Structures and Chemical Rearrangements of Benzoate Derivatives Following Gas Phase Decarboxylation

Evan H. Perez,^a Tim Schleif,^a Joseph P. Messinger,^a Anna G. Rullán Buxó,^a Olivia C. Moss,^a Kim Greis,^{a,b} Mark A. Johnson^{a,*}

^a Sterling Chemistry Laboratory, Department of Chemistry, Yale University, New Haven, CT, USA 06520
 ^b Institut für Chemie und Biochemie, Freie Universität Berlin, Arnimallee 22, 14195 Berlin (Germany); Fritz-Haber-Institut der Max-Planck-Gesellschaft, Faradayweg 4–6, 14195 Berlin (Germany)
 E-Mail: mark.johnson@yale.edu

Abstract:

Decarboxylation of carboxylate ions in the gas phase provides a useful window into the chemistry displayed by these reactive carbanion intermediates. Here we explore the species generated by decarboxylation of two benzoate derivatives: 2-formylbenzoate (2FBA) and 2-benzoylbenzoate (2BBA). The nascent product anions are transferred to a cryogenic ion trap where they are cooled to ~15 K and analyzed by their pattern of vibrational bands obtained with IR photodissociation spectroscopy of weakly bound H₂ molecules. The structures of the quenched species are then determined by comparison of these spectra with those predicted by electronic structure calculations for local minima. The 2-phenide carbanion generated by decarboxylation of 2FBA occurs in two isomeric forms that differ in the orientation of the formyl group, both of which yield a very large (~110 cm⁻¹) red-shift in the H₂ molecule bound to the anionic carbon center. Although calculated to be a local minimum, the analogous 2-phenide species could not be isolated upon decarboxylation of 2BBA. Rather, the anionic product adopts a ring-closed structure, indicating efficient nucleophilic attack on the pendant phenyl group by the nascent phenide. The barrier for ring closing is evaluated with electronic structure calculations.

I. Introduction

The ion chemistry following decarboxylation of deprotonated organic acids is mediated by highly reactive carbanions that undergo a variety of reactions when carried out in the condensed phase.¹⁻² The gas phase analogues, in which carboxylates undergo decarboxylation by collision-induced dissociation (CID), provide a useful way to isolate the carbanions and determine their intrinsic thermochemistry, structure and reactivity patterns. Early studies include mass spectrometric efforts by



Scheme 1. Decarboxylation of acids via collisioninduced dissociation.

Squires and Graul³⁻⁴ and negative ion photoelectron spectroscopic measurements by the Lineberger group in the 1990s⁶ More recent efforts involve anion vibrational spectroscopy in conjunction with electronic structure calculations to deduce rearrangements. For example, Oomens and co-workers⁷⁻⁸ used IR multiphoton dissociation of the 300-350 K CID product ions to investigate reactive pathways following decarboxylation. In this paper, we address the nature of the decarboxylation product anion formed upon CID of the 2-formyl- (2FBA, **1**) or 2-benzoylbenzoate (2BBA, **3**) depicted in Scheme 1. Both species are anticipated to generate the phenide motif after heterolytic C–C bond cleavage. The focus of this study is to determine the extent of the intramolecular interactions between the reactive carbanion center and proximal functional groups (H and phenyl, respectively) available in this benzoate scaffold. The nascent product anions are cooled to ~15 K in a cryogenic ion trap, and the structures of the cold reactant and product anions are deduced by comparing their vibrational spectra with calculated patterns for candidate structures.

II. Experimental details and computational methods.

2-benzoylbenzoic acid, 2-formylbenzoic acid, acetonitrile (HPLC grade), and water were purchased from Sigma-Aldrich and used without further purification. Helium and hydrogen were purchased from Air Liquide. The deprotonated anions were generated via electrospray ionization using 1 mM solutions of the respective acids in 1:1 acetonitrile/water. Mass spectra and photofragmentation infrared spectra were collected using a hybrid instrument that combines a commercial ThermoFisher Scientific LTQ Orbitrap Velos with a custom-built, triple-focusing time-of-flight photofragmentation mass spectrometer that has been previously described.⁹ In brief, high-resolution mass spectra were collected using the instrument's Orbitrap capability at maximum resolution ($\Delta m/m = 100,000$), though a reduced resolution of $\Delta m/m = 7,500$ was used when transferring ions to the photofragmentation spectrometer to maximize ion transmission. Collision-induced dissociation experiments were conducted using the LTQ functionality of the Velos Orbitrap and utilized a Normalized Collision Energy (NCE) of 30 au. All benzoate anions readily undergo decarboxylation in collision-induced dissociation and loss of 44 u is indeed the dominant fragment in the mass spectra as illustrated in Figs. S1 and S2. Weakly bound complexes of the analyte ions and H_2 were formed in a cryogenically cooled ion trap held at ~15 K. Vibrational spectra of the ions were obtained after photodissociation of the H₂-tagged species.

DFT was used to calculate the normal modes of the target ions at the CAM-B3LYP/6-311++G(2d,2p) level of theory. All calculations were conducted using Gaussian 09.10 Candidate structures were chosen empirically based on the match between the computed frequencies and experimentally observed absorption bands. All energies mentioned throughout the text are evaluated as the sums of the respective electronic energy plus zeropoint vibrational energies. Harmonic vibrational frequencies were scaled by 0.953 based on literature scaling factors for the employed level of theory;¹¹ the H₂ tag vibrations, given their importance for our later analyses, were treated separately by a scaling factor of 0.96 that was determined empirically from matching the calculated H₂ stretching frequency in 2FBA·H₂ to its experimental counterparts.

III. Results and discussion

IIIA. Vibrational spectra of 2FBA and 2FBA-CO₂: Structural deformations and H_2 tag shifts

To establish the spectral features associated with the parent carboxylate and phenide motifs, we first present the IR photodissociation spectra of $2FBA \cdot H_2$ (1 $\cdot H_2$) and its decarboxylate, 2FBA-CO₂·H₂ (2·H₂), in Fig. 1a and 1c, respectively. In each case, the bands highest in energy (v^{H_2} at 4078 and 3966 cm⁻¹, respectively) originate from the H₂ stretch of the tag. The former value is typical for H_2 attached to the $-CO_2$ head group. In the case of carboxylates like acetate and the anion and dianion of dodecanoic acid H₂ is bound to one of the oxygen atoms and points toward it along the H₂ bond axis with a binding energy of ~500 cm⁻¹ and frequency of ~4050 cm⁻¹.¹² The much lower frequency of the H₂



Fig. 1. Cryogenically cooled, H₂-tagged vibrational predissociation spectra of (a) 2FBA (1), (c) the decarboxylate of 2FBA (2) which was generated by collision-induced dissociation of the parent carboxylates. (b) and (d,e) depict the calculated IR spectra of 2FBA (1), and the decarboxylate (2). The CO stretch fundamentals of the carbonyl groups are highlighted in blue and the aldehyde CH stretching fundamental is highlighted in pink. Assignments indicated by band labels are detailed in Table 1. Features marked with an asterisk are not reproduced by harmonic frequency calculations (see text). All calculations were performed at the CAM-B3LYP/6-311++G(2d,2p) level of theory.

stretch in 2FBA-CO₂·H₂ (**2**·H₂), however, indicates that the H₂ molecule is more strongly bound to the carbanion. This qualitative difference is captured in the calculated structures of the two species, which are indicated in the insets in Fig. 1a and 1d. The H₂ molecule in 2FBA-CO₂·H₂ (**2**·H₂) is calculated to be slightly elongated (by 0.007 Å) relative to the value calculated for 2FBA·H₂ (**1**·H₂), accounting for the 112 cm⁻¹ red-shift in the H₂ stretch. The H₂ molecule in the 2FBA-CO₂·H₂ (**2**·H₂) complex is calculated to adopt an off-axis docking geometry (by 33 degrees relative to the C²-C⁵ bisector) with an additional tip angle of the H₂ axis (by 18 degrees as indicated in Fig. 2d). We note that the H₂ stretching region in the 2FBA-CO₂·H₂ (2·H₂) complex appears with smaller bands (labeled *) higher in energy. These are traced to combination bands with soft modes based on the results of two-color, IR-IR isomer-selective spectra (displayed in Fig. S3). The physical origin of the large H₂ red shift is discussed further in section IIIC.

Returning to 2FBA·H₂ (**2**·H₂), lower in energy, strong bands at 1329 cm⁻¹ and 1646 cm⁻¹ occur in typical positions for the symmetric and asymmetric stretching modes ($v_{sym}^{CO_2}$ and $v_{asym}^{CO_2}$, respectively) of the CO₂⁻ group. The band to the blue found at 1676 cm⁻¹ is attributed to the C=O ($v^{C=0}$) stretching mode. It should be noted that the calculated structure for 2FBA·H₂ (**2**·H₂) provided in Fig 1b was the only low-lying energetic minimum found for this ion. Here the aldehyde oxygen is pointed away from the carboxylate head group. The lowest frequency vibrational mode in the CH stretching region can be attributed to the aldehyde CH stretching fundamental ($v_{HC=0}^{C-H}$) at 2904 cm⁻¹. Typical CH stretching fundamentals for benzaldehydes appear at 2700–2900 cm⁻¹.¹³

Fig. 1c presents the vibrational spectrum of $2FBA-CO_2 \cdot H_2$ ($2 \cdot H_2$). Following decarboxylation, the carboxylate stretching modes, as expected, are absent. The $2FBA-CO_2 \cdot H_2$ ($2 \cdot H_2$) spectrum displays a close doublet around 1650 cm⁻¹. However, the harmonic calculation for this species only predicts a single feature. This suggests either anharmonic coupling due to higher-order terms in the potential or participation of a second isomer. Calculations indicate that the rotamer **2b** arising from the orientation of the formyl group relative to the carbanion center lies 7.2 kcal mol⁻¹ above the minimum energy isomer **2a**. These two isomers yield similar bands in the C=O stretching region, but the aldehyde CH modes are calculated to be significantly



Fig. 2. Cryogenically cooled, H₂-tagged vibrational predissociation spectra of (a) 2BBA (**3**), (b) the decarboxylate of 2BBA (**4**), and (c) the decarboxylate of 2FBA (**2**), the latter two of which are generated by collision-induced dissociation of the parent carboxylates. The CO stretch fundamentals of the carbonyl groups are highlighted in blue. The redshifted CH stretch of the decarboxylate of **3** (marked with an asterisk) is highlighted in red and the aldehyde CH stretches of **2** are highlighted in pink, which are significantly redshifted from the position in 2FBA·H₂ (**1**·H₂), at 2903 cm⁻¹ (Fig. 1). The arrow at 4161 cm⁻¹ indicates the H₂ stretch frequency of free H₂.⁵ Assignments indicated by band labels as well those denoted by letters a_i and b_i are detailed in Table 1.

different, with that in **2b** lying 96 cm⁻¹ below the band in **2a**. In fact, two features (at 2672 and 2764 cm⁻¹⁾ are observed in the region where aldehyde CH stretching vibrations would be expected. These are highlighted with a red asterisk in Fig. 1c, and are consistent with the two-isomer interpretation of the C=O stretch doublet. Although the two rotamers **2a** and **2b** are separated by an activation barrier of 11.8 kcal mol⁻¹, it has been shown that CID can provide enough internal energy¹⁴⁻¹⁶ to fragment ions to accommodate such a rearrangement. Subsequently, the metastable isomer can be trapped behind a significant barrier upon quenching in the cold trap. The higher-energy CH stretching region is more populated,

appearing as a dense manifold confined to the region 2975-3100 cm⁻¹ that occur in the usual range for aromatic CH groups and are observed in both experimental spectra (Fig. 1a and 1c). One of these bands falls in the location of the combination band arising from excitation of one quantum in each of the CO stretching modes arising from carboxylate. This band is denoted $v_{v1+v3}^{CO_2}$ in Fig. 1a. Although IR forbidden at the harmonic level, this transition is commonly observed in the spectra of decarboxylated anions.¹⁷

IIIB. Vibrational spectra of 2BBA and 2BBA-CO₂: Structural rearrangement of the carbanion

After assigning the spectral signatures of an ortho-carboxylate (1) and a phenide (2) produced by decarboxylation of 2FBA, we next turn to the 2BBA (3) system and its decarboxylate. This system was chosen because it provides a richer chemical landscape for intramolecular interactions and opens the possibility of chemical rearrangement. Two views of the calculated structure of 3 are presented as insets in Fig. 2a. In contrast to the 2FBA structure, the carboxylate group in **3** resides in the plane of the ring (ϕ = 4°), similar to the arrangement in the isolated benzoate anion.¹⁸ The angle between the two aromatic rings is calculated to be 80°, consistent with greater steric hindrance afforded by the phenyl group relative to that from the formyl CH. The carbonyl group is predicted to be displaced off ring axis by



Fig. 3. a) Potential energy diagram for the chemical tranformation of the phenide ion (**4**) to the fluorenone-like ring closed product (**5**). b) Experimental IR spectrum of the H₂-tagged 2BBA decarboxylate at 15 K, compared to the (scaled) computed vibrational spectrum predicted for the ring closed structure **5** (c). The CO stretch, coupled to various ring stretches, is highlighted in blue and the sp³ CH stretch is highlighted in red. Band assignments are located in Table 1. All calculations were performed at the CAM-B3LYP/6-311++G(2d,2p) level of theory.

13°, suggesting intramolecular repulsion from the proximal carboxylate charge center.

Figure 2a presents the vibrational spectrum of the 2BBA·H₂. Starting highest in energy, the H₂ stretch can be found at nearly the same position as observed for 2FBA·H₂ (8 cm⁻¹ difference). The symmetric and asymmetric carboxylate stretches are also very close to those found in 2FBA, as indicated by comparison of the experimental band positions collected in Table 1. The CH stretching region is less congested, revealing the combination band arising from the CO₂⁻ symmetric and asymmetric stretches, discussed above, as an isolated feature ($v_{v1+v3}^{CO_2}$, Fig. 2a). The carbonyl fundamental again lies just above the CO₂⁻ asymmetric stretch (by 28 cm⁻¹). The significant spectral features of these benzoates are thus preserved in the two benzoate derivatives and are nicely reproduced by the (scaled) calculated frequencies (as are the ones for the respective decarboxylates) with an average deviation of $\Delta v < 20$ cm⁻¹ (Table 1).

Table 1. Experimental (\pm 4 cm⁻¹) bands (bold), harmonic frequencies (in parentheses, scaled by 0.953), and mode character assignments/labels for the computed spectra. All calculations were performed at the CAM-B3LYP/6-311G++(2d,2p) level of theory.

Label/Assignment	0 H ⊖ H₂ 2FBA·H₂ (1)	рн о 	о Н 9 2FBA-CO ₂ ·H ₂ (2а)	H ₂
v^{H_2}	4069	4077	3964	4066
[a ₁ / b ₁]	(4044.2)	(4054.7)	(3905.8)	(4064.3)
$m{ u}_{ring}^{C-H}$ [a ₂ / b ₂]	2959, 2986, 3015, 3059, 3084 (3020.7, 3039.2, 3059.1, 3069.2)	2971-3094 (3008.0- 3088.5)	2917, 2968, 3010, 3062 (2919.5, 2978.0, 3010.2, 3029.5)	2996-3049 (2993.1-3049.7)
$ u_{HC=0}^{C-H} $ [b ₃]	2903 (2936.7)	N/A (N/A)	2672, 2764 (2803.1)	N/A (N/A)
ν ^{C=0} [a₄ / b₄]	1675 (1678.7)	1663 (1665.3)	1637, 1659 (1650.0)	1560 (1553.0)
$oldsymbol{ u}_{sp^3}^{\mathcal{C}-\mathcal{H}}$ [a3]	N/A (N/A)	N/A (N/A)	N/A (N/A)	2667 (2709.4)
$v_{ring}^{sym C=C}$	1584 (1588.2)	1559, 1584 (1565.2 <i>,</i> 1588.5)	1546 (1549.4)	1581, 1597 (1586.3, 1598.3)
ν ^{C−H} ring bend [a₅∫b₅]	1317 (1316.3)	1271 (1219.6, 1247.3, 1270.2)	1129, 1173, 1251, 1397, 1543	1439, 1464 (1440.0, 1452.7)

	(1112.5, 1164.5, 1235.3, 1393.9, 1549.4)				
v ^{asym CO} 2	1635	1635	N/A	N/A	
	(1617.7)	(1620.3)	(N/A)	(N/A)	
v ^{sym} CO ₂	1317	1326	N/A	N/A	
	(1316.3)	(1330.8)	(N/A)	(N/A)	
V ^{C-H,} aldehyde	1364	N/A	1336	N/A	
V _{bends}	(1353.8)	(N/A)	(1323.0)	(N/A)	
$v^{C-H}_{sp^3 bend}$ [a ₆]	N/A	N/A	N/A	1259, 1277	
	(N/A)	(N/A)	(N/A)	(1248.8)	

The 2BBA-CO₂·H₂ vibrational spectrum is displayed in Fig. 3b. Most notably, in contrast to the situation found in 2FBA-CO₂·H₂, the H₂ stretching frequency is essentially unperturbed from its position in the 2BBA \cdot H₂ parent spectrum (Table 1). This indicates that the carbanion scaffold is not based on the phenide functionality adopted by 2FBA-CO₂. The characteristic C=O fundamental is also missing in the 2BBA-CO₂·H₂ spectrum, and is replaced by a strong triplet, denoted a₄, centered 150 cm⁻¹ lower in energy. Moreover, a new band at 2664 cm⁻¹ (a₃) appears far below the aryl C-H stretching modes of 2BBA. The presence of the starkly redshifted $C(sp^3)$ -H stretching vibration is indicative of a relatively weakly bound hydride such as the one found in HCO₂⁻ ($v^{CH} = 2449 \text{ cm}^{-1}$).¹⁹ A strongly redshifted C–H stretch also occurs in the σ -complex resulting from protonation of benzene ($v^{\widetilde{CH}} \sim 2800 \text{ cm}^{-1}$), a binding motif that creates a local sp³ motif.²⁰⁻²¹ Meanwhile, redshifts in carbonyl C=O stretch are indicative of electron density being displaced into the antibonding π^* orbital, resulting in bond elongation. Taken altogether, these spectral changes indicate that a significant structural rearrangement has occurred in the carbanion generated by decarboxylation. Such skeletal rearrangements of closed-shell anions in gas-phase have been extensively investigated by Bowie et al. as summarized in their review,²² with the one reported here being reminiscent of the formation of fluorenylanions from Ph₂CH⁻ proposed by them.²³

An ion containing the phenide motif 4 was optimized as a local minimum, with the structure and harmonic spectrum presented in Fig. S4. This geometry is predicted to yield similar spectral features as those found in 2FBA-CO₂, e.g., with a typical carbonyl fundamental frequency and a strongly red shifted H₂ stretch. This pattern is not consistent with the observed 2BBA-CO₂ spectrum (Fig. 2c). However, an isomer **5** significantly lower in energy was found with the structure indicated in the inset of Fig. 3a. This isomer is connected to phenide 4 by an activation barrier of 11.2 kcal mol⁻¹ and stems from the nucleophilic attack of the negative charge center on the pendant phenyl ring. Similar rearrangements of the benzophenone scaffold have previously been observed in the gas phase for positively charged ions and in the condensed phase.24-25 The calculated spectrum for 5 (Fig. 3c) indeed reproduces the experimentally observed band pattern of the 2BBA-CO₂ product ion. Specifically, peak a₃, the redshifted feature at 2664 cm⁻¹, is traced to the C–H stretch of the sp³ hybridized carbon resulting from the ringclosing reaction. The triplet motif of the redshifted C=O band is accounted for by mixing with proximal C=C stretching modes. Note that **5** is the only species formed upon



Figure 4. Plots of the electrostatic potential on the respective $0.02 a_0^{-3}$ isosurfaces of the more stable rotamer of 2FBA-CO₂ (**2a**), the fluorenone-like ring-closed product (**5**) and its phenide isomer (**4**, not observed experimentally). The color code corresponds to: red, negative potential = -0.2; blue, potential = 0. Black circles indicate the calculated position of the H₂ complexes. All calculations were performed at the CAM-B3LYP/6-311++G(2d,2p) level of theory.

decarboxylation of 2BBA, indicating that the ring-closing reaction is rapid under our experimental conditions.

IIIC. Anion dependence of the H₂ frequency shifts

Having elucidated the structures of the **2a**, **4** and **5** anions, it is useful to consider the cause of the differing redshifts displayed by the H₂ adducts in each. A comparison of the electronic density maps of these species is displayed in Fig. 4. It is evident that the negative charge in **5** resides primarily on the carbonyl group, which is indeed the docking site for the H₂ molecule (see structure in Fig. 3b). This concentration of excess charge is consistent with the redshift in the C=O stretch from 2BBA (**3**) to **5**, which reflects the increase in length of the C=O bond from 1.21 to 1.24 Å. In both phenide and ring-closed anions, H₂ attaches to the position of highest negative charge density. However, the degree of charge localization is calculated to be comparable in **2**, **4** and **5** (Fig. 4), while the H₂ stretch is observed to be strongly redshifted (by ~ 100 cm⁻¹) in the phenide species **2** compared to **5** (s. Table 1). Consequently, the origin of the shift is not primarily due to electrostatics. Rather, it likely reflects a chemical interaction with the

carbanion center in the phenide motif, similar to that which drives nucleophilic attack and C-C bond formation leading to 5. In the case of H_2 , this would lead to C–H bond formation with loss of H⁻. The exit channel of the latter would correspond to the H⁻benzaldehyde ion-molecule complex. This raises an interesting speculation that reaction of H⁻ with benzaldehyde could generate 2 by elimination of H₂. To further explore this point, we calculated the potential surface corresponding to H₂ attacking the carbanion center in 2. This was accomplished by scanning the C-H distance while relaxing all other coordinates. The resulting profiles for H₂ approach to both 2FBA-CO₂ isomers (2a and 2b) are displayed in Fig. 5. These curves reveal a scenario in which, after crossing a barrier of ~ 9 kcal mol⁻¹, the systems adopt a minimum energy structure corresponding to the H⁻-benzaldehyde ionmolecule complex. In this context, the anomalous redshift in the H₂ frequencies displayed by the



Fig. 5. Potential energy curve describing the reactive encounter of H_2 with 2FBA-CO₂ (**2**). After a shallow minimum corresponding to the relatively weakly bound "tagging" regime with a minimum near 2.5 Å, a second minimum is recovered upon C–H bond formation to yield the H⁻ adduct with benzaldehyde. All calculations were performed at the CAM-B3LYP/6-311++G(2d,2p) level of theory.

phenide-H₂ adducts reflects this low-lying reaction pathway, where the H₂ "tag" actually resides in this ion-molecule reaction's entrance channel complex. Along these lines, the tricyclic structure of **5** is interestingly reminiscent of the fluorenone scaffold, which would be generated by loss of hydride from the *sp*³ carbon atom. These considerations suggest that a fruitful future direction for further research would be the characterization of the bimolecular reaction chemistry of carbanions generated by decarboxylation of the ring isomers of benzoate derivatives. Of particular interest, for example, is the nature of the 3- and 4-BBA-CO₂ species, which introduce the possibility of trapping the phenide motif depending on the barrier for circumambulatory migration of the carbanion charge center around the ring.

IV. Summary

Collisional decarboxylation of 2-benzoylbenzoate and 2-formylbenzoate anions provides a platform to characterize chemical and structural rearrangements that occur in the fragment carbanions. The structures of the decarboxylates were determined by cooling them to ~15 K in a buffer gas cooled cryogenic ion trap and recording their vibrational spectra with IR photodissociation of H₂ "tag" molecules. Decarboxylation of 2-benzoylbenzoate is observed to exclusively undergo a ring-closing reaction involving C–C bond formation to the proximal phenyl group. The fact that the metastable phenide cannot be isolated after CID indicates that this nascent species is created with internal energy significantly greater than the calculated barrier of 11.2 kcal mol⁻¹. The resulting tricyclic structure has several distinct spectroscopic features relative to those arising from the charge-localized (phenide) motif adopted by the decarboxylate of 2-formylbenzoate. These include a much smaller redshift of the weakly bound H₂ stretch and a very strongly redshifted CH stretch fundamental that is traced to the CH group located on the *sp*³ carbon center at the apex of two rings. The different spectroscopic response of the H₂ adducts is consistent with a scenario where the ring-closed product is less reactive toward the H₂ tag than the exposed charge center on a single carbon atom in the phenide motif.

Associated Content

High-resolution mass spectrometry of 2-benzoylbenzoate, 2-formylbenzoate and their respective decarboxylates (Figures S1 and S2), two-color IR laser experiment of the decarboxylate of H₂-tagged 2-formylbenzoate (Figure S3), detailed analysis of experimental IR spectrum of the decarboxylate of H₂-tagged 2-benzoylbenzoate (Figure S4), and Cartesian coordinates of calculated structures (Tables S1-S11).

Acknowledgements

We thank the Air Force Office of Scientific research under AFOSR Award No. FA9550-18-1-0213 for support of our work on the dynamics of CO₂ activation reactions. JM, EHP, ARB, OM were supported by the NSF Center for Aerosol Impacts on Chemistry of the Environment (CAICE) for support of the aspect of this work (CHE-1801971) related to the photophysics of 4BBA as a proxy molecule for sea spray aerosol radical initiation species. ARB was also supported partially by the Yale STARS program, which provides research opportunities for URM students in STEM fields. TS was supported by a Walter-Benjamin Scholarship by the Deutsche Forschungsgemeinschaft (Projektnummer 459401225). K.G. thanks the Fonds National de la Recherche, Luxembourg for funding the project GlycoCat (13549747) and the Fulbright Program for funding his research stay at Yale University.

References

1. O'Hair, R.; Rijs, N. Gas Phase Studies of the Pesci Decarboxylation Reaction: Synthesis, Structure, and Unimolecular and Bimolecular Reactivity of Organometallic Ions. *Acc. Chem. Res.* **2015**, *48* (2), 329-340.

2. Tian, Z.; Kass, S. Carbanions in the Gas Phase. *Chem. Rev.* **2013**, *113* (9), 6986-7010.

3. Graul, S. T.; Squires, R. R. On the Existence of Alkyl Carbanions in the Gas-Phase. *J. Am. Chem. Soc.* **1988**, *110* (2), 607-608.

4. Squires, R. R. Gas-Phase Carbanion Chemistry. Acc. Chem. Res. **1992**, 25 (10), 461-467.

5. Dickenson, G. D.; Niu, M. L.; Salumbides, E. J.; Komasa, J.; Eikema, K. S. E.; Pachucki, K.; Ubachs, W. Fundamental Vibration of Molecular Hydrogen. *Phys. Rev. Lett.* **2013**, *110* (19), 193601.

6. Lineberger, W.; Johnson, M.; Martinez, T. Once upon Anion: A Tale of Photodetachment. *Annu. Rev. Phys. Chem.* **2013**, *64*, 21-36.

7. Paul, M.; Peckelsen, K.; Thomulka, T.; Neudorfl, J.; Martens, J.; Berden, G.; Oomens, J.; Berkessel, A.; Meijer, A.; Schafer, M. Hydrogen tunneling avoided: enol-formation from a charge-tagged phenyl pyruvic acid derivative evidenced by tandem-MS, IR ion spectroscopy and theory. *Phys. Chem. Chem. Phys.* **2019**, *21* (30), 16591-16600.

8. Schafer, M.; Peckelsen, K.; Paul, M.; Martens, J.; Oomens, J.; Berden, G.; Berkessel, A.; Meijer, A. Hydrogen Tunneling above Room Temperature Evidenced by Infrared Ion Spectroscopy. *J. Am. Chem. Soc.* **2017**, *139* (16), 5779-5786.

9. Menges, F. S.; Perez, E. H.; Edington, S. C.; Duong, C. H.; Yang, N.; Johnson, M. A. Integration of High-Resolution Mass Spectrometry with Cryogenic Ion Vibrational Spectroscopy. *J. Am. Soc. Mass Spectrom.* **2019**, *30* (9), 1551-1557.

10. Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Mennucci, B.; Petersson, G. A.; Nakatsuji, H.; Caricato, M.; Li, X.; Hratchian, H.

P.; Izmaylov, A. F.; Bloino, J.; Zheng, G.; Sonnenberg, J. L.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; J. A. Montgomery, J.; Peralta, J. E.; Ogliaro, F.; Bearpark, M.; Heyd, J. J.; Brothers, E.; Kudin, K. N.; Staroverov, V. N.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Rega, N.; Millam, J. M.; Klene, M.; Knox, J. E.; Cross, J. B.; Bakken, V.; Adamo, C.; Jaramillo, J.; Gomperts, R.; Stratmann, R. E.; Yazyev, O.; Austin, A. J.; Cammi, R.; Pomelli, C.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Zakrzewski, V. G.; Voth, G. A.; Salvador, P.; Dannenberg, J. J.; Dapprich, S.; Daniels, A. D.; Farkas, Ö.; Foresman, J. B.; Ortiz, J. V.; Cioslowski, J.; Fox, D. J. *Gaussian 09, Revision D.01*, Gaussian, Inc.: Wallingford, CT, 2009.

11. Kashinski, D. O.; Chase, G. M.; Nelson, R. G.; Di Nallo, O. E.; Scales, A. N.; VanderLey, D. L.; Byrd, E. F. C. Harmonic Vibrational Frequencies: Approximate Global Scaling Factors for TPSS, M06, and M11 Functional Families Using Several Common Basis Sets. J. Phys. Chem. A **2017**, *121* (11), 2265-2273.

12. Kamrath, M. Z.; Relph, R. A.; Guasco, T. L.; Leavitt, C. M.; Johnson, M. A. Vibrational Predissociation Spectroscopy of the H₂-Tagged Mono- and Dicarboxylate Anions of Dodecanedioic Acid. *Int. J. Mass Spectrom.* **2011**, *300* (2-3), 91-98.

13. Tolstorozhev, G. B.; Skornyakov, I. V.; Bel'kov, M. V.; Shadyro, O. I.; Brinkevich, S. D.; Samovich, S. N. IR spectra of benzaldehyde and its derivatives in different aggregate states. *Opt. Spectrosc.* **2012**, *113* (2), 179-183.

14. Louris, J. N.; Cooks, R. G.; Syka, J. E. P.; Kelley, P. E.; Stafford, G. C.; Todd, J. F. J. Instrumentation, Applications, and Energy Deposition in Quadrupole Ion-Trap Tandem Mass-Spectrometry. *Anal. Chem.* **1987**, *59* (13), 1677-1685.

15. Molesworth, S.; Leavitt, C. M.; Groenewold, G. S.; Oomens, J.; Steill, J. D.; van Stipdonk, M. Spectroscopic Evidence for Mobilization of Amide Position Protons During CID of Model Peptide Ions. *J. Am. Soc. Mass Spectrom.* **2009**, *20* (10), 1841-1845.

16. Harrilal, C. P.; DeBlase, A. F.; Fischer, J. L.; Lawler, J. T.; McLuckey, S. A.; Zwier, T. S. Infrared Population Transfer Spectroscopy of Cryo-Cooled Ions: Quantitative Tests of the Effects of Collisional Cooling on the Room Temperature Conformer Populations. *J. Phys. Chem. A* **2018**, *122* (8), 2096-2107.

17. Shin, J.-W.; Hammer, N. I.; Johnson, M. A.; Schneider, H.; Gloss, A.; Weber, J. M. An Infrared Investigation of the $(CO_2)_n^-$ Clusters: Core Ion Switching from Both the Ion and Solvent Perspectives. *J. Phys. Chem. A* **2005**, *109*, 3146-3152.

18. Woo, H. K.; Wang, X. B.; Kiran, B.; Wang, L. S. Temperature-dependent photoelectron spectroscopy of methyl benzoate anions: Observation of steric effect in o-methyl benzoate. *J. Phys. Chem. A* **2005**, *109* (50), 11395-11400.

19. Gerardi, H. K.; DeBlase, A. F.; Su, X.; Jordan, K. D.; McCoy, A. B.; Johnson, M. A. Unraveling the Anomalous Solvatochromic Response of the Formate Ion Vibrational Spectrum: An Infrared, Ar-Tagging Study of the HCO_2^- , DCO_2^- , and HCO_2^- ·H₂O Ions. *J. Phys. Chem. Lett.* **2011**, *2* (19), 2437-2441.

20. Solcà, N.; Dopfer, O. Protonated Benzene: IR Spectrum and Structure of C6H7+. *Angew. Chem., Int. Ed.* **2002,** *41* (19), 3628-3631.

21. Douberly, G. E.; Ricks, A. M.; Schleyer, P. v. R.; Duncan, M. A. Infrared Spectroscopy of Gas Phase Benzenium Ions: Protonated Benzene and Protonated Toluene, from 750 to 3400 cm–1. *J. Phys. Chem. A* **2008**, *112* (22), 4869-4874.

22. Eichinger, P. C. H.; Dua, S.; Bowie, J. H. A Comparison of Skeletal Rearrangement Reactions of Even-Electron Anions in Solution and in the Gas-Phase. Int. J. Mass Spectrom. **1994**, *133* (1), 1-12.

23. Currie, G. J.; Bowie, J. H.; Massywestropp, R. A.; Adams, G. W. Collision-Induced Dissociations of Substituted Benzyl Negative-Ions in the Gas-Phase - the Elimination of C4h4. J. Chem. Soc., Perkin Trans. 2 **1988**, (3), 403-408.

24. Amundson, L. M.; Owen, B. C.; Gallardo, V. A.; Habicht, S. C.; Fu, M. K.; Shea, R. C.; Mossman, A. B.; Kenttamaa, H. I. Differentiation of Regioisomeric Aromatic Ketocarboxylic Acids by Positive Mode

Atmospheric Pressure Chemical Ionization Collision-Activated Dissociation Tandem Mass Spectrometry in a Linear Quadrupole Ion Trap Mass Spectrometer. *J. Am. Soc. Mass Spectrom.* 2011, 22 (4), 670-682.
Cheng, K.; Zhao, B. L.; Qi, C. Z. Silver-catalyzed decarboxylative acylation of arylglyoxylic acids with arylboronic acids. *Rsc Adv.* 2014, 4 (89), 48698-48702.

For Table of Contents Use Only



structurally characterized by Cryogenic Ion Vibrational Spectroscopy

Structures and Chemical Rearrangements of Benzoate Derivatives Following Gas Phase Decarboxylation

Evan H. Perez, Tim Schleif, Joseph P. Messinger, Anna G. Rullán Buxó, Olivia C. Moss, Kim Greis, Mark A. Johnson*

The Table of Contents graphic summarizes the major findings of this paper in two reaction equations: Decarboxylation of 2-formylbenzoate yields 2-formylphenide while decarboxylation of 2-benzoylbenzoate results in a fluorenone-like structure via an intramolecular nucleophilic ringclosing, presumably via the intermediary phenide. The structural assignments in this paper were performed via Cryogenic Ion Vibrational Spectroscopy.