of minimum-energy deprotomer structures were identified for each deprotontated carboxylic acid, both for the isolated (gaseous) and solution-phase environments. Table 1 lists the zero-point-corrected gas- and solution-phase energies obtained, with the corresponding structures and geometric parameters included in the Supporting Information (Scheme S1 and Tables S1–S9).

We found that the relative energies of the carboxylate and phenoxyde deprotomers were strongly geometry-dependent and, consequently, dependent on the level of theory employed. This is challenging since accurate experimentally determined geometries are not available routinely for these anionic systems. For this reason, we focus here on the qualitative trends predicted by our calculations (Table 1). The calculations suggest that the phenoxyde deprotomers exist at lower relative energies than the carboxylate deprotomers for the gaseous CMA, CA, and FA anions, but this trend inverts in solution (EtOH and MeCN), albeit with smaller energy differences between the two deprotomeric forms.

Given the results of the calculations, we anticipate that both carboxylate and phenoxyde deprotomers of CMA and CA will be produced with approximately equal abundances upon ESI from solutions of EtOH and MeCN. More speciﬁcally, we predict carboxylate:phenoxyde ratios of 1:1 and 1:2 for [CMA-H]- and [CA-H]-, respectively. For [FA-H]-, we do not expect signiﬁcant contributions from the \(\text{C}_3\)-symmetric phenoxyde deprotomer in EtOH or MeCN; this rotamer of the lower-energy \(\text{C}_1\)-symmetric phenoxyde deprotomer exists at considerably higher energy (+73.4 and +61.2 kJ mol\(^{-1}\)) above the
lowest-energy carboxylate deprotomer in EtOH and MeCN, respectively; Table 1. We predict that the remaining carboxylate and phenoxide deprotoners of [FA-H]− produced from EtOH and MeCN occur with approximately abundances of 2:1 in EtOH and 1:1 in MeCN.

3.2. Gas-Phase UV–Vis Absorption Spectra of [CMA-H]−, [CA-H]−, and [FA-H]−. ESI readily produces the deprotonated forms of the coumaric acids [CMA-H]−, [CA-H]−, and [FA-H]− as gas-phase ions with m/z 163, 179, and 193, respectively.

3.2.1. [CMA-H]−. Figure 1a displays the gas-phase photodepletion (PD) spectra of [CMA-H]− across the 2.5–5.5 eV spectral range. The solid blue and red lines in (a) and (c) are five-point adjacent averages of the data points. The theoretical gas-phase absorption spectrum of [CA-H]− displayed in Figure 2b (black line) and was constructed using a 1:1 ratio (i.e., as an equally weighted linear combination) of the two deprotoners of [CMA-H]−. Comparison of the experimental and theoretical spectra clearly shows that both deprotoners must be present in the gas-phase ion ensemble, given that the simulated spectra in the band I and III regions in particular require contributions from both deprotoners. This is in line with the predicted relative energies of [CMA-H]− (Table 1). Band I (λmax ~ 2.9 eV) in the experimental PD spectrum can be unambiguously assigned to a bright π* ← π transition from the phenoxide deprotomer (A′, 2.87 eV), while bands II and III were assigned to π* ← π transitions of the phenoxide (A′, 3.90 eV) and carboxylate (A′, 4.70 eV) deprotomers. Oomens and co-workers have previously used in situ IR spectroscopy of electrosprayed [CMA-H]− to identify its deprotomers and identified similar ratios of carboxylate:phenoxide isomers as we observe here.

We next turn to exploring the photofragment ions that are associated with excited state decay. [CMA-H]− produces a single, dominant ionic photofragment at m/z 119 (Figure 1c) corresponding to the loss of m/z 44 (−CO2) from the parent ion. Oomens and co-workers were able to use in situ IR spectroscopy to identify this fragment as the para-vinyl-phenoxide anion. The production profile of the m/z 119 photofragment peaks at ca. 2.9 eV (eq 3a). Table 2 lists the ionic m/z of the major ionic photofragments of the deprotonated coumaric acids, along with their accompanying neutral fragments. The m/z 119 photofragment was observed following excitation of [CMA-H]− produced via electrospray from both EtOH and MeCN.

\[
\text{[CMA-H]}^- + h\nu \rightarrow m/z \ 119 + \text{CO}_2 \quad \text{(3a)} \\
\rightarrow \text{e}^- + \text{[CMA-H]}^- \quad \text{(3b)}
\]

Additional minor photofragments of [CMA-H]− were observed for the anions produced from electrospray in EtOH at m/z 121, 117, and 93 (Figure S7) and from electrospray in MeCN at m/z 145, 117, 93, and 91 (Figure S10). The minor photofragments can only be clearly observed through the band I region, likely due to electron detachment competing more effectively against ionic fragmentation as excitation energy increases. The VDE of the carboxylate and phenoxide deprotoners of CMA are calculated to be 3.73 and 2.42 eV (Table 1), respectively, so any electronic excitations lying above this energy occur within the electron detachment continuum. The propensity for electron detachment versus photofragmentation is discussed further in Section 3.3.

3.2.2. [CA-H]−. Figure 2a displays the UV–vis laser PD spectrum of [CA-H]− electrosprayed from EtOH and MeCN. Five broad absorption bands (I–V) are observed, with an initial strong feature (I) between 2.6 and 3.4 eV, followed by a cluster of somewhat less intense bands (II–V) from the UVB/UVC regions that peak at ca. 4.0, 4.4, 4.8, and 5.2 eV, respectively.

The theoretical gas-phase absorption spectrum of [CA-H]− is displayed in Figure 2b (black line) and was constructed using a 1:2 carboxylate:phenoxide weighted linear combination of the spectral contributions for the two deprotoners. Band I corresponds to a bright π* ← π transition (A′, 2.79 eV) of the para-phenoxide deprotomer, clearly revealing the presence of the phenoxide deprotomer in the experimental spectrum. The somewhat broader bands (II–V) are more challenging to assign unambiguously, but bands II and III appear to result from the overlap of a number of bright π* ← π transitions originating...
from the carboxylate and para- and meta-phenoxide deprotoners, while band IV appears to be dominated by excitations of the carboxylate deprotomer ($A', 4.81$ eV). (We used a ratio of 5:1 for the para:meta conformers to obtain the best fit to the experimental spectrum. It should be noted that these conformers can interconvert via proton transfer, so likely equilibrate with the more stable para-conformer then dominating.) Comparison of the experimental spectrum obtained by electrospraying CA from both EtOH and MeCN with the theoretical summed spectrum clearly shows that the experimental spectrum must result from both isomers being present in the electrosprayed ion ensemble in the ratio of $\sim 1:2$ carboxylate:phenoxide.

Photoexcitation of [CA-H]$^-$ is associated with two significant ionic photofragmentation channels (eqs 4a and 4b) that correspond to the production of the $m/z$ 135 and 134 photofragments (Figures 2b and 2c; Table 2) associated with loss of neutral CO$_2$ and H + CO$_2$, respectively. Both of these photofragments are produced predominantly through the band I region, corresponding to photoexcitation of the lowest-lying singlet excited state of the para-phenoxide deprotomer. Relatively minor photofragments of [CA-H]$^-$ can be observed at $m/z$ 133, 117, 109, 107, and 91 for electrospray from EtOH and at $m/z$ 133, 117, 109, and 107 for electrospray from MeCN (Figures S8 and S11).

$$[\text{CA-H}^-] + h\nu \rightarrow m/z 135 + CO_2 \quad (4a)$$
$$\rightarrow m/z 134 + CO_2 + *H \quad (4b)$$
$$\rightarrow e^- + [\text{CA-H}]^- \quad (4c)$$

The propensity for electron detachment versus photofragmentation is discussed further in Section 3.3.

3.2.3. [FA-H]$^-$ Figure 3a displays the gas-phase UV–vis PD spectra of [FA-H]$^-$. In contrast to that of [CMA-H]$^-$ and [CA-H]$^-$, the PD spectrum of [FA-H]$^-$ changes considerably depending on whether FA is electrosprayed in EtOH or MeCN. Using EtOH as the solvent (blue spectrum), we observe three absorption features (labeled I–III) with an absorption onset at the lowest energy band (I) of 2.6 eV. This band relatively weak and broad and is observed to extend from the visible region into the UV (2.6–3.4 eV). A similarly broad band (II) is observed between 3.4 and 4.25 eV, with a maximum at ca. 4.0 eV. Band III appears as a rather strong feature with an absorption extending between 4.3 and 5.5 eV and peaking prominently at 4.8 eV. In contrast when FA is electrosprayed from MeCN (red spectrum), band I dominates the PD spectrum of [FA-H]$^-$, while bands II and III are significantly less intense.

In [CMA-H]$^-$ and [CA-H]$^-$, band I was unambiguously assigned to a bright $\pi^* \rightarrow \pi$ transition originating from a phenoxide deprotomer, with the higher-energy bands being assigned to $\pi^* \rightarrow \pi$ transitions originating from carboxylate deprotoners. Our calculations predict the same assignments for [FA-H]$^-$; band I arises from the bright $\pi^* \rightarrow \pi$ transition of the $C_1$-symmetric phenoxide deprotomer ($A', 2.79$ eV), while bands II and III are associated with transitions of the carboxylate deprotoners. The greater changes observed for the PD spectra upon electrospraying from EtOH and MeCN for FA reveal that production of deprotoners for this molecule is a more sensitive function of solvent than for CMA and CA. (The distribution of deprotoners will be a function of differences in the relative stability of the deprotonated form for this molecule is a more sensitive function of solvent than for CMA and CA. The distribution of deprotoners can be observed at $m/z$ 133, 117, 109, 107, and 91 for electrospray from EtOH and at $m/z$ 133, 117, 109, and 107 for electrospray from MeCN (Figures S8 and S11).

$$[\text{FA-H}^-] + h\nu \rightarrow m/z 178 + [\text{CH}_3]^* \quad (5a)$$
$$\rightarrow m/z 149 + CO_2 \quad (5b)$$
$$\rightarrow m/z 134 + [\text{CH}_3]^* + CO_2 \quad (5c)$$
$$\rightarrow e^- + [\text{FA-H}]^- \quad (5d)$$

We note that pathways 5a and 5c both result in the production of free radical species. Similar free radical products have been observed for the deprotonated forms of the sunscreens oxybenzone, 2-phenylbenzimidazole-S-sulfonic acid, and benzophenone-4 in recent work.36–38
Figures 4b–4e present the action spectra for the four major ionic photofragments of [FA-H]− electrosprayed from EtOH (blue spectrum). The precursor [FA-H]− (m/z 193) PD spectrum is reproduced in Figure 4a to facilitate comparison. It is evident from Figure 4 that the relative production yield of the four major photofragments of [FA-H]− at m/z 178, 149, 134, and 117 varies significantly as a function of photon energy. The production spectra of m/z 178 and 149 (Figures 4b and 4c), respectively, show that both photofragments are primarily produced across the 2.6–3.6 eV (visible-UVA) region, with m/z 178 produced only in low yield around 4.0 and 4.6 eV. The action spectrum of m/z 134 (Figure 4d), however, shows that the fragment is produced across the entire spectral range with production peaking close to the band maxima I−III (Figure 4a). We note that production of m/z 134 is enhanced through band II. The production of the photofragment at m/z 117 (Figure 4e) is almost 10-fold less intense than the m/z 149 and 134 ions and is produced only through bands II and III.

The photofragment production spectra obtained from [FA-H]− when it is electrosprayed from MeCN are presented in red on Figures 4b–4e. Electrospraying from MeCN dramatically reduced ionic fragment production at ca. 3.5 eV, with the m/z 134 photofragment ion (the major photofragment when FA is electrosprayed from MeCN) almost entirely absent.

Figure S12 presents the relative ion yield spectra of the photofragments of [FA-H]− electrosprayed from both EtOH and MeCN, providing an overview of their relative production. These plots show that production of m/z 149, which corresponds to the loss of CO2 from [FA-H]−, is strongly enhanced through the band I region, which is associated with the phenoxide deprotoner. The m/z 134 photofragment (eq 5c) shows a generally increasing trend in production with photon energy, although the resulting production profile does vary for electrospray from the two solvents. Production of m/z 178 can...
be seen to decrease as production of $m/z$ 134 increases, suggesting that the $m/z$ 134 ion may be a secondary photofragment produced when $m/z$ 178 has high internal energy.

The propensity for electron detachment versus photofragmentation is discussed further in the next Section 3.3.

### 3.3. Electron Detachment Yield vs Photodepletion Yield Interpretation

The electron detachment yield spectra of $[\text{CMA-H}]^-$, $[\text{CA-H}]^-$, and $[\text{FA-H}]^-$ are displayed in Figures 5a–5c and 6, with the corresponding photodepletion spectra for comparison. Electron loss is not directly measurable in our instrument, so these spectra are obtained by assuming that any photodepleted ions not detected as ionic photofragments are lost through electron detachment. (This is true for the situation where ionic fragments with $m/z < 50$ represent only minor decay pathways.) Further discussion of electron detachment yield spectra can be found in ref. 4242.

Figures 5a–5c present the electron detachment yield spectra obtained when electrospraying in ethanol, while Figures 6a–6c display spectra obtained when the solvent is acetonitrile. For both sets of spectra, the electron detachment yield curves display very similar profiles to the corresponding photodepletion spectra, indicating that electron detachment is the main photodepletion pathway. Ionic fragmentation can be seen to be relatively more significant through the band I regions (2.5–3.25 eV), with electron detachment becoming increasingly more dominant as the photoexcitation energy increases. We note that the calculated VDEs for the phenoxide and carboxylate isomers of the $[\text{CMA-H}]^-$, $[\text{CA-H}]^-$, and $[\text{FA-H}]^-$ anions are predicted to be around 2.4 and 3.7 eV, respectively (Table 1). The entire photodepletion region scanned here therefore lies at or above the detachment energies for the phenoxide deprotoners, with the band II and II regions lying above the VDEs of the carboxylate isomers. Despite the fact that the photodepletion spectra are being acquired above the detachment energies, the

![Figure 4](https://doi.org/10.1021/acs.jpca.1c04880)

![Figure 5](https://doi.org/10.1021/acs.jpca.2100000)

**Figure 4.** (a) Gas-phase photodepletion spectrum of $[\text{FA-H}]^-$ electrosprayed in EtOH (blue) and MeCN (red). (b–e) Photofragment production spectra of the four major photofragments of $[\text{FA-H}]^-$ with $m/z$ 178, 149, 134, and 117, respectively, as electrosprayed in EtOH (blue) and MeCN (red). The solid lines are a five-point adjacent average of the data points.

**Figure 5.** Electron detachment yield (ED$^*$; red) vs photodepletion yield (PD$^*$; blue) spectra of (a) $[\text{CMA-H}]^-$, (b) $[\text{CA-H}]^-$, and (c) $[\text{FA-H}]^-$ when electrosprayed in EtOH, respectively. The solid lines are a five-point adjacent average of the data points. The overlaid arrows represent the calculated VDEs of the phenoxide (solid gray) and carboxylate (dotted gray) deprotomers, as outlined in Table 1.

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**Table 1.**

| Ion          | Phenoxide VDE (eV) | Carboxylate VDE (eV) |
|--------------|--------------------|----------------------|
| $[\text{CMA-H}]^-$ | 2.4                | 3.7                  |
| $[\text{CA-H}]^-$   |                    |                      |
| $[\text{FA-H}]^-$   |                    |                      |

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**Section 3.3.**

Electron Detachment Yield vs Photodepletion Yield Interpretation. The electron detachment yield spectra of $[\text{CMA-H}]^-$, $[\text{CA-H}]^-$, and $[\text{FA-H}]^-$ are displayed in Figures 5a–5c and 6, with the corresponding photodepletion spectra for comparison. Electron loss is not directly measurable in our instrument, so these spectra are obtained by assuming that any photodepleted ions not detected as ionic photofragments are lost through electron detachment. (This is true for the situation where ionic fragments with $m/z < 50$ represent only minor decay pathways.) Further discussion of electron detachment yield spectra can be found in ref. 4242.

Figures 5a–5c present the electron detachment yield spectra obtained when electrospraying in ethanol, while Figures 6a–6c display spectra obtained when the solvent is acetonitrile. For both sets of spectra, the electron detachment yield curves display very similar profiles to the corresponding photodepletion spectra, indicating that electron detachment is the main photodepletion pathway. Ionic fragmentation can be seen to be relatively more significant through the band I regions (2.5–3.25 eV), with electron detachment becoming increasingly more dominant as the photoexcitation energy increases. We note that the calculated VDEs for the phenoxide and carboxylate isomers of the $[\text{CMA-H}]^-$, $[\text{CA-H}]^-$, and $[\text{FA-H}]^-$ anions are predicted to be around 2.4 and 3.7 eV, respectively (Table 1). The entire photodepletion region scanned here therefore lies at or above the detachment energies for the phenoxide deprotoners, with the band II and II regions lying above the VDEs of the carboxylate isomers. Despite the fact that the photodepletion spectra are being acquired above the detachment energies, the
3.4. Higher-Energy Collisional Dissociation vs Photo-fragmentation. To investigate the thermal fragmentation pathways of the deprotonated antioxidants on their electronic ground states, higher-energy collisional dissociation (HCD) was employed (Figures 7a−7c and 8a−8c; Table 2). These measurements are essential to identify ions that are secondary fragments, i.e., ionic fragments that are formed when a precursor species breaks apart at high internal energy (ref 43 gives a full discussion of how the HCD energies can be related to ion internal energies obtained upon photoexcitation.) In addition, these measurements are important as any photofragments not observed in HCD studies can be identified as purely photochemical products.

Collisional activation of [CMA-H]− produced from both EtOH and MeCN (Figures 7a and 8a, respectively), produces a single major HCD product with m/z 119. (A m/z 93 ion is produced as an extremely minor fragment at the very highest collisional energies measured.) The m/z 119 ion is the only significant photofragment observed for [CMA-H]−.

For [CA-H]− electrosprayed from both EtOH and MeCN (Figures 7b and 8b), HCD produces the m/z 135 ion (loss of CO2) as the dominant fragment. The ion at m/z 134 is observed as a very low intensity fragment >30% HCD energy. m/z 135 and 134 are the major and minor photofragments of [CA-H]−, and our HCD results suggest that m/z 134 photofragment is a secondary photofragment produced at high internal energies of m/z 135 (since m/z 134 increases over the same energy range that m/z 135 decreases). For both [CMA-H]− and [CA-H]−, the similar profile of HCD fragment production when the deprotonated molecule is electrosprayed from either solvent is used. Collisional activation of [FA-H]− from both EtOH and MeCN (Figures 7c and 8c) results in the m/z 134 fragment (loss of [CH3]• + CO2) as the dominant product, with m/z 178 (loss of [CH3]•) and 149 (loss of CO2) appearing as medium intensity fragments. The intensity of both m/z 178 and 134 decreases at higher collisional energies, indicating that m/z 178 is fragmenting into m/z 134 at higher internal energy. (m/z 117 and 89 are observed as extremely minor fragments >35% HCD energy.) Notably, there is a significant difference in the relative ion production intensities of the m/z 178 and 134 fragments from [FA-H]− for electrospray from the two solvents. When EtOH is employed as the electrospray solvent, m/z 134...
dominates as the most intense ion across the HCD energies scanned (e.g., at 17.4% HCD energy: m/z 134 = 34% and m/z 178 = 23%). In contrast, when MeCN is used as the electrospray solvent, m/z 178 is the main ion observed between 0 and 40% HCD energy (e.g., at 17.4% HCD energy: m/z 134 = 17% and m/z 178 = 34%). This is consistent with the proportion of the two possible deprotomers varying when different electrospray solvents are employed.

Finally, we note that the major HCD fragments mirror the major gas-phase photofragments, which suggests that these deprotonated coumaric acids are relaxing by predominantly statistical (ergodic) decay over the spectral range (2.5−5.5 eV) studied.

4. CONCLUSIONS

Recent time-resolved laser spectroscopic and theoretical studies of the CMA, CA, and FA have explored their photodynamics, revealing that the molecules can undergo rapid nonradiative excited state decay via a cis−trans isomerization process. A limited number of studies have explored how these photodynamics are affected by pH and, hence, the extent of deprotonation, finding that deprotonation had little effect on excited state decay. This result is consistent with our results here for the deprotonated coumaric acids, since the ionic fragmentation patterns for UV−vis laser and HCD excitation are consistent with statistical fragmentation which is associated with rapid ultrafast decay. This is perhaps not surprising given that the coumaric acid deprotonation sites are not the key parts of the molecule that effect the nonradiative transition from the excited state back to the electronic ground state. However, the results presented here provide confirmation that the action of coumaric acid based antioxidants are likely to perform well as UV filters even in mildly alkaline environments such as seawater.

Two recent studies have used a custom-built ion mobility mass spectrometer to study deprotonated coumaric acids formed via electrospray. The main focus of these studies was on the photoisomerisation dynamics and excited state proton transfer. Carboxylate and phenoxide deprotonomers were identified in these studies through their differential collisional cross sections. Ion mobility separation of deprotoners will be advantageous for systems where the optical transitions associated with a pair of deprotoners are not well separated. While ion mobility detection is clearly complementary to the UV−vis laser photodissociation spectroscopy within the commercial mass spectrometer presented here, it is important to have a range of approaches available since what is crucial is that they can be applied in situ to identify deprotomerics species. This is of particular importance for the reaction studies discussed in the introduction, where the type of mass spectrometer available for ESI detection may vary.

The results presented above show that laser photodissociation spectroscopy within a laser-interfaced mass spectrometer combined with theoretical wave function calculations can identify the phenoxide and carboxylate deprotoners of coumaric acids produced by ESI. For deprotonated CA and CMA, we find that the ratio of phenoxide:carboxylate deprotoners is relatively insensitive to electrospray in both protic and aprotic solvents. The ratio of deprotoners of FA is considerably more strongly affected by the electrospray solvent employed, with electrospray from aprotic MeCN leading to a higher proportion of the carboxylate deprotoner, which is favored in solution. Overall, our results show unequivocally that a mixture of carboxylate:phenoxide deprotoners is formed upon electrospray for all three coumaric acids studied here. This leads us to conclude that this is likely to be a general phenomenon when deprotonated coumaric acids are produced via electrospray.

While a number of studies have now used UV−vis laser photodissociation to identify protomers, to our knowledge our study represents the first where photodissociation supported by advanced quantum chemical calculations has been used to definitively identify deprotomerics ions. Given that there are a growing number of fundamental studies being conducted on gaseous ions produced via electrospray, as well as electrospray mass spectrometry being used as an analytic tool to monitor solution-phase reactions, it is important to have tools available to spectroscopically identify deprotomerics (or protomerics) structures. Although in situ IR spectroscopy has been used for this purpose, free electron lasers have typically been employed in these experiments, which leads to challenges due to access limitations. Our results demonstrate an accessible alternative in situ approach using adapted commercial

Figure 8. Parent ion dissociation curves for solutions of (a) [CMA-H]− (m/z 163), (b) [CA-H]− (m/z 179), and (c) [FA-H]− (m/z 193) electrosprayed in MeCN, respectively, for their most intense ionic fragments formed upon HCD between 0 and 40% energies. The curved lines are a three-point adjacent average of such data points and are provided as a viewing guide, to emphasize the profile for each individual fragment.
instrumentation that can successfully determine deprotonation ratios in electrosprayed ions with UV chromophores.

**ASSOCIATED CONTENT**

1 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jPCA.1c04880.

Deprotomer structures of [CMA-H]−, [CA-H]−, and [FA-H]−, optimized Cartesian coordinate tables for [CMA-H]−, [CA-H]−, and [FA-H]−; photodepletion laser power dependence measurements; additional photofragment action spectra; an ion yield plot for [FA-H]−; proposed structures of the major ionic fragments; excited state assignments obtained from the calculations; key molecular orbitals (PDF).

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Notes
The authors declare no competing financial interest.

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