Surface $pK_a$ of octanoic, nonanoic, and decanoic fatty acids at the air–water interface: applications to atmospheric aerosol chemistry
Surface \(pK_a\) of octanoic, nonanoic, and decanoic fatty acids at the air–water interface: applications to atmospheric aerosol chemistry†

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There exists large uncertainty in the literature as to the \(pK_a\) of medium-chain fatty acids at the air–water interface. Via surface tension titration, the surface-\(pK_a\) values of octanoic (C8), nonanoic (C9), and decanoic (C10) fatty acids are determined to be 4.9, 5.8, and 6.4, respectively. The surface-\(pK_a\) determined with surface tension differs from the bulk value obtained during a standard acid–base titration. Near the surface-\(pK_a\) of the C8 and C9 systems, surface tension minima are observed and are attributed to the formation of surface-active acid–soap complexes. The direction of the titration is shown to affect the surface-\(pK_a\) of the C9 system, as the value shifts to 5.2 with NaOH titrant due to a higher concentration of Na+ ions at pH values close to the surface-\(pK_a\). As the reactivity and climate-relevant properties of sea spray aerosols (SSA) are partially dictated by the charge and surface activity of the organics at the aerosol–atmosphere interface, the results presented here on SSA-identified C8–C10 fatty acids can be used to better predict the health and climate impact of particles with significant concentrations of medium-chain fatty acids.

Introduction

The air–water interface provides a unique environment for chemistry due to its asymmetry and 2-D intermolecular interactions.1–2 As the local environment at the interface is different from the bulk, one would expect the \(pK_a\) (an expression of acidity) of surface-active molecules to deviate from bulk values. This deviation can be attributed to increased interactions between molecular groups as a result of van der Waals forces between longer hydrocarbon chains,3 a change in the local pH at the surface due to the charge density of the surface film,4–6 or a change in the effective dielectric constant of the medium surrounding the headgroups of molecules at the interface.7–10

Fatty acids constitute a common class of surface-active molecules which are studied at the air–water interface. Some studies have shown that as the hydrophobic chain length of the acid increases, and the molecules become more surface active, the \(pK_a\) also increases.3,11 However, it has been commonly, and incorrectly, reported in literature that the \(pK_a\) of medium-chain fatty acids (C8–C10) at the interface is the same as that of their bulk counterpart.12–14 As the surface-\(pK_a\) of these medium-chain fatty acids seem ill-defined, one might be inclined to examine the trends of other physical properties (solubility and critical micelle concentration (CMC)) against carbon chain length for these saturated compounds (Fig. 1).15,16 Between two and six carbons, the acids are very soluble (from \(\infty\) to 0.97 g/100 g H2O) and have higher CMC values (indicating that...
a large concentration is necessary for the required molecular associations to form micelles). For acids with chain lengths of twelve or more carbon atoms, solubility and CMC remain relatively constant at low values. Between these two regimes lies the class of medium-chain fatty acids, which are on the cusp of being considered in the same context as their insoluble, surface-active (C_m ≥ 12) counterparts as opposed to predominantly bulk species (C_m ≤ 6).

While fatty acids have implications in many fields such as biological membranes,17–19 mineral flotation,20,21 and food science;22–34 of particular interest is aerosol chemistry. Breaking waves at the ocean surface generate and release sea spray aerosols (SSA) to the neighboring atmosphere. During this process, the organic molecules enriched in the sea surface microlayer (SSML) are transferred to the atmosphere as coatings on SSA. Although C_{36} and C_{18} fatty acids are found to be the most abundant in these organic films,13,25,26 C_{9}−C_{10} saturated fatty acids have also been identified in significant concentrations.13 Moreover, the concentrations of these shorter acids are likely to become more prevalent as the aerosol ages due to the oxidation of longer-chain fatty acids.27–30 SSA reactivity,31–34 lifetime,35,36 light-scattering37,38 and nucleating abilities39 are affected by the organic coatings at the aerosol–atmosphere interface. Surface tension is one of the governing parameters of several of these properties, and can be affected by the pH of the aerosol, and in turn, the surface-pK_a of the molecules at the interface. Upon entering the atmosphere, SSA will be at a pH close to that of the ocean (8.1),40 and any short- or medium-chain fatty acids have been thought to exist in their deprotonated state, and therefore would have low surface activity. However, upon aging in the atmosphere, reactions with gas phase acidic species (i.e. HCl, HNO_3) will cause these aerosols to become acidified,41 thus increasing the surface activity of the fatty acids and decreasing the surface tension at the interface. The pH at which this transformation (deprotonated to protonated) occurs is dictated by the surface-pK_a of the species at the interface, and will affect the surface-tension-dependent properties of the aerosol. For example, separation relative humidity (SRH) is a metric that can be used to determine the phase state of an aerosol, and is affected by pH and the pK_a of the organic fraction of SSA. The SRH of model SSA particles decreases at high pH as a result of the increased solubility of the organic components.12 As particle phase state has a large influence on the chemical and physical properties of an aerosol, an understanding of the pH at which the organic components become deprotonated is crucial.43

The primary goal of this work is to determine the surface-pK_a of medium-chain fatty acids at the air–water interface, which we expect to differ from their commonly used bulk values (~4.8). We utilize a surface tension titration technique to determine these values, and evaluate our results within the context of similar, yet controversial, studies in the literature.

**Experimental methods**

**Materials**

Octanoic acid (C_{8}, ≥ 99%, M_w 144.21 g mol^{-1}), nonanoic acid (C_{9}, ≥ 97%, M_w 158.24 g mol^{-1}), and decanoic acid (C_{10}, ≥ 98%, M_w 172.26 g mol^{-1}) were purchased from Sigma Aldrich (St. Louis, MO) and used without further purification. HCl (Trace Metal Grade) was purchased from Fisher Scientific (Waltham, MA), and NaOH pellets (99%) were purchased from Mallinckrodt (Paris, KY). A Barnstead Nanopure filtration system (D4741, Thermolyne Corporation, Dubuque, IA) with additional cartridges to remove organic material (D5026 Type I ORGANIC-free Cartridge Kit, Pretreat Feed) provided ultrapure water with a resistivity of at least 18.0 MΩ cm. Solutions of the C_{8}−C_{10} acids were prepared at concentrations of 1 mM at pH 12 for surface tension (γ) versus pH titrations with a 0.10 M HCl titrant. The C_9 system was also prepared at 1 mM in a pH 2 solution for titration with 0.10 M NaOH. 1 mM C_9 solutions were prepared in ultrapure water for standard acid–base titrations with 0.005 M NaOH. All experiments were conducted at 21 ± 1 °C and at a measured relative humidity of 43 ± 2%.

**Surface tension measurements**

Surface tension was probed with a platinum Wilhelmy plate and calculated via

\[
\gamma = \frac{F}{l \times \cos \theta}
\]  

where the force (F) is obtained from the measured difference in mass of the plate, θ is the contact angle between the plate and solution (assumed to be 0°),44 and l is the perimeter of the wetted surface (39.24 mm). Readings were obtained with a KSV NIMA (Biolin Scientific, Espoo, Finland) tensiometer and interface unit equipped with LayerBuilder software. The plate was heated by flame prior to measurements to remove any organic residue from the surface.

**Surface tension titration**

The use of surface tension to determine surface-pK_a was inspired by the work Dickhaus & Priefer and Sugawara et al.45,46 For the C_{8}−C_{10} acids, surface tension versus pH data was obtained via titration of a 50 mL aliquot of the C_9 solution with either acid or base while monitoring γ and pH (Accumet AB15 pH meter, Fisher Scientific). The pH probe was calibrated before the measurements with pH 1, 4, 6, 7, and 10 buffer solutions. The γ measurements were taken continuously as titrant was added to a 1 mM aliquot of fatty acid solution, allowing for the mass reading to stabilize before a measurement was recorded. Stirring was used to ensure efficient mixing of the added titrant. For titrant additions after which the mass reading did not change significantly, a wait time of 2 minutes was allotted prior to recording the surface tension mass reading.

Along more dynamic areas of the titration curves, an equilibration time of at least 10 minutes was required for the mass readings to stabilize. A reading was deemed stable when the mass only changed by approximately 2 mg around a central value. Dilution effects17 on the measured γ of the Gibbs monolayer were minimized with only a ~10% increase in volume from titrant addition (Fig. S1 and S2, ESI†). To compensate for the rising height of solution due to the addition of titrant, the plate was periodically raised to maintain a constant depth in solution. The error bars reported are dominated by the uncertainties from...
the pH meter measurements and the error of the surface tension data fits. Errors reported represent one standard deviation from the average of triplicate measurements.

Infrared reflection absorption spectroscopy

To probe the structure and ordering of the surfactant films at the interface, IRRAS spectra of the C₉ systems were collected using a custom built setup housed in an FTIR spectrometer (Spectrum 100, Perkin Elmer). The setup consists of two gold mirrors. The first mirror directs the incoming IR beam to the sample surface at an incidence angle of 46° relative to surface normal. The second mirror directs the reflected beam to a liquid nitrogen-cooled MCT detector. As IRRAS spectra are plotted in terms of reflectance–absorbance (RA = −log(R/R₀)), a background spectrum was collected of a bare pH 2, 5.6 (water), or 12 surface and used as R₀. This background solution was aspirated from the Petri dish and replaced with a 1 mM C₉ solution at the appropriate pH. The spectrum of this solution was used as R. Spectra of the C₉ systems were compared to that of a stearic acid (C₁₈) monolayer. Each spectrum is the result of coaveraging 300 scans over the full range (4000–450 cm⁻¹) at a resolution of 4 cm⁻¹. Spectra shown have been baseline-subtracted by a third order polynomial and are the average of at least 3 individual spectra.

Results & discussion

C₉₄ titration with 0.10 M HCl

To experimentally determine the surface-pKₐ of the C₆–C₁₀ medium-chain fatty acids, 1 mM solutions were titrated with 0.10 M HCl from a pH of 12 to a pH of 2. Representative surface tension vs. pH titration curves for the three acids are shown in Fig. 2. At a pH of 12, the C₉ acids are fully deprotonated, and thus relatively surface inactive. The surface tension curves begin at pH 12 and at surface tensions of approximately 70 mN m⁻¹, slightly depressed relative to pure water. As 0.10 M HCl is titrated into the solutions, the surface tension decreases, indicative of the formation of the more surface-active protonated species. The C₈ and C₉ curves develop an interesting feature from pH 6 to 4.5, hereafter referred to as the surface tension dip, where the surface tension decreases to a minimum value then rises to reach a plateau. The physical interpretation of this dip is discussed later.

Upon titration to pH < 4, the surface tension curves approach constant values. The values of surface tension at this point reflect the relative surface activity of each C₉ acid. C₁₀ approaches a surface tension value of 33.1 ± 0.3 mN m⁻¹ (the most surface active), followed by C₉ at 34.3 ± 1.6 mN m⁻¹, and C₈ at 52.9 ± 2.7 mN m⁻¹. The greater surface activity with chain length can be attributed to increasing van der Waals forces between the longer hydrocarbon chains.

A mathematical model is applied to the collected surface tension titration data to calculate the surface-pKₐ (pKₐₛ) of the selected C₉ fatty acids. This model (described in Appendix A) is adapted from that of Cratin⁴⁸ in his work on the interfacial pKₐ of stearic acid at the oil–water interface, and is expressed in terms of the change in surface tension with pH.

\[
\Delta \gamma = \frac{\Delta \gamma_{\text{max}}}{1 + 10^{(pH_p - pK_{a,s})}}
\]

When this surface activity model is applied to the systems studied here, a trend is observed between the surface-pKₐ and chain length. The fits of the data are shown in Fig. 3 and the surface-pKₐ values of the C₉ systems are presented in Table 1. As can be seen from the values in Table 1, the surface-pKₐ of the fatty acid increases with chain length from 4.9 to 6.4. This trend can be understood to be an additional consequence of increased van der Waals forces between fatty acid molecules with longer chains. Greater forces between the hydrophobic groups at the interface result in increased interactions between the carboxylate headgroups, thus making the acidic hydrogen more difficult to remove. However, the surface-pKₐ for the C₈ system (4.9 ± 0.2) is within the range of the accepted value of its bulk pKₐ (4.89). When a 1 mM C₈ system, prepared in water, is titrated with NaOH in a standard weak acid–strong base titration (Fig. S3, ESIF), a bulk pKₐ of 4.97 ± 0.05 is determined from the pH at the half-way point, which also is in close agreement with its literature pKₐ value of 4.96. The significant deviation of surface-pKₐ from the C₈ bulk value demonstrates that our surface tension titration is actually probing the surface-pKₐ at the air–aqueous interface. The surface-pKₐ values presented here should be used when discussing surface speciation.

Although the trend observed with chain length is in accordance with accepted trends in the literature, the surface-pKₐ values obtained via our surface tension titration are lower than those which have been reported for the C₈ and C₁₀ acids by Kanicky et al. Throughout the literature, there are many values reported for the pKₐ of both medium- and long-chain fatty acids, although less have been reported for the former. A summary of these results, in addition to those determined here, are represented in Fig. 4.
These values have been determined using a variety of techniques, cover a wide range, and are only sometimes specifically designated as the surface-pK$_a$. Acid–base titration techniques, not including surface tension, were utilized by Kanicky, McLean, and Cistola to determine the pK$_a$ of fatty acids with chain lengths up to eighteen carbons. McLean studied high concentrations of fatty acids to determine "colloidal pK$_a$," and obtained values that are significantly lower than those obtained by Kanicky, particularly at shorter carbon chain lengths. However, a linear trend of increasing pK$_a$ with chain length is consistent between the works.

Other techniques for determining the protonation state of surface-active fatty acids can also be applied to calculate surface-pK$_a$. For example, using X-ray photoelectron spectroscopy, Prisle et al. found the fraction of 0.03 M decanoic acid in a decanoic acid/decanoate system to be only ~10% at a bulk pH of 7.1. When the Henderson–Hasselbalch equation is applied to the published data, a surface-pK$_a$ value of 6.1 is calculated for their C$_{10}$ system, which agrees well with the value reported here. Monolayer relaxation data and surface potential-area isotherms were used by Aveyard et al. and Glazer et al. to calculate the surface-pK$_a$ of stearic acid as 9.3 and 8.0, respectively. Overall, these results suggest inconsistencies in the determination of surface-pK$_a$ between measurement techniques.

The work of Kanicky, McLean, and Prisle also imply that concentration can impact the determination of surface-pK$_a$, as concentrations of the systems vary amongst the experiments and sometimes are not specified in the literature. Additionally, in those experiments in which acid–base titration is used to determine pK$_a$, concentration variations between the studies may affect the activity coefficient of the ionized species in such a way that could potentially account for discrepancies in the reported pK$_a$ values. It has been shown that upon dilution of medium-chain fatty acid systems, and subsequent extrapolation of the data to an extremely dilute limit, the pK$_a$ of these acids approach a value of approximately 5.0. The 1 mM concentrations used in the experiments presented here lie at the solubility limit for the C$_8$–C$_{10}$ acids in water. Therefore, the values reported in Table 1 represent the upper limit to the surface-pK$_a$ of these systems.

Fig. 3 The surface activity model (dashed line along data points) applied to C$_8$ (a), C$_9$ (b), and C$_{10}$ (c) systems. The vertical dashed line is the average surface-pK$_a$ and includes an error bar of ± one standard deviation based on the calculation of surface-pK$_a$ for three separate trials. Surface tension data has been transformed to $\Delta\gamma = \gamma_{\text{max}} - \gamma$.

Table 1 Surface-pK$_a$ of C$_8$–C$_{10}$ fatty acids titrated from pH 12 with 0.10 M HCl

<table>
<thead>
<tr>
<th>C$_n$</th>
<th>Surface-pK$<em>a$ (pK$</em>{a(s)}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C$_8$</td>
<td>4.9 ± 0.2</td>
</tr>
<tr>
<td>C$_9$</td>
<td>5.8 ± 0.1</td>
</tr>
<tr>
<td>C$_{10}$</td>
<td>6.4 ± 0.2</td>
</tr>
</tbody>
</table>

Fig. 4 Survey of surface-pK$_a$ values from the literature. Dashed lines are linear fits of selected data sets and are drawn to guide the eye.
the respective surface-pK_a of the acid systems. From knowledge of the oleic acid systems, we also assume that the cause of the minima in the C_8 and C_9 systems is the presence of RCOOH:RCOO^- complexes at the interface. Interestingly, a dip was not observed for the C_10 system. It is important to note that the data points along the titration curve that are associated with the surface tension dip for the C_8 and C_9 systems were removed prior to fitting the data with eqn (2). This is due to an assumption within the surface activity model that all species, protonated or deprotonated, exist ideally in solution or at the interface with only weak interactions. As it is assumed that this dip is due to the strong interaction of acid-soap species, the deviation from the fit of the data with eqn (2) is further evidence of the formation of RCOOH:RCOO^- complexes near the surface-pK_a.

C_9 titration with 0.10 M NaOH

The titration curves obtained in this work are not the result of measuring several solutions made at discrete pH values. Therefore, the impact of the direction of the titration was considered by titrating a 1 mM C_9 solution at pH 2 with 0.10 M NaOH to approach pH 12. A representative titration of this kind is depicted in Fig. 5. The fully protonated C_9 solution has a surface tension of 31.4 ± 2.0 mN m^{-1} prior to the titration. The similarity between this starting value and the final value around pH 2 of C_9 in Fig. 2 supports the claim that dilution of the C_n solutions over the course of the HCl titration has a minimal impact on the overall results. As the solution becomes more basic during the titration, the surface tension rises as the C_9 species become deprotonated and are removed from the interface. At high pH values, the surface tension approaches a value of approximately 69 mN m^{-1}, in agreement with the starting point of the titration of C_9 with 0.10 M HCl.

Noticeably absent from Fig. 5a is the surface tension dip observed in the C_9 titration curve in Fig. 2. As stated previously, this dip in surface tension is due to the formation of RCOOH:RCOO^- complexes near the surface-pK_a. However, as a pH 2 solution of C_9 is titrated with 0.10 M NaOH, the concentration of Na'^+ ions in the resulting solution is much greater than the concentration of Na^- that is diluted in the titration of a pH 12 solution with 0.10 M HCl. For example, the concentration of Na^- at pH ~ 5 is roughly double in the titration with 0.10 M NaOH compared to that with 0.10 M HCl, respectively 18.5 ± 0.3 mM and 9.1 ± 0.1 mM. The increase in Na^- concentration leads to a disruption in the hydrogen bonding network between protonated and deprotonated fatty acids, similar to a recent study of the effect of sodium on palmitic acid monolayers at varying degrees of deprotonation. This change in bonding environment prevents the complexation, thus eliminating the observed dip.

When eqn (2) is applied to the C_9 titration with 0.10 M NaOH (Fig. 5b), a surface-pK_a value of 5.21 ± 0.02 is determined. When compared to the surface-pK_a obtained using the acidic titrant, the results have a ΔpK_a of ~0.6. The likely cause for such a change in the surface-pK_a is the higher concentration of Na'^+ in solution. While the Gouy–Chapman model was not used to calculate the surface-pK_a of these weakly packed (Fig. 6 and Fig. S5, ESI) Gibbs monolayer systems, the Poisson–Boltzmann description of ion distribution at the interface sheds light on the reduced surface-pK_a in the presence of additional Na'. According to Gouy–Chapman theory, at a given pH, the addition of competing monovalent ions to a system will result in a decreased amount of H'^+ ions accumulating at the interface. Therefore, there will be an increased tendency for the acidic groups at the interface to deprotonate. As there is a greater concentration of Na' in this direction of titration at pH values near the surface-pK_a, the shift of the surface-pK_a to a lower value is to be expected. This effect of ions on surface-pK_a has been observed elsewhere in the literature. In the work of Aveyard et al., the surface-pK_a of stearic acid was found to decrease by 2 pH units upon the addition of 10 mM NaCl (roughly the same amount as the difference between the acidic and basic titrant systems here). By tracking changes in the vibrational modes associated with the hydrogen-bonded carbonyl (C==O) and the deprotonated carboxylate (CO_2^-) headgroup of acid and deprotonated forms, respectively, spectroscopic investigations, utilizing IRRAAS and vibrational sum frequency generation (VSFG), have also revealed that ions (Na'^+, Ca^{2+}, Mg^{2+}, Zn^{2+}, etc.) can affect the protonation state of fatty acid monolayers at...
the air–aqueous interface. This is, again, due to a reduction in the surface-pK\(_a\) as a result of ion affinity for the negative electron density of the carboxylate headgroups at the surface. Therefore, it is important to acknowledge that the concentration of ions in solution will have an effect on the surface-pK\(_a\) determined via surface tension titration and other methods.

### Conclusions

We have used surface tension titration to determine the surface-pK\(_a\) of C\(_{\text{8}}\)–C\(_{\text{10}}\) medium-chain fatty acids at the air–water interface. The surface tension versus pH data was fit to a surface activity model\(^{48}\) which determined surface-pK\(_a\) values of 4.9 for octanoic acid, 5.8 for nonanoic acid, and 6.4 for decanoic acid. Due to the solubility limit of the acids in water, these values represent the upper limit to the surface-pK\(_a\), which decreases upon dilution of the system to the value of the bulk pK\(_a\). The octanoic and nonanoic acid systems exhibited a surface tension dip near the surface-pK\(_a\) due to the formation of acid–soap complexes, which have greater surface activity than the pure acid form. Additionally, it was shown that even small amounts of Na\(^+\) can lower the surface-pK\(_a\) of the C\(_{\text{9}}\) acid system (5.2), and disrupt the bonding network between acid and soap species. This general result is consistent with the ion-induced deprotonation of various fatty acid systems due to the distribution of ions at a charged interface.\(^{61,64–66}\) Upon comparison of the results presented here with those currently in the literature,\(^{3,8,11,49}\) there still exists controversy.

The surface-pK\(_a\) of fatty acids deviates significantly from the bulk value of acetic acid starting at a chain length of nine carbon atoms. Moreover, the surface-pK\(_a\) of these fatty acids are not equivalent to their values in the bulk, as evidenced by the weak acid–strong base titration of nonanoic acid which resulted in a bulk pK\(_a\) value of 4.97, consistent with what has been reported in the literature.\(^{14}\) Therefore, the values of the surface-pK\(_a\) of the SSA-prevalent soluble fatty acids determined here should be utilized for aerosol reactivity and CCN models to more accurately assign the speciation, and surface activity, of organic components at the surface of SSA.\(^{39,67,68}\) Such advances in the understanding of the fundamental properties of these interfacial systems will help to improve certainty in the impacts of aerosols on the climate.

### Conflicts of interest

The authors declare no conflict of interest for this manuscript.

### Appendix A. Surface activity model derivation

If one considers the equilibrium between the protonated (LH) and deprotonated (L\(^-\)) forms of a C\(_n\) acid at the surface,

\[
\text{LH} \leftrightarrow \text{L}^- + \text{H}^+ (\text{aq})
\]

the following expression for the surface equilibrium constant, \(K_{\text{a(s)}}\), is obtained.

\[
K_{\text{a(s)}} = \frac{[\text{L}^-][\text{H}^+]_{\text{s}}}{[\text{LH}]_{\text{s}}}
\]

From the equilibrium, the fraction of protonated (\(f_{\text{LH}}\)) and deprotonated (\(f_{\text{L}^-}\)) species can be written as

\[
f_{\text{LH}} = \frac{[\text{LH}]_{\text{s}}}{[\text{L}^-]_{\text{s}} + [\text{LH}]_{\text{s}}}
\]

\[
f_{\text{L}^-} = \frac{[\text{L}^-]_{\text{s}}}{[\text{L}^-]_{\text{s}} + [\text{LH}]_{\text{s}}}
\]

and rearranged to be expressed in terms of \(K_{\text{a(s)}}\) and [H\(^+\)]\(_s\).

\[
f_{\text{LH}} = \frac{1}{1 + K_{\text{a(s)}}[\text{H}^+]_{\text{s}}^{-1}}
\]

\[
f_{\text{L}^-} = \frac{K_{\text{a(s)}}[\text{H}^+]_{\text{s}}^{-1}}{1 + K_{\text{a(s)}}[\text{H}^+]_{\text{s}}^{-1}}
\]

Or, using the definitions of pH and pK\(_{\text{a(s)}}\), eqn (A4a) and (A4b) can be rewritten as

\[
f_{\text{LH}} = \frac{1}{1 + 10^{\text{pH} - pK_{\text{a(s)}}}}
\]

\[
f_{\text{L}^-} = \frac{10^{\text{pH} - pK_{\text{a(s)}}}}{1 + 10^{\text{pH} - pK_{\text{a(s)}}}}
\]

It is important to note that in order to apply Cratin’s surface activity model\(^{48}\) (originally developed to describe stearic acid dissociation at the oil–water interface where a completely dissociated acid would result in an interfacial tension of ~0 mN m\(^{-1}\)) to these systems, a simple difference must be taken within the data such that at high pH, the surface tension is associated with a value of 0 mN m\(^{-1}\). As such, the surface tension data collected during the titration is presented as \(\Delta \gamma\) against pH, where \(\Delta \gamma = \gamma_{\text{max}} - \gamma\), and \(\gamma_{\text{max}}\) is the surface tension of the C\(_n\) system at a pH of 12. The surface activity (\(a\)) for these medium-chain fatty acids at the air–water interface is then expressed in terms of the lowering of this surface tension difference relative to a maximum value (\(\Delta \gamma_{\text{max}}\) obtained at low pH).

\[
a = \frac{(\Delta \gamma_{\text{max}} - \Delta \gamma)}{\Delta \gamma_{\text{max}}}
\]

One of the bounds of this model is that all species, protonated or deprotonated, must coexist independently with only weak interactions in order to express their total activity as a simple summation. Under this assumption, the total activity (\(a_t\)) of the system can be expressed as the sum of the activity and fraction of the protonated and deprotonated species.

\[
a_t = \sum_i a_i f_i = a_{\text{L}^-} f_{\text{L}^-} + a_{\text{LH}} f_{\text{LH}}
\]
Equating eqn (A6) and (A7) yields
\[
\frac{(\Delta \gamma_{\text{max}} - \Delta \gamma)}{\Delta \gamma_{\text{max}}} = a_L f_L - a_{LH} f_{LH} \tag{A8}
\]
From examination of a \(\Delta \gamma\) vs. pH plot for these systems, one can deduce that \(a_{LH} = 0\), and \(a_L = 1\) (see ESI\(^\dagger\)). From this, eqn (A8) can be written as
\[
\frac{(\Delta \gamma_{\text{max}} - \Delta \gamma)}{\Delta \gamma_{\text{max}}} = f_L \tag{A9}
\]
which can be arranged and expressed as eqn (A10).
\[
\Delta \gamma = \Delta \gamma_{\text{max}} (1 - f_L) \tag{A10}
\]
Inserting the equation for \(f_L\) from eqn (A5\(b\)), the above equation can be written in terms of the surface pH (pH\(_S\)) and \(pK_a(s)\) of the system.
\[
\Delta \gamma = \Delta \gamma_{\text{max}} \left(1 - \frac{10^{(pH_s - pK_{a(s)})}}{1 + 10^{(pH_S - pK_{a(s)})}}\right) \tag{A11}
\]
Finally, eqn (A11) can be simplified to yield
\[
\Delta \gamma = \frac{\Delta \gamma_{\text{max}}}{1 + 10^{(pH_S - pK_{a(s)})}} \tag{A12}
\]
If one assumes that pH\(_S\) is equal to the bulk pH (pH\(_b\)), eqn (A12) becomes
\[
\Delta \gamma = \frac{\Delta \gamma_{\text{max}}}{1 + 10^{(pH_b - pK_{a(s)})}}
\]
While charged monolayers at the interface can create an electric double layer with hydrogen ion concentrations enhanced relative to the bulk, a 1 mM C\(_9\) solution, for example, represents a very loosely packed sub-monolayer system. Fig. 6 shows IRRAS spectra of 1 mM C\(_9\) solutions at pH 2, 12, and of samples extracted from the titration experiment at the surface tension dip. No CH stretching modes are observed at pH 12 indicating that the C\(_9\) molecules are not found above the detection limit at the interface. CH stretching modes are present for the 1 mM C\(_9\) solution at pH 2 and in the extracted dip region sample. However, the intensities of the modes are very weak, and the CH\(_2\) stretching frequencies are blue shifted compared to more ordered and surface active monolayer of stearic acid (Fig. S5, ESI\(^\dagger\)), indicating that the C\(_9\) system is loosely packed and exhibits significant gauche defects. Therefore, the assumption that pH\(_S\) is approximately equal to pH\(_b\) holds for the calculation of the surface-pK\(_a\) of the medium-chain fatty acids.\(^{63}\)

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**References**


