Proton affinities and ion enthalpies*

John L Holmes1, Nick A van Huizen2,3 and Peter C Burgers2

Abstract

Proton affinities of a number of alkyl acetates (CH3–C(=O)–OR) and of methyl alkanoates (R–C(=O)–OCH3, R=H, alkyl) have been assembled from the literature or measured using the kinetic method. It was observed that the proton affinities for the isomeric species CH3–C(=O)–OR and R–C(=O)–OCH3 are almost identical, an unexpected result as the charge in these protonated ester molecules is largely at the keto carbon atom and so this site should be more sensitive to alkyl substitution. Analysis of the data, including those from lone pair ionisation and core-electron ionisation experiments available from the literature, indicate that after protonation, extensive charge relaxation (or polarisation) takes place (as is also the case, according to the literature, after core-electron ionisation). By contrast, after lone pair ionisation, which results in radical cations, such relaxation processes are relatively less extensive. As a consequence, changes in ion enthalpies of these protonated molecules follow more closely the changes in neutral enthalpies, compared with changes in enthalpies of the corresponding radical cations, formed by electron detachment. Preliminary analyses of published energetic data indicate that the above finding for organic esters may well be another example of a more general phenomenon.

Keywords

Proton affinity, kinetic method, ionisation energy, core-electron ionisation, ion stabilisation, tandem mass spectrometry, gas-phase ion chemistry

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Introduction

The ionisation energy (IE) of a molecule M is given by

\[ \text{IE}(M) = \Delta_f H^0(M^+) - \Delta_f H^0(M) \]  

(1)

From appropriate IE measurements, the enthalpies of formation of M^+, \( \Delta_f H^0(M^+) \), may be assessed.

The proton affinity (PA) of the molecule is given by

\[ \text{PA}(M) = \Delta_f H^0(M) + \Delta_f H^0(H^+) - \Delta_f H^0(MH^+) \]  

(2)

From PA measurements \( \Delta_f H^0(MH^+) \) may be obtained.

The gas phase enthalpies of formation of a significant number of organic cations (also called ion enthalpies) have been determined from IE and PA measurements and also by appropriate appearance energy (AE) determinations.\(^1\) Thus for example, \( \Delta_f H^0[\text{t-C}_4\text{H}_9]^+ \) has been derived from the IE of the t-butyl radical, the PA of isobutene and also by computation, leading to \( \Delta_f H^0[\text{t-C}_4\text{H}_9]^+ = 713 \pm 3 \text{ kJ/mol} \).\(^2,3\) Similarly, \( \Delta_f H^0[\text{CH}_3\text{C}^+\text{OHCH}_3]^+ \) has been derived from AE measurements and the PA of acetone, \( \Delta_f H^0 = 499 \pm 3 \text{ kJ/mol} \).\(^4\) Based on such measurements, it has been well established in a number of publications\(^5-13\) that plots of \( \Delta H_f \) for a series of organic cations, in which an electron releasing group is successively added to the formal charge bearing site, are a simple exponential function of ion size \( [\ln(n)] \), where \( n \) is the number of atoms in the ion and is commonly assumed to relate directly to ion size. These \( \Delta H_f \) versus \( \ln(n) \) plots have been shown to be linear for ethyl,\(^5-8\) methyl,\(^5-8,9\) hydroxy,\(^5,7-9\), methoxy,\(^5,7-9\) and amino\(^5,7-9\) substitutions at the formal charge site in species such as methane,\(^5,12\) the methyl cation,\(^5,7,8,13\) olefinic,\(^5,7,8,13\) and aromatic hydrocarbons\(^5,11\) and compounds containing N,\(^5,7,8,10\) Si, Ge, Sn and Pb.\(^12\)

In addition to providing a method by which thermochromic data may be estimated, such plots provide...
physicochemical information. For example, from these observations the location of greatest charge density in an ion may be assigned. In many textbooks, the ions \(+\text{CH}_2\text{OH}, \text{CH}_3\text{CHOH} and \text{CH}_2\text{OCH}_3\) are displayed as oxonium ions, with the formal charge on oxygen, e.g. \(\text{CH}_2=\text{O}^+\text{H}\). However, the \(\Delta H^0\) values for these ions, 711, 592 and 667 kJ/mol, respectively, show that the stabilising effect of the methyl substituent at carbon in \(+\text{CH}_2\text{OH}\) is much greater than at oxygen. It follows that the charge density is greater at C than at O.

It has been established, from a comparison of oxygen ls core ionisation energies,\(^{14,15}\) that organic acids and esters, \(\text{R}_1\text{C}(=\text{O})\text{O}-\text{R}_2\) (for example the isomers acetic acid and methyl formate), protonate at the keto oxygen. With the formal charge on the keto carbon, e.g. \(\text{CH}_3=\text{C}(\text{OH})_2\text{H}\) and \(\text{H}^+\text{C}(\text{OH})\text{OCH}_3\), it is not surprising that the former has a significantly lower \(\Delta H^0\) (314 vs. 391 kJ/mol), because in \(\text{CH}_3=\text{C}(\text{OH})_2\text{H}\) three electron donating substituents are attached to the formal charge bearing site, compared with two for \(\text{H}^+\text{C}(\text{OH})\text{OCH}_3\). However, this stabilisation is not reflected in the PAs of acetic acid and methyl formate, which are, within experimental error, the same, PA = 783 ± 1 kJ/mol.\(^1\) This is also unexpected in that the IEs are different, with acetic acid having the lower IE (10.62 ± 0.02 eV, compared to 10.84 ± 0.02 eV for methyl formate) and consistent with the formal charge at the keto group in the radical cation. Indeed, it appears from a survey of the literature (and present work) that the PA of a number of \(\text{R}_1\text{C}(=\text{O})\text{O}-\text{R}_2\) and \(\text{R}_2\text{C}(=\text{O})\text{O}-\text{R}_1\) isomers are remarkably similar, despite the charge of the protonated species being at (the carbon atom of) the keto group.

The purpose of this paper is to establish whether the PAs of \(\text{R}_1\text{C}(=\text{O})\text{O}-\text{R}_2\) and \(\text{R}_2\text{C}(=\text{O})\text{O}-\text{R}_1\) are indeed similar for a wide variety of substituents \(\text{R}\), and if so, to provide a rationale. During the course of this work we have compiled and compared the IEs and PAs for a large number of compounds and we have also analysed our data using core-electron ionisation energies available from the literature.

**Experimental**

Collision-induced dissociation (CID) experiments were performed using a Bruker Esquire ESI ion trap mass spectrometer as described in Jobst et al.,\(^{16}\) van Huizen et al.\(^{17}\) and Burgers et al.\(^{18}\) The esters were dissolved in methanol at a concentration of 0.01 M. The proton bound ester dimers were generated by infusion of the ester solutions with an acidified (0.1% trifluoroacetic acid) water/methanol (50/50) mixture in a ratio of 10: 190 at an infusion rate of 240 \(\mu\text{L} \text{h}^{-1}\). The kinetic method\(^{19,20}\) was employed to determine relative PA values of the esters. Product ion intensities (R) from proton bound heterodimers were obtained at 50% survival yield\(^{21}\) and the obtained lnR values were plotted against the known PA values of reference ester molecules (methyl acetate and octyl acetate\(^{17}\)) to obtain the new PA values. We\(^{16-18}\) and others\(^{22}\) have observed that the product ion ratio observed in ion trap experiments depends only slightly on the amplitude and so using low amplitudes (corresponding to high survival yields) does not give more accurate relative PA data, but leads to loss of signal strength only. Further details can be found in Jobst et al.\(^{16}\) van Huizen et al.\(^{17}\) and Burgers et al.\(^{18}\) The D-labelled esters \(\text{CD}_3\text{C}(=\text{O})\text{O}-\text{R}\) and \(\text{CH}_3\text{C}(=\text{O})\text{O}-\text{CD}_3\) were prepared by small scale esterification of \(\text{CD}_3\text{C}(=\text{O})\text{OH}\) with ROH and of \(\text{CH}_3\text{C}(=\text{O})\text{O}-\text{CD}_3\) with \(\text{CD}_2\text{OH}\), respectively.

**Results and discussion**

**The PAs of organic esters**

In the following we discuss the PAs of compounds of the type \(\text{R}_1\text{C}(=\text{O})\text{O}-\text{R}_2\) and \(\text{R}_2\text{C}(=\text{O})\text{O}-\text{R}_1\) but have limited ourselves to \(\text{R}_1=\text{CH}_2\text{C}(=\text{O})\text{OH}\), \(\text{HC}(=\text{O})\text{OCH}_3\) and \(\text{CH}_3\text{C}(=\text{O})\text{OH}\), and methyl formate, \(\text{HC}(=\text{O})\text{OCH}_3\) are remarkably similar, 783 ± 1 kJ/mol.

In a preliminary experiment we generated the proton bound dimer [\(\text{CD}_3\text{C}(=\text{O})\text{OH}][\text{HC}(=\text{O})\text{OCH}_3]\] and observed that it dissociated to \(\text{m}/\text{z}\) 64 ([\(\text{CD}_3\text{C}(\text{OH})_2\text{H}\)]) and to \(\text{m}/\text{z}\) 61 ([\(\text{HC}(\text{OH})\text{OCH}_3\)]) in a ratio of 2.12 ± 0.20. It has been established that in such kinetic method experiments for these classes of compounds, the equation \(\Delta\text{PA} (\text{kJ/mol}) = 2\ln R\) holds.\(^{16-18}\) where R is the ratio of the product ions, in this case 2.12. This leads to a difference in PAs of acetic acid and methyl formate of only 1.5 ± 0.4 kJ/mol and so the PAs of acetic acid and methyl formate are indeed closely similar. The complete list of our measured PA values for the alkyl acetates and methyl alkanoates is shown in Table 1.

This table also includes some relevant values for branched alkyl groups and values from the literature.\(^1\) The first three columns in Table 1 give PA data for the alkyl acetates and the last three columns show the PA data for the isomeric methyl alkanoates (in a given row), for example the PA of hexyl acetate is 840.5 kJ/mol, while that of its isomer, methyl heptanoate, is 841.3 kJ/mol. Some of these observations deserve extra comment. Generally speaking, our PA values compare well with the literature values, see for example the values for propyl acetate, methyl pentanoate and methyl cyclohexanecarboxylate (MCC). However, some other values differ by ±2 kJ/mol or more. For example, our PA for isopropyl acetate (841.7 kJ/mol) is 5.1 kJ/mol larger than the literature value (836.6 kJ/mol).
However, according to the literature (see Table 1), the PA of propyl acetate and isopropyl acetate are the same. This could reasonably be expected, because the charge in the protonated species will be on the carbonyl group, and so branching remote from the charge site will have little effect. Our results for the PA determinations of propyl acetate and isopropyl acetate are given in Figure 1. Here, the PA of both compounds have been measured against the bases ethyl-, butyl-, hexyl- and octyl acetate (see Table 1) and it can be seen that the PA of isopropyl acetate is 4.8 kJ/mol larger than that of propyl acetate. In agreement with our values is the finding that the PA of isopropyl formate (811 kJ/mol) is 6 kJ/mol larger than the PA of propyl formate (805 kJ/mol). These matters will be discussed below.

Similarly, our derived PA value for methyl benzoate is 6.8 kJ/mol lower than the literature value. In a control experiment, methyl benzoate was measured against MCC and from this result it followed that the PA of methyl benzoate is 2.1 kJ/mol lower than that of MCC, not 4.7 kJ/mol higher. To evaluate the internal consistency of the data presented in Table 1, several such control experiments were performed; for example methyl nonanoate was also measured against methyl 2-methylbutyrate which have the same PAs. The observed intensity ratios R for [(methyl nonanoate) + H⁺]/[(methyl 2-methylbutyrate) + H⁺] formed from the proton bound dimer was found to be 1.1 (lnR = 0.1) leading to ΔPA = (0.2 ± 0.4) kJ/mol.

It was found that the PA of phenyl formate was much lower than all the other ester compounds listed in Table 1, and so the PA of phenyl formate was measured against some nitriles as reference bases (butane-, pentane-, heptane-, nitrile and tert butylcyanide). One particular example deserves further mention. We could not generate protonated tert butyl acetate in our ion trap experiments; rather we observed extensive formation of m/z 61, protonated acetic acid. We tentatively conclude that this species is formed by the hydrolysis of transient protonated tert butyl acetate by residual water present in the ion trap; CH₃–C(OH)–OR + H₂O → CH₃–C(OH)₂ + ROH, where R = t-buty1; such hydrolysis reactions have been observed previously in ion trap experiments.

| Alkyl acetates | Methyl alkanoates |
|----------------|-------------------|
| Homologue      | PA¹               | PA²               | Homologue      |
|----------------|-------------------|-------------------|----------------|
| Formic acid    | 742.0             | 742.0             | Formic acid    |
| Acetic acid    | 783.7             | 782.5             | Methyl formate |
| Methyl acetate | 821.6             | 821.6             | Methyl acetate |
| Ethyl acetate  | 839.0             | 839.0 ± 0.4       | 830.2          | 832.1 ± 0.4   | Methyl propanoate |
| Propyl acetate | 836.6             | 836.9 ± 0.5       | 836.4          |               | Methyl butanoate |
| Butyl acetate  | 839.0 ± 0.6       | 839 ± 1           | 838.9 ± 0.6    | Methyl pentanoate |
| Pentyl acetate | 839.8 ± 0.7       | 840.5 ± 0.7       | 841.3 ± 0.7    | Methyl hexanoate |
| Hexyl acetate  | 836.9 ± 0.5       | 837.9 ± 0.5       | 838.9 ± 0.6    | Methyl heptanoate |
| Heptyl acetate | 841.0 ± 0.8       | 844 ± 1           | 841.8 ± 0.8    | Methyl octanoate |
| Octyl acetate  | 841.3 ± 0.8       | 841.3 ± 0.8       | 842.4 ± 0.8    | Methyl nonanoate |
| Nonyl acetate  | 841.8 ± 0.9       | 842.9 ± 0.9       | 843.0 ± 0.9    | Methyl decanoate |
| Decyl acetate  | 842.4 ± 0.9       | 843.4 ± 0.9       | 844.0 ± 0.9    | Methyl undecanoate |
| Isopropyl acetate | 836.6        | 841.7 ± 0.7       | 836.6          | 838.5 ± 0.7   | Methyl isobutyrate |
| sec. butyl acetate | 844.5 ± 0.5     | 842.4 ± 0.5       | 843.5 ± 0.5    | Methyl 2-methylbutyrate |
| iso butyl acetate | 839.0 ± 0.5     | 839.5 ± 0.5       | 841.0 ± 0.5    | Methyl isovalerate |
| tert. butyl acetate | 845.2 ± 1.6³     | 845.5             | 843.3 ± 0.6    | Methyl pivalate |
| Cyclohexyl acetate | 846.1 ± 0.5    | 846.2             | 845.8 ± 0.5    | Methyl cyclo C₆H₁₃ (MCC) |
| Phenyl acetate | 837.7 ± 0.5       | 850.5             | 843.7 ± 0.5    | Methyl benzoate |
| Vinyl acetate  | 813.9             | 825.8             | 829.3 ± 0.5    | Methyl acrylate |
| Phenyl formate | 802.6 ± 0.4       | 821.1             | 821.1          | Benzoic acid |

PA: proton affinity.
¹From Lias et al. ³This work.
²Using Na⁺ bound dimer, see text.
PA and methyl group substitutions

It can be seen from Table 1, that except for the isomeric pairs vinyl acetate/methyl acrylate and phenyl formate/ benzilic acid, all other paired isomers have very similar PA values (minor, secondary effects, will be discussed later). This means that for saturated and branched R-chains, the PAs of the isomers R–C(=O)–OCH₃ and CH₃–C(=O)–OR are almost equal, despite the finding that protonation of organic esters occurs exclusively at the keto group and that the charge is predominantly at the keto C atom. A confirmation of this result comes from labelling experiments: the labelled proton bound dimer of methyl propanoate [CH₃CH₂C(=O)OCH₃] show a ratio for m/z 92 and m/z 89 of 1.01, whereas the labelled proton bound dimer of methyl propanoate [CH₃CH₂C(=O)OCH₃] show a ratio for m/z 92 and m/z 89 of 1.05.)

The above finding, namely that the PAs of the isomers R–C(=O)–OCH₃ and CH₃–C(=O)–OR are almost equal, can be extended to include organic acids and alkyl formates as far as their PAs have been determined. Starting from formic acid as the prototype molecule, we can substitute the C–H hydrogen or the O–H hydrogen by a methyl group and then perform further homologous substitutions. We can then list, for each substitution, the increments in PA, leaving constant; these matters will be discussed in detail elsewhere.

In addition to the observed correlations between PA and IE, good correlations between PAs and core-electron ionisation energies for certain categories of molecules have also been established; these have been referred to as the Martin–Shirley correlations. For example, for alcohols, it was found that a plot of PA versus IE does not yield a slope of −1.0 (as would be expected from equation (3)), but a less negative slope (ca. −0.6). For example, consider successively methyl group substitutions in NH₃: NH₃ → CH₃NH₂ → (CH₃)₂NH → (CH₃)₃N where the plot of PA against IE yields a straight line with a slope of −0.44, see Figure 2(a). According to equation (3), a slope less negative than −1 indicates that the N–H homolytic bond dissociation energy decreases as the PA increases, see Figure 2(b), see also Choi and Boyd. If a plot of the PA against IE gives a slope of close to −1 (as is the case for the phosphine methyl group substitutions), then the homolytic bond dissociation energy remains constant; these matters will be discussed in detail elsewhere.
Figure 2. (a) PA as a function IE for the amines NH$_3$, CH$_3$NH$_2$, (CH$_3$)$_2$NH and (CH$_3$)$_3$N. The slope is $-0.44$. (b) Homolytic bond dissociation energies (D) of the protonated amines as a function of PA.

IE: ionisation energy; PA: proton affinity.
that the change in PA from one molecule to another is almost exactly equal to the change in core IE, i.e. a plot of the oxygen 1s IE against the PA gives a slope of $-1$. It was argued that adding a positive charge to a nucleus (core ionisation) and adding a positive charge adjacent to the nucleus (protonation) will give rise to similar molecular electronic relaxation effects. The proton attachment reaction can be split into two hypothetical steps. In the first, the proton attaches itself to an atom (for example oxygen) without flow of charge in the molecular framework; shifts in energy of this “reaction” are due to differences in the electron density about the oxygen in the ground state and are inductive effects. In the second hypothetical step, the excess charge is distributed over the whole molecule to minimise Coulombic repulsion (relaxation or polarisation effects). Several groups agree that differences in relaxation energies (rather than differences in inductive effects) dominate both core ionisation and protonation processes. Furthermore, if linear relationships exist between PA and core ionisation energies and between PA and IE, then a linear relation should also exist between IE and core ionisation energies. This is illustrated in Figure 3 for the amines discussed above, in which the PAs and IEs are plotted against the N(1s) binding energies of the amines \cite{Mills15, Martin30} (The N(1s) binding energies are averages from Mills et al. \cite{Mills15} and Martin and Shirley \cite{Martin30}). It can be seen that both IE and PA correlate linearly with the N(1s) binding energy. (Following Mills et al. \cite{Mills15} relative binding energies are negative.) However, the slope for the IE curve is significantly larger (2.9) than the slope of the PA curve (1.3) and this parallels the observation that the slope of the PA versus IE curve is significantly less negative than $-1$, see above. We thus conclude that for this system, the changes in PA are largely governed by relaxation effects, but that changes in IE also reflect changes in inductive effects. Similar plots can be made for other systems, and when such plots are made for organic esters, a remarkable result ensues, see Figure 4.

![Figure 3](image1.png)

**Figure 3.** IE and PA as a function of N(1s) ionisation energies for the amines. Values relative to NH$_3$ = 0. IE: ionisation energy; PA: proton affinity.

![Figure 4](image2.png)

**Figure 4.** IE and PA as a function of O(1s) ionisation energies for esters. Values relative to methyl formate = 0. IE: ionisation energy; PA: proton affinity.
where we show that the curve for PA against oxygen 1s IE has a slope of even less than 1 (0.73), indicating that after protonation, extensive relaxation effects operate and that inductive effects are virtually non-existent. Even for the IE versus oxygen 1s IE curve, the slope is only 1.05 indicating that for the radical cations too, relaxation effects are important. In this respect it should be mentioned that relaxation energies have a tendency to increase with molecular size.30 Thus we conclude that for the above organic esters the PA values are indeed largely governed by molecular size, whereas this is less so for the IE values. In such cases, changes in the heat of formation of the protonated species closely follow those of the neutral species and so any conclusions drawn from these PA data as to charge location in the protonated species should be viewed critically.

It can be seen from Table 1 that the PA of methyl acrylate is close to (and slightly lower than) that of methyl propanoate. This is in sharp contrast to other $\alpha,\beta$ unsaturated keto compounds whose PA values are significantly larger (by 10–60 kJ/mol)25 than their corresponding saturated analogues. This stabilisation is thought to arise from participation of canonical resonance structures, but clearly such structures are not involved in the case of protonated methyl acrylate. Again, we conclude that the PAs of methyl propanoate and methyl acrylate are primarily determined by size.

We also tried to measure the PA of methyl propiolate, HC≡C–C(=O)–O–CH3 but this experiment failed: no proton bound dimers could be formed from this compound, it undergoing rapid trimerisation in our ESI experiments to produce m/z 253. The latter ion loses CH3OH and 44 Da and so it is most probably not a proton bound trimer, and we propose benzenetricarboxylic acid, trimethyl ester.

**Comparison of ion stabilisation effects from IE and PA measurements**

An alternative way to view ion stabilisation effects arising from methyl substitution at a formal charge bearing site versus a non-charge bearing site is presented in Figure 5. Starting from CH3OH, we can substitute the O–H hydrogen by a methyl group to yield CH3OCH3. The stabilisations (measured by a decrease in IE or an increase in PA) are given by the bars, and it can be seen that the increase in PA is about one half the decrease in IE, paralleling the effects discussed above. These stabilisations are those arising from methyl substitution at the charge bearing site. When we substitute the hydrogen at a non-charge bearing site, we find as expected, much smaller stabilisation energies, and again the increase in PA is smaller than the decrease in IE.

A similar situation pertains to ionised acetaldehyde, CH3CH=O. Here again, methyl substitution at the charge bearing site (to produce acetone) is associated with larger stabilisation energies (both from IE and PA measurements) than methyl substitution at the non-charge bearing site (to yield propanal), and once more the changes in PA are significantly smaller than the changes in IE.

Similar plots can also be produced for methyl substitutions in formic acid: substitution at the formal charge-bearing site yields acetic acid, while substitution at the non-charge bearing site gives methyl formate, the compounds discussed above. Three features then emerge from Figure 5(c):

1. The differences in IE are consistent with the formal location of the charge on the keto group, but the effect is relatively small;
2. The differences in IEs and PAs are also relatively small, and
3. There is almost no difference in the PA values, see also above. Such very small differences in PA values could erroneously be taken to show that protonation in organic esters also occurs at the ether oxygen, whereas the cause of this effect lies, see Figure 4, in the dominant relaxation processes occurring after protonation (PA), a process.
which is of relatively lesser importance after lone pair ionisation (IE). Indeed, according to DFT and MP2 ab initio calculations, the positive charge in both protonated methyl formate and protonated acetic acid is largely at the carbonyl C (protonated methyl formate: \(+0.735\) (DFT), \(+0.923\) (MP2); protonated acetic acid: \(+0.926\) (DFT), \(+1.110\) (MP2); Mayer, Personal communication, 2017).

Based on these above findings, we have collected all the IE and PA values available in the NIST website, together with additional data found in the literature. We will report on a detailed analysis of these data in a future publication, but a generalisation (already apparent from the literature) rapidly emerges, namely that for a given category of molecules, the change in PA is usually smaller than the change in IE (with the notable exception of successive methyl substitutions in PH\(_3\)). Thus for a given category of molecules, the changes in ion enthalpies of the protonated species more closely follow those in the enthalpies of the neutral molecules, compared to changes in the ion enthalpies of the radical cations. Another feature of the above ab initio calculations is that the incoming proton, once attached to the molecule, retains ca 58\% of its charge, with the remainder spread over the other atoms in the molecule. This effect has been observed previously, for example in nitriles,\(^{33}\) oxygen containing compounds,\(^{34}\) amines\(^{34}\) and even amino acids.\(^{35}\) Whether this phenomenon also relates to the observation that ion enthalpies of protonated species tend to more closely follow neutral enthalpies (which would appear logical) is also a matter of current investigation.

Secondary effects upon protonation of organic esters

It can be seen from Table 1 that both phenyl formate and vinyl acetate have lower PAs than their isomers, benzoic acid and methyl acrylate, respectively. Scaled molecular models (for example Dreiding ball-and-stick models) show that the vinyl group in vinyl acetate is very close to the CO function, whereas there is no such interference in the isomer. A similar situation obtains for the phenyl analogues. Also, in protonated vinyl acetate there is a very close proximity between the terminal CH\(_2\) group and the protonated carbonyl moiety. Such post protonation effects could well affect the stability of the protonated species, but in the absence of more ab initio calculations such conclusions must remain speculative. As for secondary effects, it is interesting to note that the PA of ethyl acetate (833.7 kJ/mol) is larger than that of methyl propanoate (832.1 kJ/mol) but with longer alkyl chains the PAs cross (at the pair butyl acetate/methyl pentanoate) to become slightly smaller for the acetates having long chains. This is shown in Figure 6(a), which plots the difference in PA for the isomeric alkyl acetates and methyl alkanoates as a function of chain length, data from Table 1. More precise values for the relative PAs of these isomeric esters can be obtained by performing the kinetic method experiments with isotopically labelled isomers, by observing the product ions from proton bound dimers \([\text{CD}_3\text{C}(=\text{O})\text{OR}]^{+}\)/\([\text{RC}(=\text{O})\text{OCH}_3]^{-}\), see Figure 6(b). First, these result again show that the PAs of the isomeric species CH\(_3\)–C(=O)–OR and of R–C(=O)–OCH\(_3\) are very similar indeed. Second, as the chain gets longer, the PA of the alkyl acetate relative to that of the isomeric methyl alkanoate drops and this behaviour becomes asymptotic. This could indicate that after protonation, some H-bonding takes place between the O–H proton and the hydrocarbon chain, an effect that will be relatively more pronounced in the smaller chain alkyl acetate ions (for example for protonated ethyl acetate such bonding would result in a six-membered ring, but for methyl propanoate, a more strained five-membered ring would ensue). Such effects might also explain why the PA of isopropyl acetate is significantly larger (by 3.2 \(\pm\) 0.5 kJ/mol) than that of its isomer.

Figure 6. Differences in PA for the isomeric alkyl acetates (CH\(_3\)–C(=O)–OR) and of methyl alkanoates (R–C(=O)–OCH\(_3\)) as a function of R: (a) from laddering experiments, see Table 1; (b) from the isotope labelled dimers \([\text{CD}_3\text{C}(=\text{O})\text{OR}]^{+}\)/\([\text{RC}(=\text{O})\text{OCH}_3]^{-}\), PA: proton affinity.
methyl isobutyrates. Again, in the absence of high level ab initio calculations such an interpretation must remain speculative.

Summary

The PAs of the isomeric alkyl acetates CH$_3$–C(=O)–OR and methyl alkanoates R–C(=O)–OCH$_3$ were found to be almost identical (to within 2 kJ/mol) for a large number of substituents R. Despite the charge in the protonated species being largely on the carbonyl C atom (as indicated by ab initio calculations), and so this site would be expected to be more sensitive to stabilisation effects by alkyl substitution. From a comparison of PA data and core-electron ionisation energies, it is concluded that after protonation of the ester molecules, extensive charge relaxation or polarisation takes place, overwhelming any inductive effects. Thus differences in the heat of formation of these protonated species closely follow differences in heat of formation of the neutral species. Whether the observed behaviour for the organic esters is an (extreme) example of a more general phenomenon is currently being investigated.

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