2013

Rapid differentiation of isomeric lipids by photodissociation mass spectrometry of fatty acid derivatives

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Publication Details
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Abstract
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METHODS Herein we describe the derivatization of a structurally diverse suite of fatty acids as 4-iodobenzyl esters (FAIBE). Electrospray ionization of these derivatives in the presence of sodium acetate yields abundant \([M + Na]^+\) ions that can be mass-selected and subjected to laser irradiation \((\lambda = 266 \text{ nm})\) on a modified linear ion-trap mass spectrometer.

RESULTS Photodissociation (PD) of the FAIBE derivatives yields abundant radical cations by loss of atomic iodine and in several cases selective dissociation of activated carbon–carbon bonds (e.g., at allylic positions) are also observed. Subsequent CID of the \([M + Na – I]^+\) radical cations yields radical-directed dissociation (RDD) mass spectra that reveal extensive carbon–carbon bond dissociation without scrambling of molecular information.

CONCLUSIONS Both PD and RDD spectra obtained from derivatized fatty acids provide a wealth of structural information including the position(s) of unsaturation, chain-branching and hydroxylation. The structural information obtained by this approach, in particular the ability to rapidly differentiate isomeric lipids, represents a useful addition to the lipidomics tool box.

Keywords
differentiation, isomeric, lipids, photodissociation, rapid, derivatives, mass, acid, spectrometry, fatty, GeoQuest, CMMB

Disciplines
Medicine and Health Sciences | Social and Behavioral Sciences

Publication Details

This journal article is available at Research Online: http://ro.uow.edu.au/smhpapers/512
Rapid differentiation of isomeric lipids by photodissociation mass spectrometry of fatty acid derivatives

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Keywords: lipids, lipidomics, mass spectrometry, photodissociation, ion activation, radical ions

Running Heading: Photodissociation of derivatized fatty acids
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Methods: Herein we describe the derivatization of a structurally diverse suite of fatty acids as 4-iodobenzyl esters (FAIBE). Electrospray ionization of these derivatives in the presence of sodium acetate yields abundant [M + Na]+ that can be mass-selected and subjected to laser irradiation (λ = 266 nm) on a modified linear ion-trap mass spectrometer.

Results: Photodissociation (PD) of the FAIBE derivatives yields abundant radical cations by loss of atomic iodine and in several cases selective dissociation of activated carbon-carbon bonds (e.g., allylic positions) are also observed. Subsequent collision-induced dissociation of the [M + Na - I]+ radical cations yields radical-directed dissociation (RDD) mass spectra that reveal extensive carbon-carbon bond dissociation without scrambling of molecular information.

Conclusions: Both PD and RDD spectra obtained from derivatized fatty acids provide a wealth of structural information including the position(s) of unsaturation, chain-branching and hydroxylation. The structural information obtained by this approach, in particular the ability to rapidly differentiate isomeric lipids, represents a useful addition to the lipidomics tool-box.
Introduction

Traditional mass spectrometric analysis of fatty acids (FA) involves: (i) a chemical derivatization step to enhance the volatilization of these lipids; (ii) separation by gas-chromatography; and (iii) ionization, most commonly by electron ionization (EI). By far the most widely used method of chemical modification is to convert free fatty acids to fatty acid methyl esters (FAMEs), which have desirable chromatographic properties.\textsuperscript{1,2} Observation of the molecular ion in a conventional EI (70 eV) spectrum of a FAME allows the assignment of carbon-chain length and degree of unsaturation, complementing information derived from GC retention time. Unfortunately, the molecular ions arising from EI of FAMEs are typically low in abundance with much of the signal intensity found in peaks corresponding to dissociation and/or rearrangements of the molecular framework, \textit{e.g.}, a base peak at \textit{m/z} 74 is common and arises from the McLafferty rearrangement.\textsuperscript{1,3} Furthermore, even when the stoichiometry of the lipid can be determined from the EI mass spectrum, distinguishing it from possible isomeric variants is difficult due to scrambling of molecular information driven by the high energy of the EI process. For example, EI cannot distinguish unsaturated FAME isomers due to the migration of double bonds prior to dissociation of the molecular ion.\textsuperscript{3} Although such effects can be modulated in some instances by reducing the EI energy,\textsuperscript{4} in general, distinguishing between isomeric FAMEs relies on their separation by GC and comparison of retention times with authentic compounds. This too presents challenges where appropriate reference compounds are not available or chromatographic resolution is poor resulting in overlapped peaks.\textsuperscript{5,6} The inability to assign fatty acid structures unambiguously or even the possibility of misassignments is thus a significant limitation of current technologies. In particular, such limitations mask the full extent of natural variation in lipid structure (\textit{e.g.}, the positions of branched motifs or site(s) of unsaturation) and thus, the distinct biochemical and biophysical roles of individual lipid species are more difficult to uncover.\textsuperscript{7,8,9}
As a result these aforementioned limitations in EI analysis of FAMEs, other derivatization strategies have emerged to enhance the structure-specificity of fatty acid analysis by GC-MS. Notable among these are the use of picolinyl esters and 4,4-dimethyloxazoline (DMOX) derivatives. These modifications introduce a nitrogen heterocycle that when subjected to conventional EI can effectively sequester the charge thus enhancing the abundance of the molecular ion and facilitating the formation of more structurally diagnostic product ions. For example, the observation of a 12 Th (or in some cases 26 Th) peak spacing in EI spectra of picolinyl ester or DMOX derivatives can be used to assign carbon-carbon double bond position(s) in unsaturated fatty acids. Although this represents a significant improvement over FAME derivatives, there are still limitations in this approach. For example, EI of DMOX derivatives do not yield diagnostic details of methyl branching points within saturated acyl chains. Branched lipids, especially methyl branched fatty acids, are widespread in nature where they are thought to have essential roles in metabolism. Thus mass spectrometric methods capable of also revealing lipid isomerism arising from chain-branching are desirable. One approach, pioneered by Zirrolli and Murphy, involved mass-selection of the low abundant molecular ions arising from EI of FAMEs and subjecting these to low energy collision-induced dissociation (CID) on a tandem mass spectrometer. The resulting EI-CID mass spectra successfully differentiated non-branched and branched acyl chains. In the case of the straight-chain variants, almost identical product ion abundances arising from the cleavage of every carbon-carbon bond were observed in the EI-CID spectrum and resulted in regular peak spacing of 14 Th. In contrast, for the branched-chain species, carbon-carbon bond cleavage was enhanced on either side of the tertiary carbons leading to a characteristic gap of 28 Th that allowed the position(s) of methyl substitution to be assigned. The mechanism of EI-CID fragmentation was studied by isotope-labelling experiments which revealed that radical-driven fragmentation was the most probably responsible for these spectral patterns. Recently, EI-CID was extended by Brenna and co-
workers\textsuperscript{18} to examine a series of saturated branched-chain FAME synthetic standards. They were able to establish an exhaustive look-up table of diagnostic product ions for determining the methyl branching points. It was noted however, that the sensitivity in detecting these features reduces as the position of methyl-substitution approaches the ester moiety. Notably therefore, the predicted peak assigning a branch point at the C3-position is often absent in the EI-CID spectrum. Although powerful, one disadvantage of EI-CID is the inherently low abundance of the required molecular ions such that only a very small number of ionized molecules retain the necessary structural information for tandem MS analysis.

Due to recent advances in fast and efficient liquid chromatography (LC) protocols, LC-MS is gaining wider acceptance as an alternative approach to the identification and quantification of fatty acids in biological extracts.\textsuperscript{19,20} Location of double bonds within a particular acyl chain is found to significantly affect LC providing desirable chromatographic resolution of isomers – particularly in the presence of silver ions\textsuperscript{21,22} – however, the unambiguous assignment of this structural feature still relies on retention times and comparison with standard compounds (\textit{cf.} GC discussion above). Therefore ion activation methods that produce diagnostic mass spectra and are also compatible with LC protocols are of increasing interest. Given the presence of the carboxylate moiety, [M - H]\textsuperscript{-} ions generated by electrospray ionization (ESI) are common targets for fatty acid identification in LC-MS protocols. Conventional (low energy) CID mass spectra of these ions are typically dominated by loss of carbon dioxide and thus do not reveal details of the structure of the hydrocarbon chain although recent work suggests the relative abundance of these ions can be a useful probe.\textsuperscript{23} Hsu and Turk\textsuperscript{24} have shown that the addition of lithium salts can give rise to abundant [M - H + Li\textsubscript{2}]\textsuperscript{+} ions that upon CID undergo carbon-carbon bond cleavage at vinylic and allylic positions that can be exploited in localizing double bond positions. This, and related approaches, have been successfully deployed for the LC-MS identification of unusual fatty acids in complex mixtures.\textsuperscript{25} An alternative approach to promote even-electron
fragmentation at carbon-carbon double bonds is to undertake chemical modification of the motif itself. For example, ozonolysis of lipids in a thin film or pre-treatment of extracts with osmium tetroxide (OsO₄) modify the targeted functional group by oxidation making it susceptible to cleavage upon CID. Such chemical derivatizations do increase the complexity of the mixture and may cause significant and undesirable changes in LC behaviors depending on how many sites are modified. Alternative methods exploit selective chemistries in the gas phase, such as ozone-induced dissociation and covalent adduct chemical ionization. These are powerful methods for the identification of carbon-carbon double bond positions in unsaturated lipids and both are compatible with LC-MS workflows but are unable to obtain information on other structural features in the lipid such as position(s) of chain branching.

Conceptually it is attractive to exploit radical-driven dissociation (cf. EI-CID) but in combination with high ion yields, well-defined dissociation energetics and in a manner compatible with LC-MS. In this vein we have recently introduced radical-directed dissociation (RDD) to the structural analysis of complex lipids. In this approach even-electron complexes are formed between the target lipids and a suitable radical initiator during ESI. Mass-selection of the complex and subsequent laser irradiation gives rise to odd-electron ions, which upon further activation by low energy CID results in extensive dissociation of the carbon-carbon bonding framework. RDD is thus able to unambiguously identify double-bond position(s) as well as differentiate between isomeric branched and non-branched acyl chains within complex lipids. As we will show however, the non-covalent attachment of radical initiators to lipids is not applicable to the structural analysis of simple lipids such as isomeric fatty acids. Here we describe an alternative strategy tailored to fatty acid analysis that exploits a standard chemical derivatization procedure to covalently attach a chromophore to the lipid that incorporates a UV-labile carbon-iodine bond. Examination of a suite of so-modified fatty acids by both PD and RDD reveals highly sensitive and selective fragmentation that can
successfully differentiate isomeric fatty acids differing only in location of carbon-carbon double bond(s) or branching point(s).

Methods

Chemicals and reagents

HPLC grade methanol, chloroform, pentane and acetonitrile solvents were obtained from Ajax Finechem (Sydney, NSW, Australia) and were used without further purification. The following fatty acid standards (~99% purity) were purchased from Nu-Chek Prep (Elysian, Minnesota, USA): linoleic acid FA (9Z,12Z-18:2); petroselinic acid FA (6Z-18:1); petroselaidic acid FA (6E-18:1); oleic acid FA (9Z-18:1); elaidic acid FA (9E-18:1); cis-vaccenic acid FA (11Z-18:1); ricinoleic acid 12-OHFA (9Z-18:1); and 12-hydroxy stearic acid 12-OHFA (18:0). Two saturated (non-branched and branched chains) fatty acid isomers were purchased from Sigma-Aldrich (Castle Hill, Australia): arachidic acid (99%) FA (20:0) and phytanic acid (96%) FA (4Me16:0). Other reagents for derivatization including boron trifluoride (BF₃) 10% in methanol solution, 4-iodoaniline (98%), 4-iodobenzyl alcohol (97%) and sodium acetate and sulphuric acid (98%) were also purchased from Sigma-Aldrich (Castle Hill, Australia).

In this manuscript, it is sometimes instructive to indicate the double bond position within a fatty acid or derivative without specific reference to the stereochemistry. So here we adopt the commonly used Δⁿ nomenclature indicating that the double bond is located at the \(^n\)th carbon-carbon bond counting from the carboxylate moiety, \(^m\) e.g., the carbon-carbon double bond in oleic acid is Δ⁹.

Sample preparation

A chemical reaction between a fatty acid and 4-iodobenzyl alcohol is initiated in the presence of a 2% sulphuric acid catalyst (Scheme 1). The esterification procedure was
conducted as follows: approximately 1 mg of fatty acid and 5 mg of 4-iodobenzyl alcohol were dissolved in 1 mL acetonitrile in a 3 mL glass vial. Concentrated sulphuric acid (20 µL) was added to the mixture to obtain a final concentration of 2% (v/v) in solution, followed by heating at 80°C for 30-45 minutes in a water bath. After cooling, water (0.5 mL) and pentane (1 mL) were sequentially added to the solution. The reaction mixture was shaken several times to separate the aqueous and organic components: the desired ester is partitioned into the non-polar organic layer. The upper phase (i.e., pentane) was then collected with a fatty acid 4-iodobenzyl ester (FAIBE) concentration of approximately 3 mM. Prior to ESI-MS experiments, samples were prepared by diluting the pentane extracts to 10-20 µM of fatty acid iodobenzyl derivatives in methanol. Sodium (or lithium) acetate was added to the sample solutions to a final concentration of 50 µM. FAIBE yields achieved under these conditions were estimated to be > 90% with the exception of the hydroxyl fatty acids where yields were ca. 10-50%.

**Scheme 1:** Derivatization of fatty acids to produce fatty acid 4-iodobenzyl esters (FAIBE).

Fatty acid methyl ester derivatives of arachidic acid FA (20:0) and phytanic acid FA (4Me16:0) were prepared by treatment with 10% BF₃ in methanol. The samples used in preparation of non-covalent adducts between FAMEs and 4-iodoaniline were obtained by extracting FAME (20:0) and FAME (4Me16:0) into pentane and diluting this organic layer to a final concentration of 10-20 µM of FAMEs in methanol, then adding 4-iodoaniline 10 µM and 0.2% formic acid.
Instrumentation

All mass spectra were acquired on a modified LTQ linear ion trap mass spectrometer (Thermo Fisher Scientific, San Jose, CA). The sample solutions of fatty acid derivatives were introduced into the electrospray ionization source by direct infusion to generate the gaseous sodium adducts. Typical source parameters were: sample flow rate 3 µL min⁻¹, spray voltage +4.5 kV, capillary temperature 250°C, tube lens voltage 129 V, and the capillary voltage 49 V. Nitrogen gas served as the sheath (arbitrary flow units between 5 and 20), auxiliary and sweep gases (between 0 and 5), and helium gas served as the buffer gas. Ions were mass-selected with a window of 1-3 Th and subjected to photodissociation (PD) as described below. Radical-directed dissociation (RDD) mass spectra were acquired by subsequent mass-selection of radical ion photoproducts followed by collision-induced dissociation using standard conditions (i.e., manufacturers parameters of normalized collision energy and activation time are set to 20-25 % and 30 ms, respectively). All spectra presented represent an average of 50-100 scans.

The linear ion trap was previously modified³¹ to enable PD experiments following the experimental configuration described by Ly and Julian.³² Briefly, a quartz window (MDC Vacuum Products, Hayward, CA) was installed on the posterior plate of the vacuum housing to allow transmission of 266 nm laser pulses (~30 mJ cm⁻²) from a flash lamp-pumped Nd:YAG laser (Continuum, Santa Clara, CA). All laser experiments herein are conducted at λ = 266 nm and the pulse-width of the laser is approximately 5 ns. The laser beam was directed into the trap via two right-angle prisms, which can be adjusted to optimize alignment with the ion cloud. Pulses were synchronized to the beginning of the activation step of a typical MSⁿ experiment by feeding a TTL pulse from the instrument to the laser via a digital delay generator (Berkeley Nucleonics, San Rafael, CA). Only one laser pulse irradiates the selected ions per mass spectral cycle.
Results and Discussion

Saturated straight-chain and branched fatty acids

Figure 1 shows photodissociation (PD) mass spectra of protonated 4-iodoaniline (IA) at m/z 220 and the two non-covalent complexes of this adducting agent with two saturated methyl esters, forming [FAME (20:0) + IA]⁺ and [FAME (4Me16:0) + IA]⁺ precursor ions both seen at m/z 546.

**Figure 1:** PD mass spectra of (a) protonated 4-iodoaniline, and non-covalent complexes between protonated 4-iodoaniline with FAMEs of (b) phytanic acid, FAME (4Me16:0) and (c) arachidic acid, FAME (20:0). Structures of the precursor ions are indicated above each spectrum.
All three PD spectra in Figure 1 show only one major product ion at $m/z$ 93 corresponding to an aniline radical cation and significantly, in the case of the lipid adduct ions, no radical ion incorporating the lipid is observed. The formation, and subsequent collisional activation of lipid radical ions derived from PD of such non-covalent complexes was central to the radical-directed dissociation (RDD) strategy we have previously employed for complex lipids.\textsuperscript{29} The data shown here, however, indicate that at least in the case of saturated FAMEs, our non-covalent RDD approach is unable to provide any structurally diagnostic fragmentation let alone discrimination between branched and straight chain fatty acid isomers.

In contrast to the non-covalent complexes described above, the PD mass spectra of covalent 4-iodobenzyl ester derivatives of the same straight-chain and branched fatty acid isomers show rich fragmentation and significant points of differentiation (Figure 2a and c, respectively). These spectra were obtained by isolating $m/z$ 551 precursor ions for [FAIBE (20:0) + Na]$^+$ and [FAIBE (4Me16:0) + Na]$^+$ and subjecting them to a single laser pulse ($\lambda = 266$ nm). For both isomers the major ionic photoproduct is observed at $m/z$ 424 and corresponds to homolysis of the carbon-iodine bond and formation of a $[M + Na - I]^+$ radical cation. Subsequent mass-selection and collision-induced dissociation of this ion in each case gave rise to the RDD mass spectra shown in Figures 2(b) and (d). Similar PD and RDD spectra were obtained for the analogous $[M + Li]^+$ ions (Supporting Information, Figure S1) suggesting that dissociation is predominantly radical-directed rather than charge-directed in these experiments. Proposed radical-directed fragmentation pathways for the branched acyl chain are shown in Scheme 2.
Figure 2: Mass spectra of saturated non-branched vs. branched fatty acid 4-iodobenzyl ester (FAIBE) derivatives. (a) and (b) PD and RDD mass spectra acquired from arachidic acid, FAIBE (20:0); (c) and (d) PD and RDD spectra acquired from phytanic acid FAIBE (4Me16:0).
Scheme 2: Proposed fragmentation pathways for FAIBE (4Me16:0) where the $m/z$ of product ions correspond to those shown in Figure 2(d).
Comparing the PD spectra shown in Figures 2(a) and (c) reveals a difference in peak spacing in the low-mass region of the spectra. Specifically, product ions at m/z 172 and 185 ($\Delta m = 13$ Da) are observed for the straight-chain FAIBE (20:0), while ions at m/z 172 and 199 ($\Delta m = 27$ Da) are associated with methyl-branched chain of FAIBE (4Me16:0). Mechanisms to account for these fragments in the latter case are indicated in Scheme 2 (pathways vii, viii). In this proposal, hydrogen atom abstraction by the nascent phenyl radical results in relocation of the radical to the C2- and C4-positions on the fatty acyl chain. Subsequent $\beta$-scission from these positions can yield the radical cation at m/z 172 or the closed shell olefin at m/z 199. Applying the same mechanisms to arachidic acid (not shown) yields the identical product ion at m/z 172 but, in the absence of the C3-methyl branch, $\beta$-scission from C2 yields a product ion at m/z 185. These features of the PD spectra alone are sufficient to differentiate the two isomers and further spectral differences are also noted among the lower abundance product ions. These low intensity signals are enhanced by subsequent collisional activation and the resulting RDD spectra are discussed below.

The regular peak spacing observed in the RDD spectrum of the straight-chained arachidic acid in Figure 2(b) suggest that hydrogen abstraction by the phenyl radical occurs almost uniformly at all methylene positions along the carbon chain. Subsequent $\beta$-scission processes from these carbon-centred radicals give rise to a homologous series of radical cations (resulting from neutral alkene elimination) and even-electron olefinic cations (resulting from alkyl radical losses). The superposition of these processes that differ in product ion mass by 1 Da results in the apparent peak doubling observed in Figure 2(b), for example, m/z 297 and 298 ions result from loss of C$_9$H$_{19}^*$ and alkene C$_9$H$_{18}$, respectively. The presence of methyl branching however, leads to more selective fragmentation patterns in the RDD of FAIBE (4Me16:0) (Figure 2d) where product ion abundance is found to vary more substantially. The preference for chain cleavage at some positions over others is attributed to the scenarios where either (i) a stabilized tertiary radical located at a branch point undergoes
dissociation via $\beta$-scission (e.g., Scheme 2 pathway iii) or (ii) dissociation adjacent to a branch point yields a secondary radical product (e.g., Scheme 2 pathway ii). This rationale is illustrated for all major product ions in Scheme 2 and explains the characteristic alternation between even and odd mass product ions observed in Figure 2(d). It is also noted that the peak spacing observed at each methyl branching point is not always the same. Particularly, $\Delta m = 27$ Da between $m/z$ 199 and 172 is diagnostic for the methyl branch at C3, closest to carbonyl group (*vide supra*), while $\Delta m = 28$ Da between $m/z$ 409 and 381 is observed for the iso-branching point adjacent to the methyl terminus. For other branching points in the middle of the carbon chain, the peak spacing is $\Delta m = 29$ Da at each position (e.g., $m/z$ 340 and 311, and $m/z$ 270 and 241). If found to be consistent in other branched fatty acids, such peak spacings in RDD spectra may prove useful in the differentiation of iso- and anteiso-branched chain FA.\textsuperscript{18} Importantly, the radical dissociation processes in the RDD spectra of the FAIBE derivatives clearly differentiate straight-chain FA (20:0) from its methyl-branched FA (4Me16:0) isomer (*cf.* Figure 2b and d). Moreover in the latter case, the fragmentation patterns can be used to localize the positions of methyl substituents.

**Monounsaturated fatty acids**

Three isomeric monounsaturated fatty acids, namely FA (11Z-18:1), FA (9Z-18:1) and FA (6Z-18:1) were derivatized as 4-iodobenzyl esters and subjected to electrospray ionization in the presence of sodium acetate. The resulting [M + Na]$^+$ ions were then irradiated by a single laser pulse ($\lambda = 266$ nm) and the resulting PD mass spectra are shown in Figure 3. While in all three instances the most abundant product ions are assigned as the loss of atomic iodine (*i.e.*, [M + Na - I]$^+$ at $m/z$ 394) the next most abundant product ions are distinct in each case (*i.e.*, $m/z$ 324, 296 and 254 in Figures 3a-c, respectively). Although some magnification is required to visualise these product ions, the PD spectra display excellent signal-to-noise and thus these diagnostic signals can readily be used to differentiate between the three isomers.
Furthermore, the neutral losses of the radicals $\text{C}_5\text{H}_{11}^\cdot$, $\text{C}_7\text{H}_{15}^\cdot$ and $\text{C}_{10}\text{H}_{21}^\cdot$ in the spectra shown in Figures 3(a-c), respectively are consistent with homolytic cleavage of the allylic carbon-carbon bond implicating a common mechanism in each case.

**Figure 3:** Photodissociation (PD) mass spectra acquired from monounsaturated isomers [FAIBE (18:1) + Na$^+$] derivatized from (a) FA (11Z-18:1), (b) FA (9Z-18:1) and (c) FA (6Z-18:1). The symbol * indicates the diagnostic product ion formed selectively from each double bond positional isomer.
Energetically, the initial phenyl radical formed following photolysis of the carbon-iodine bond can abstract any hydrogen atom from the carbon chain due to the higher carbon-hydrogen bond dissociation energy of benzene (~113 kcal mol\(^{-1}\)) compared to aliphatic carbon-hydrogen bond energies (~96 kcal mol\(^{-1}\)).\(^{33}\) As a result, facile hydrogen atom transfer from a range of sites on the fatty acyl chain is expected and several representative structures resulting from this initial rearrangement are indicated in Scheme 3. In some instances the radical will be at a stabilized position such as the \(\alpha\)-position (adjacent to the carbonyl moiety) or the allylic positions (adjacent to the carbon-carbon double bond), examples of which are indicated in Schemes 3(b) and (d), respectively. Even in instances where the radical does not transfer directly to these positions, subsequent hydrogen atom transfer events will also be thermodynamically favorable leading to an eventual cascade to stabilized positions. Indeed, Ly and Julian have estimated that in RDD of peptides and proteins up to three such sequential hydrogen atom transfers may occur until the radical becomes localized at energetically favored positions.\(^{34}\) As a consequence of these processes, the bulk of the ion population that survives the initial photolysis event is expected to be made up of stabilized radical cations and it is these species which will be isolated for subsequent CID in RDD experiments (see later).

In contrast, if the initial hydrogen atom abstraction occurs adjacent to an activated bond then direct \(\beta\)-scission results. Two such examples are illustrated for FAIBE \((9Z-18:1)\) in Scheme 3(a) and (c) and correspond to the formation of resonance stabilized ester enolate \((m/z 172)\) and allyl \((m/z 296)\) radical cations, respectively. The selectivity of the latter pathway makes this a particularly useful diagnostic for double bond position and thus provides an alternative means of selective identification and differentiation of unsaturated fatty acids.
Scheme 3: Proposed mechanism radical rearrangement and dissociation resulting from PD of 4-iodobenzyl ester derivative of oleic acid. Note that only representative examples of isomerisation and/or dissociation pathways of the radical cation are shown.

Subsequent CID performed on the abundant radical ions at m/z 394 from each of the monounsaturated FAIBE isomers gave rise to rich RDD fragmentation as illustrated in Figure 4. At a glance, the base peaks observed in all three RDD spectra are distinct (i.e., m/z 337, 309 and 239 in Figure 4a-c, respectively) thus representing a characteristic ion for each isomer. In Figure 4(a), the base peak at m/z 337 corresponds to a neutral loss of a C₄H₉• radical (-57 Da), which likely involves β-scission from an allyl radical adjacent the Δ₁¹ double bond (Scheme 4a). By extension of this mechanism, when the position of unsaturation is at Δ⁹, as for FAIBE (9Z-18:1), a 28 Th shift of the most abundant RDD fragment is predicted. This corresponds to a C₆H₁₃• loss (Scheme 4b), and indeed the ion produced via this mechanism is observed at m/z 309 in Figure 4(b). Similarly, the same process occurring in FAIBE (6Z-18:1) would give rise to a product ion at m/z 267 from loss of a C₉H₁₉• radical. While this fragment is observed in the RDD spectrum, it is of low abundance and the spectrum shown in Figure 4(c) is instead
dominated by the product ion at \( m/z \) 239 suggesting a competing pathway exists for this isomer that results in loss of a \( \text{C}_{11}\text{H}_{23} \cdot \) radical. One possible explanation for the preferential formation of the \( m/z \) 239 signal for this isomer is the process outlined in Scheme 4(c). In this mechanism, activation of the stabilized ester enolate radical results in cyclization and addition to the \( \Delta^6 \)-double bond to form a six-membered ring. Subsequent \( \beta \)-scission from this intermediate could give rise to the observed \( \text{C}_{11}\text{H}_{23} \cdot \) radical loss. The formation of a six-membered ring for FAIBE (6Z-18:1) is expected to be favorable, while the same process will become disfavored for double bond positions beyond \( \Delta^6 \) due to entropic constraints.

In contrast to the formation of a single dominant product ion, as observed for FAIBE (6Z-18:1) (Figure 4c), the RDD spectra of FAIBE (9Z-18:1) and FAIBE (11Z-18:1) isomers (Figure 4a and c) reveal extensive fragmentation of the hydrocarbon chain. The product ions observed correspond to dissociation arising from almost all positions suggesting that dissociation is preceded by radical migration. By analogy to saturated fatty acids discussed earlier (cf. Scheme 2), these product ions can be rationalized as resulting from a series of alkyl radical or alkene losses. The regular peak spacing of 14 Th that arises from these processes, is interrupted by the carbon-carbon double bond providing a distinctive 12 Th spacing as indicated by the symbols \( \blacklozenge \) in Figure 4, e.g., \( m/z \) 309 and \( m/z \) 297 in Figure 4(a). The characteristic 12 Th peak spacing suggests that RDD spectra of FAIBE derivatives could be used to locate position(s) of unsaturation within unknown lipids. Interestingly, this peak spacing pattern is consistent with that reported for unsaturated fatty acid derivatives upon \( \text{EI}^{12,13} \) and high energy CID.\(^{35,36} \)
Figure 4: RDD spectra resulting from the PD-generated \([M + Na - I]^+\) radical cation \(m/z\) 394 from 4-iodobenzyl ester derivatives: (a) FAIBE (11Z-18:1); (b) FAIBE (9Z-18:1); and (c) FAIBE (6Z-18:1). Product ions with a spacing of 12 Th indicative of carbon-carbon double bond position are labelled with symbols (◆).
Scheme 4: Proposed mechanism of RDD fragmentation observed from (a) FAIBE (11Z-18:1); (b) FAIBE (9Z-18:1); and (c) FAIBE (6Z-18:1).

The sensitivity of both PD and RDD processes to the configuration of the carbon-carbon double bond was also investigated. FAIBE derivatives of the elaidic acid FA (9E-18:1) and petroselaidic FA (6E-18:1) acids were prepared, both of which have trans stereochemistry about their double bonds. The PD and RDD spectra obtained from the sodium adduct ions of these derivatives are provided as supporting information (Figures S2 and S3, respectively) and are indistinguishable from those acquired from their cis counterparts, oleic and petroselinic shown in Figures 3 and 4. While the inability to differentiate stereoisomers by this approach was disappointing it was not altogether unexpected given that the thermodynamic preference for hydrogen atom abstraction from the allylic position(s) in
unsaturated lipids is largely independent of stereochemistry. Indeed, the formation of allylic radicals such as those indicated in Schemes 3 and 4 result in a loss of stereochemical information.

Polyunsaturated fatty acid

In order to investigate the photo-fragmentation behavior of polyunsaturated fatty acids, 4-iodobenzyl ester derivatives of linoleic acid were prepared and subjected to both PD and RDD (Figure 5a and b, respectively). PD of the mass-selected [M + Na]$^+$ ion of FAIBE (9Z,12Z-18:2) at m/z 519 gives rise to predominantly the [M + Na - I]$^+$ ion at m/z 392 (Figure 5a). Several lower abundant photo-fragments are also observed in this spectrum. The product ions at m/z 336 and 296 correspond to neutral loss of atomic iodine followed by losses of C$_4$H$_8$ and C$_7$H$_{12}$ alkenes, respectively. By analogy with the mono-unsaturated systems discussed above, these fragments can be rationalized as resulting from dissociation of activated carbon-carbon bonds and formation of a radical cation incorporating the stabilized allyl radical moiety (cf. Scheme 3c). In contrast, the product ion at m/z 322, corresponding to the neutral loss of a C$_5$H$_{10}$ alkene with cleavage at the vinylic position appears to be characteristic of the homoallylic diene motif. The RDD spectrum shown in Figure 5(b) was acquired from CID of the PD-generated radical ions at m/z 392. The fragmentation pattern of the polyunsaturated fatty acid is clearly distinguishable from those of the monounsaturates previously discussed. Notably, the product ion spacing of 12 Th (labelled with ♦ in Figure 5b) is repeated twice in the RDD spectrum of the linoleic acid derivative. The first of these is noted between m/z 321 and 309, corresponding to C$_5$H$_{11}$• and C$_6$H$_{11}$• neutral alkyl and alkenyl radical losses resulting from vinylic and allylic carbon-carbon bond cleavages and indicative of the $\Delta^{12}$ double bond. Similarly, the second distinctive ion pair observed at m/z 281 and 269 coincides with the location of the second $\Delta^9$ double bond. An interesting “double peak” feature is also observed in Figure 5(b) with abundant ions at both m/z 309 and 307 suggestive
of competing pathways for cleavage at the $\Delta^{12}$-double bond that result in neutral losses of unsaturated ($C_6H_{11^\cdot}$) and saturated radicals ($C_6H_{13^\cdot}$), respectively. The approximately equal abundance of these two product ions appears to be a characteristic feature of the 18:2 acyl chain as the equivalent ions are of low or negligible abundance for both the monounsaturated 18:1 analogues (e.g., note the low abundance of $m/z$ 295 relative to $m/z$ 297 in Figure 4a) and indeed, the equivalent C9 losses from 18:2 (i.e., no $m/z$ 267 partners the $m/z$ 269 in Figure 5b) While the mechanistic origins of this “double peak” remain to be demonstrated, there is some analogy to our prior observations in the RDD spectra of complex lipids. Overall, these data confirm that the key structural features of the polyunsaturated fatty acid are retained in the radical ion formed upon photodissociation of the 4-iodobenzyl ester derivative. Moreover, the subsequent dissociation of this species gives rise to a fragmentation pattern from which the initial structure can be gleaned.
Figure 5: (a) The PD spectrum of the \([M + Na]^+\) ion of FAIBE (9Z,12Z-18:2) linoleic acid. (b) The RDD spectrum acquired by collisional activation of \(m/z\) 392 radical ion formed in (a). Adjacent peaks with 12 Th spacing indicated in the position of carbon-carbon double bonds are labelled with (◆) symbols.

Hydroxy fatty acids

Hydroxy fatty acids (OHFA) are an abundant lipid class found in foodstuffs,\(^{37,38}\) along with bacteria and fungi.\(^{39,40}\) The specific location of hydroxylation is reported to affect metabolic functionality,\(^{41,42}\) necessitating analytical methods capable of elucidating the location of such motifs. Shown in Figure 6(a) is a PD spectrum generated following photolysis of mass-selected \([M + Na]^+\) ions (at \(m/z\) 539) of the 4-iodobenzyl ester derivative of 12-hydroxy stearic acid. The most abundant product ion in this spectrum is observed at \(m/z\)
412 and corresponds to the radical cation formed upon loss of atomic iodine. Subsequent CID of this species yields the RDD spectrum shown in Figure 6(b). Two major product ions can be seen in both PD and RDD spectra and represent the losses of C_7H_{14}O (m/z 298) and C_6H_{13} (m/z 327) from the [M + Na - I]^+ radical cation. Both product channels could derive from β-scission reactions directed from an alkoxy radical at the 12-position as indicated in Scheme 5. The dominance of these fragments over those arising from carbon-centred radicals, points to a preference for hydrogen abstraction from the hydroxyl moiety. Simple enthalpic arguments do not account for this selectivity however, with the oxygen-hydrogen bond dissociation energies of alcohols typically higher (~105 kcal mol^{-1}) compared to carbon-hydrogen bonds (~96 kcal mol^{-1}).^{33} Rather we propose that this selectivity may arise from the proximity of the hydroxyl group to the phenyl-iodide moiety in the three-dimensional structure of the gas phase ion. It seems plausible that in such a structure the hydroxyl group, along with the ester oxygens, would interact directly with the sodium cation placing the former in the vicinity of the benzyl ester moiety and thus the nascent phenyl radical upon liberation by photolysis. For comparison, the unsaturated hydroxy fatty acid 12-OHFA (9Z-18:1) – a major component of castor oil^{43} – was also investigated. PD and RDD mass spectra obtained from the 4-iodobenzyl ester derivatives are shown in Figure 6(c) and (d), respectively. While the PD spectrum is more complex than that of the corresponding saturated analogue (cf. Figures 6a and c), ions indicative of the location of the hydroxyl moiety and the double bond are clearly observed at m/z 296 (-C_7H_{14}O) and 325 (-C_6H_{13}). Indeed the RDD spectrum, obtained by subsequent CID on the [M + Na - I]^+ at m/z 410, is dominated by these two product ions. In summary, both hydroxy fatty acids investigated here show abundant free radical-driven dissociation at the site of the hydroxyl moiety. These findings suggest that either PD or RDD could be deployed to describe the site(s) of hydroxylation in an unknown hydroxyl fatty acid.
Figure 6: Mass spectra acquired from 4-iodobenzyl ester derivatives of the hydroxyl fatty acids, 12-OHFA (18:0) and 12-OHFA (9Z-18:1). (a) The PD spectrum of the [M + Na]$^+$ adduct ion of the 12-OHFA (18:0) derivative at m/z 539 and (b) the RDD spectrum acquired by performing subsequent CID on the m/z 412 radical ion. (c) The PD spectrum of the [M + Na]$^+$ adduct ion of the 12-OHFA (9Z-18:1) derivative at m/z 537 and (b) the RDD spectrum acquired by performing subsequent CID on the m/z 410 radical ion.
**Scheme 5:** A proposed dissociation mechanism to account for the formation of major product ions at m/z 298 and 327 resulting from decomposition of the radical cation at m/z 412 generated by PD of the 4-iodobenzyl ester derivative of 12-OHFA (18:0).

**Conclusion**

The standard esterification procedure employed here to produce the required FAIBE derivatives was successfully demonstrated for a variety of structurally diverse fatty acids. In general the esterification was high-yielding with the exception of the hydroxy fatty acids for which competing reactions (likely intramolecular lactonization) reduced the efficiency of the desired conversion. Nonetheless even in this case sufficient FAIBE was formed to demonstrate the effectiveness of the photodissociation strategy and alternative esterification protocols could be employed to improve this yield if required. Ionization of the FAIBE derivatives as sodium adducted ions and subsequent laser-photolysis at 266 nm was found to be efficient in the generation of the desired radical ions with yields of between 30 and 100% of the precursor ion abundance. In addition to the expected iodine loss, several of the derivatized fatty acids showed distinct fragmentation patterns upon photolysis. In particular, cleavage of activated carbon-carbon bonds was observed, such as homolysis of allylic carbon-
carbon bonds in unsaturated fatty acids. This selective fragmentation upon photoactivation may find application in selective screening of complex lipid mixtures for specific motifs such as double bond position(s) or sites of hydroxylation. For example, undertaking PD on a mass spectrometer of triple quadrupole geometry (such as that recently described by Dugourd, Lemoine and their co-workers\(^{44}\)) would allow multiple reaction monitoring or neutral loss scans to be undertaken targeting a particular structural motif or individual fatty acid isomer present within a complex mixture.

Subsequent CID of the \([M + Na - I]^+\) radical cations formed from these FAIBE derivatives gives rise to rich, radical-driven fragmentation and the resulting RDD mass spectra were able to distinguish between isomeric fatty acids. The usual mass difference between peaks is representative of the succession of methylene groups of the carbon chain. Thus, RDD of derivatized fatty acids containing saturated, non-branched acyl chains are characterized by spectra with regular peak spacings of 14 Th between successive neutral losses in the alkyl radical or alkene series. In the same way, any increase in the spacings between adjacent groups of peaks provides evidence for the presence of branching features or even hydroxylation in the acyl chain. For example, spacings of 27/28/29 Th were shown to locate both (i) the positions of methyl branched chain for the phytanic acid derivative (cf. Figure 2d) and (ii) the position of hydroxylation in 12-hydroxy stearic acid (cf. Figure 6b). Similarly, a decrease in peak spacing to 12 Th, was observed to arise at the position(s) of carbon-carbon double bonds within unsaturated fatty acids.

In conclusion, we have presented here a novel approach to lipid structure elucidation that is particularly suited to describing simple lipids such as fatty acids. The requirement to incorporate a chromophore and a photolabile moiety were readily satisfied in this instance by incorporation of 4-iodobenzyl alcohol by standard esterification procedures. As noted in the introduction however, a wide range of fatty acid derivatization procedures have previously been described (e.g., DMOX derivatives) and many of these could be readily adapted to
satisfy the requirements of RDD. It is thus attractive to consider exploitation of alternative
derivatives that may allow for improved derivatization and/or ionization efficiencies and even
greater selectivity in radical-directed fragmentation.

Acknowledgements:
S.J.B., A.J.T. and T.W.M. are grateful to the Australian Research Council (DP0986628 and
DP120102922) and the University of Wollongong for funding. H.T.P. was supported by a
matching scholarship from the University of Wollongong, T.W.M. is an Australian Research
Council Future Fellow (FT110100249) and S.J.B. and A.J.T are supported by the Australian
Research Council’s Centre of Excellence for Free Radical Chemistry and Biotechnology
(CE0561607).

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