A rapid and sensitive profiling of free fatty acids using liquid chromatography electrospray ionization tandem mass spectrometry (LC/ESI-MS/MS) after chemical derivatization†

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Free fatty acids (FFAs) have diverse roles in cellular energy and signaling and they are critical molecules in various biological states. Due to the poor ionization efficiency of FFAs under electrospray ionization mass spectrometry (ESI-MS) conditions, it is a challenging aspect to construct a robust platform for profiling of various FFAs in biological samples using liquid chromatography ESI-MS. In the present study, we applied trimethylsilyldiazomethane (TMSD) derivatization to improve ionization efficiencies in the profiling of FFAs. Multiple reaction monitoring (MRM) was used for the selective quantification of methylated FFAs. The optimal TMSD methylation was validated for a reliable FFA profiling. Furthermore, the high-throughput analysis of FFAs was successfully performed in short analysis and derivatization times. To verify the utility and effectiveness of the developed method, we compared both methylation and nonmethylation (intact FFA) data in the profiling of FFAs in mice liver and plasma. It is noteworthy that the methylation derivatization provided better results in FFA profiling. Further, we performed statistical data analysis where HBV and mock mice tissues were discriminated when the methylated FFAs data were used. In the lipidomics field, the present method can also be applied for the profiling of FFAs in biological samples for biomarker discovery. The present validated LC/ESI-MS/MS assay method may also be used for FFA profiling modeling studies in other biomedical samples.

Introduction

Free fatty acids (FFAs), categorized as a class of lipids, provide an important energy source as nutrients, and they also function as signalling molecules in various cellular processes including insulin secretion. FFAs are biomedical indicators of the abnormal lipid metabolism in various metabolic diseases including diabetes mellitus. Several researchers have applied FFA profiling to find biomarkers for the diagnosis and characterization of various metabolic diseases. In some studies, FFAs were also profiled to find the potential biomarkers of Alzheimer’s disease, coronary heart disease and other diseases.

Mass spectrometry (MS) has been widely used for the profiling of FFAs in biological samples. Gas chromatography (GC) coupled to electron ionization (EI) MS was usually applied to analyse fatty acid methyl ester (FAME) which is one of the most common fatty acid derivatives. By using this technique, the separation of positional and geometrical (cis/trans) isomers was also achieved in approximately 20 min. In addition to GC-MS, liquid chromatography (LC)-MS has also been established as an effective analytical technique in the profiling of FFAs with short run times. The use of electrospray ionization (ESI) MS alone, which is a soft ionization technique, only provides the information of molecular ions. Thus, tandem MS (MS/MS) is generally applied for the sensitive and selective analysis of FFAs. FFAs have been analysed in the negative ion mode and there is a limitation to the ionization efficiency. In the previous study, barium acetate was used as the cationization agent for the sensitive profiling of FFAs in the positive ion multiple reaction monitoring (MRM) mode using triple quadrupole mass spectrometer. Some researchers, Lee et al., and Pettinella et al., have demonstrated that trimethylaminoethyl ester iodide derivatization provided good sensitivity in the analysis of FFAs. Zhou et al. has also demonstrated that isotope-labeling derivatization by using 2,4-dimethoxy-6-piperazin-1-yl pyrimidine (DMPP) could improve ionization efficiency for analysing of FFAs. Yang et al. derivatized...
FFAs with 2-bromo-1-methylpyridinium iodide and 3-carbino-l-1-methylpyridinium iodide, forming 3-acloxyxymethyl-1-methylpyridinium iodide (AMMP). They observed that the detection sensitivity was approximately 2500-folds higher in positive-ion mode ionization than negative-ion mode.23 Similarly, Bollinger et al. have reported a method on conversion of FFAs carboxylic acid group in to an amide bearing permanent positive charge, N-(4-aminomethylphenyl)pyridinium (AMPP) and that method showed approximately 60 000-folds increased detection sensitivity in positive ion mode ionization when compared to negative ion mode.23 However, those methods are complex and time-consuming processes to derivatize FFAs.

In particular, a rapid and sensitive profiling method is required for the analysis of FFAs in clinical samples. The chemical derivatization can significantly increase the sensitivity and specificity of GC-MS and LC/ESI-MS method to analyse highly acidic compounds such as phosphopeptides, phospholipids and FFAs.24–29 Particularly, trimethylsilyldiazomethane (TMSD) derivatization method is extremely simple and it was applied for analysis of phosphopeptides26 and phospholipids29 to improve detection sensitivity. Recently, Lee et al. also demonstrated TMSD derivatization method that could improve peak shape and detection limits of many lipid classes.30 In the present study, LC/ESI-MS/MS method with TMSD methylation was used for the profiling of FFAs. The utility and effectiveness of the developed method was verified by the profiling of FFAs in plasma and liver samples.

Liver is a core organ which plays a key role in lipid metabolism and has various enzymes related to lipid synthesis such as fatty acid synthase (FAS), a key enzyme for FFA.31 Hepatitis B virus can cause various liver diseases such as cirrhosis and hepatocellular carcinoma.32 It is known that HBV virus disrupts various lipid synthesis-related enzymes in liver.33–34 Therefore, we have applied our developed method to HBV and mock mice tissues to identify FFA changes.

**Experimental**

**Materials**

HPLC-grade acetonitrile, methanol, water and isopropanol were purchased from J.T. Baker (Avantor Performance Material, Inc, PA, USA). Trimethylsilyldiazomethane (TMSD) reagent was purchased from Sigma-Aldrich, St. Louis, MO, USA. The FFAs, palmitic acid (16:0), oleic acid (18:1) and arachidonic acid (20:4) were purchased from Sigma-Aldrich St. Louis, MO, USA. The FFA standard, arachidonic acid-d8 (20:4-d8) was purchased from Cayman chemical, Ann Arbor, MI. The 26 G needle with 3 mL syringes were purchased from KOVAX syringe, Korea vaccine CO, Seoul, Korea. Zoletil and xylazine were purchased from Virbac S.A, France and Sigma-Aldrich Co. LLC., USA, respectively. Dulbecco’s phosphate-buffered saline (DPBS) was purchased from Thermo Fisher Scientific Inc., USA.

**Animal samples**

All mice used for our study are maintained in Konkuk University of Seoul, Republic of Korea according to standard animal care protocols and fed normal laboratory chow and tap RO water. All mice related experiments were carried out in accordance with Konkuk University IACUC (Institutional Animal Care and Use Committees). The four week aged female mice (C57BL/6) were purchased from Nara biotech co., ltd (Seoul, Korea). Animals were housed at temperature 23 ± 2 °C, with relative humidity of 50 ± 10% and light controlled environment. To obtain plasma samples, mice were anesthetized by intraperitoneal injection of 200 µL of zoletil-xylazine (40 mg mL−1)–DPBS solution (8 : 2 : 90; v/v) and plasma was collected by cardiac puncture and stored at −80 °C. In addition, livers particularly right lateral lobe were also excised from the same set of mice and washed with PBS to remove the plasma and connective tissue. After that they bled dry with filter paper wiper and then stored at −80 °C until analysis.

**Sample preparation**

Each lipid standard was dissolved in methanol and stored at −20 °C. We compared three common lipid extraction methods to select the better one for analysing FFAs.

First, we applied Bligh and Dyer method which is the most common lipid extraction method.35 In this method, a 750 µL of chloroform/methanol (1 : 2; v/v) was added to 50 µL of plasma samples and 50 mg of liver tissues which were taken in different Eppendorf tubes (1.5 mL). Next, a 10 µL of arachidonic acid-d8 (internal standard (IS)) solution was added to above plasma and tissue samples at a concentration of 10 ng µL−1. In the case of liver tissues, an additional step of homogenization was performed after the addition of chloroform and methanol. After vortexing for 1 min, samples were incubated in ice for 10 min. After that a 250 µL of chloroform and 450 µL of water were added to both plasma and tissue sample tubes. After the centrifugation (13 800 × g, 2 min at 4 °C), the organic phases were collected into different Eppendorf tubes.

Secondly, we performed extremely simple lipid extraction method where we used only methanol as extraction solvent. For about 50 µL of plasma sample, 10 µL of IS followed by 1 mL of ice cold methanol was added. After that, the sample mixture was vortexed for 30 s and incubated for 10 min in ice. After centrifugation (10 000 × g, 5 min at 4 °C), about 1 mL of supernatant was collected into another Eppendorf tube. In the case of liver...
All samples were stored at −80 °C before analysis. For TMSD methylation, the samples were dried in a SpeedVac concentrator and reconstituted with 100 μL methanol.

### TMSD methylation

A simple and rapid TMSD methylation derivatization method was used for the analysis of FFAs in the plasma and liver samples. A solution of TMSD (2 mol L$^{-1}$) in hexane (50 μL) was mixed in equal proportions with sample and standard (v/v). After vortexing for 30 s, mixture was incubated at 30 °C for 10 min under optimized conditions which has been described in the succeeding text. For quenching the derivatization reaction, glacial acetic acid (6 μL) was added as described previously.

### LC-ESI/MS/MS equipment and conditions

The HPLC analysis was performed on an Agilent 1290 infinity series HPLC instrument (Agilent Technologies, USA) equipped with binary pump (G4220A, USA), an autosampler (G4226A, USA), a column compartment (G1316C, USA) and a thermostat (G1330B, USA). The temperature of column oven and autosampler was set at 40 °C and 4 °C, respectively. For the separation of FFAs, Hypersil GOLD column (2.1 mm × 150 mm ID; 1.9 μm Thermo scientific) was used. The mobile phase solvent A consisted of a acetonitrile/methanol/water mixture (19 : 19 : 1) with 0.1% (v/v) formic acid and 20 mmol L$^{-1}$ ammonium
formate; and the mobile phase solvent B consisted of iso-
propanol with 0.1% (v/v) formic acid and 20 mmol L⁻¹ ammone-
nium formate. The flow rate of the mobile phase was 0.25 mL
min⁻¹ and the injection volume was 3 μL. A 20 min lipid elution
gradient was performed as follows: first 10 min, solvent
composition was set at 95% A and 5% B to elute FFAs; followed
by a linear gradient to solvent 90% A and 10% B for 2 min and
kept for 3 min for elution of other lipids. Finally, the column
was equilibrated at 5% solvent B for 5 min before reuse.

LC-MS analysis was performed on a triple quadrupole mass
spectrometer (QQQ LC-MS 6490 series, Agilent Technologies,
USA) equipped with an ESI source which provides high sensi-
tivity by iFunnel technology that consists of three components:
a hexabore capillary, Agilent Jet Stream technology, and a dual
ion funnel. The typical operating source conditions for MS
scan in the positive and negative ion ESI mode were optimized
as follows: capillary voltage 4000 V, nozzle voltage 500 V. The
nebulizer was set at 40 psig and the nitrogen drying gas was set
at a flow rate of 13 L min⁻¹ and the temperature was main-
tained at 250 °C. For collision-induced dissociation (CID)
experiments, the precursor ion of each FFA species was
selected using the quadrupole analyser and the product ions
were analysed using another quadrupole analyser. Ultra-pure
nitrogen was used as collision gas. The collision energies for
methylated and nonmethylated FFAs (intact FFA) were also
optimized by using FFA standards (8 eV for intact FFAs and 10
eV for methylated FFAs). All the spectra of FFAs were recorded
under optimized experimental conditions and the quantitative
analysis was performed in multiple reaction monitoring
(MRM) mode using computed transitions for methylated and
intact FFAs (Table 1).

Result and discussion

FFA profiling by LC/MS/MS with TMSD methylation

In this study, TMSD methylation was applied to increase the
sensitivity of FFA analysis. To the best of our knowledge, this is
the first attempt to analyse FFAs in plasma and liver samples
with TMSD methylation by LC/MS/MS. Thus, we established the
MRM transition (m/z value of precursor ion (Q1) > m/z value of
product ion (Q3)) for the profiling of methylated FFAs. The ESI-
MS conditions were also optimized in the analysis of methyl-
ated FFA standards (palmitic acid-saturated FFA, oleic acid-
monounsaturated FFA and arachidonic acid-polyunsaturated
FFA). The methylated FFAs were analysed as [M + H]⁺ ions in
positive ion mode whereas intact FFAs were predominately
analysed as [M − H]⁻ ions in negative ion mode. The m/z value of
the molecular ions of methylated FFAs showed the mass shift
of 15 Da to their intact molecular weights which corresponds to
the addition of methyl group (Fig. 1). The fragmentation pattern
of methylated and intact FFAs were confirmed by using tandem
MS analysis. In the negative ion mode, intact FFAs showed
similar fragmentation behaviour that was reported earlier by
Nagy et al., and other researchers.15,39-42 The diagnostic product
ion observed in the negative ion mode MS spectra of FFAs is [M
− H-44]⁻ which corresponds to the neutral loss of CO₂. On the
other hand, the positive ion mode analysis of methylated FFAs
showed [M + H-32]⁺ ion as the base peak which corresponds to
the loss of CH₃OH. The peak corresponds to the loss CH₃OH
was selected as Q3 transition in the MRM analysis of all FFAs
(Fig. 1). For the profiling of various FFAs, we fixed the MRM
transitions of each species based on their abundant character-
istic product ions of [M − H − CO₂]⁻ and [M + H − CH₃OH]⁺ for
intact and methylated FFAs, respectively.

Further, we also optimized the tested time (10, 15, 20, 25 and
30 min) and temperatures (10, 20, 30, 40, and 50 °C) for TMSD
methylation of FFAs. In these experiments, we analysed FFA
standards, C16:0 (palmitic acid-saturated FFA), C18:1 (oleic
acid-monounsaturated FFA) and C20:4 (arachidonic acid-
polyunsaturated FFA) with different derivatization time and
temperatures. The arachidonic acid-d₈ standard was used as an
IS for this analysis. According to different derivatization time
and temperature, the efficiency of methylation was estimated by
the peak area of FFA standards normalized by IS. 10 min of
derivatization time was enough for methylation of FFAs and 30
°C of reaction temperature showed the best efficiency for the
derivatization (Table 2). Thus, 10 min and 30 °C were selected as
the optimized conditions for methylation of FFAs.

The derivatization efficiency of TMSD methylation for FFA
profiling is given in Table 3. The methylated and intact FFAs
showed difference in ionization efficiency which may presum-
ably be due to their structural differences. To evaluate the effi-
ciency of TMSD methylation, we compared the peak areas
of intact FFAs in both methylated and nonmethylated mixtures.
The TMSD methylation efficiency of each FFA was described as
follows: the percentage of methylated FFAs ((peak area of
compound in nonmethylated FFAs – peak area of non-
methylated compound in methylated FFAs)/peak area of a
compound in a nonmethylated FFAs × 100 (%)) (Table 3). All
FFA standards showed high TMSD derivatization efficiency as
shown in Table 3 (the lowest efficiency was 82.9%).

Validation study

The performance of TMSD methylation was validated to
confirm whether the methylation is applicable to FFA profiling
in biological samples (Table 4). First, the reproducibility of

Fig. 1 Formation of TMSD derivative of arachidonic acid (C20:4) and its proposed fragmentation pattern.
methylation was estimated by calculating the relative standard deviation (% RSD) of 1 ng of FFA standards in intra- and interday experiments. In the analysis of four FFA standards, the % RSDs of the retention time and peak areas showed less than 15.1% and 1.6% RSD in both intra- and inter-day studies, respectively (Table 4). Second, the sensitivity of the FFA profiling with and without methylation were also compared. The limitation of detection (LOD) and coefficient ($R^2$) were calculated in terms of methylated and nonmethylated conditions. As a result, both methylated and intact FFA standards showed good coefficient (higher than 0.975). However, in the case of LOD, methylated standards showed better sensitivity than nonmethylated standards. It can be that noted that the intact oleic acid showed 25 fold higher LOD than the corresponding methylated standard (Table 4). Furthermore, sensitivity was significantly increased in case of palmitic acid and oleic acid standards (Fig. 2). Even though the intact arachidonic acid showed same LOD and sensitivity as similar to its methylated form, its average peak area was smaller than TMSD acid showed same LOD and sensitivity as similar to its methylated form, its average peak area was smaller than TMSD acid. The results of lower LOD, good coefficient and low % RSD of inter and intraday variation supported the reproducibility and reliability of our developed method. The matrix effect has been evaluated, where the matrix factor was ranged from 0.86–1.05. These data indicate that there was no significant endogenous interference. The stabilities of samples were within the limits and the mean percentage changes were less than ±10% from their nominal concentrations. Our results indicate that the samples were stable at 4 °C for 24 h in autosampler at –80 °C for 30 days. The samples were stable at room temperature for 6 h.

**Comparison of extraction methods of FFA**

For the optimization of extraction method, we compared three common lipid extraction methods (Method 1: Bligh and dyer; Method 2: extraction with methanol; Method 3: Dole’s procedure) with liver sample from C57BL/6 mice. A 50 mg each of liver tissue was used for each experiment to compare the extraction efficiency. As shown in Fig. 3, the overall area of FFAs extracted from methods 1 and 2 showed higher value than method 3. When we compared the extraction methods 1 and 2, they showed similar results in terms of the number of identified FFAs (14 FFAs in both methods) and relative area (area of FFA/area of IS) of FFAs. As these extraction methods
Table 4  Data of validation of methylated fatty acids and LODs

| Species               | RT\(^a\) (min) | Intraday variation | Interday variation | Coefficient (\(R^2\)) | LOD (ng) | Linearity range (ng) |
|----------------------|-----------------|--------------------|--------------------|------------------------|----------|----------------------|
| Palmitic acid        | 1.59            | 1.6                | 5.6                | 1.1                    | 15.1     | 0.9931 0.9869 1600 800 800–25000 |
| Oleic acid           | 2.8             | 0.2                | 5                  | 0.4                    | 7.1      | 0.9819 0.9927 1000 40 40–25000 |
| Arachidonic acid     | 2.15            | 0.4                | 4.6                | 0.4                    | 4.8      | 0.9909 0.9943 100 100 100–100000 |
| (d8) arachidonic acid-IS | 2.09          | 0.9                | 1.7                | 0.6                    | 15       | 0.991 0.975 100 100 100–100000 |

\(^a\) Average of retention time (\(n = 9\)). \(^b\) % RSD (\(n = 9\)) of intraday variation. \(^c\) % RSD (\(n = 27\)) of interday variation (3 days).

Fig. 2  Improved sensitivity of FFA standards by TMSD methylation. Panel (A) MRM chromatogram of intact FFA standards (C16:0 (palmitic acid-saturated FFA), C18:1 (oleic acid-monounsaturated FFA) and C20:4 (arachidonic acid-polyunsaturated FFA)); Panel (B) MRM chromatograms of methylated FFAs where they showed different RT due to methylation (addition of methyl group to compound).

Fig. 3  Comparison among three common FFA extraction methods (Method 1: Bligh & dyer; Method 2: extraction using methanol; Method 3: Dole’s mixture).
are very complex and tedious time consuming processes which may cause the experimental handling error, we moved to the simple extraction method which involves methanol as extraction solvent. Finally, methanol extraction method was selected due to its simplicity and good performance (Fig. 3).

**Application of TMSD methylation for profiling of FFAs in mice liver and plasma**

In order to validate the utility of TMSD methylation for the profiling of FFAs in biological samples, we applied our validated method for analysing mice liver and plasma (C57BL/6 mice). The arachidonic acid-d8 (20:4) was used as the IS for the normalization of each FFA species. The relative area (area of FFA/area of IS) of FFA species was characterized for the detailed phenotype of mice samples. Each sample (plasma and tissues) was analysed four times (n = 4). Furthermore, the results of FFA profiling with and without methylation were compared. Compared to intact FFAs, the RT of methylated FFAs was slightly delayed due to the formation of methyl ester structure (Fig. 2 and 4). It can be noted that the methylated FFAs were eluted within 5 min even though their RTs were delayed. Thus, it is appropriate to perform the high-throughput profiling of FFAs. Upon methylation, 11 FFAs in mice plasma and 16 FFAs in mice liver were successfully analysed. On the other hand, in the case of intact FFAs, six FFAs in mice liver and plasma were detected (Table 5). It is noteworthy that five methylated FFAs (C14:1, 18:1, 18:2, 18:3, and 18:4) were additionally analysed in mice plasma when compared to intact FFAs. Furthermore, ten FFAs (C16:1, 18:0, 18:1, 18:2, 18:3, 18:4, 20:1, 20:2, 22:2, and 22:3) were only detected by TMSD methylation in mice liver. The average relative areas of methylated FFAs are much higher than the intact FFAs (Fig. 4 and Table 5). All the detected FFAs were also validated by tandem mass spectrometric experiments (Fig. 5 and S1†).

**Application of TMSD methylation in the FFA profiling in HBV infected mice**

We also applied the developed method for profiling of FFAs in three samples (HBV infected samples and its mock mice liver tissue). In these experiments, each tissue was analysed four times (n = 4). The peak area of individual FFA species was divided by internal standard for normalization and relative quantification. The normalized FFA data of mice samples were subjected to principal component analysis (PCA) and hierarchical clustering analysis (HCA) (heat map, dendrogram) which were performed through the web-based data analysis software, MetaboAnalyst 3.0. In addition, the uses of FFA profiling with and without methylation were also compared in this study. When the data used that was obtained by

![Fig. 4](image_url) Comparison between intact and methylated FFAs in liver tissues. Panel (A) MRM chromatograms of intact FFAs, C18:1, C18:2, C18:3 and C18:4 in liver (no FFAs were detected). Panel (B) The MRM chromatograms of methylated FFAs (the data indicated that the sensitivity was significantly increased after methylation derivatization).
nonmethylation, HBV and mock mice tissues were not well separated in the PCA score plot and HCA (Fig. 6A). On the other hand, HBV and mock mice tissues were clearly distinguished when the data of methylated FFAs was used in the statistical analysis (Fig. 6B). As it can be seen from Fig. 6B, many FFAs were decreased in HBV induced mice and finally we found some significantly changed FFAs (C16, C16:1, C20:1, C20:2, C20:3, C20:4, C20:5, C22:1, C22:3, C22:4 and C22:5; \( P \)-value < 0.05) in methylated conditions. These results indicated that the use of methylation can provide the comprehensive information of FFA lesions with higher intensity and more number of identifications than the nonmethylation.

**Table 5** Fatty acid species analysed in mice liver and plasma by QqQ with and without methylation

| Species           | RT\(^a\) (min) | Liver\(^b\) | Plasma\(^b\) |
|-------------------|----------------|------------|-------------|
|                   | Intact | Methylated | Intact | Methylated | Intact | Methylated |
| C8:0              | —      | —          | —      | —          | —      | —          |
| C10:0             | —      | —          | —      | —          | —      | —          |
| C12:0             | —      | —          | —      | —          | —      | —          |
| C14:0             | —      | —          | —      | —          | —      | —          |
| C14:1Δ9           | —      | 1.6        | —      | —          | —      | 0.48 ± 0.05\(^c\) |
| C16:0             | —      | 3          | —      | —          | —      | —          |
| C16:1Δ9           | —      | 2.3        | —      | 1.09 ± 0.12| —      | —          |
| C18:0             | —      | 3.7        | —      | 0.09 ± 10  | —      | —          |
| C18:1n9           | —      | 3          | —      | 17.7 ± 1.65| —      | 1.47 ± 0.22 |
| C18:2n6           | —      | 2.5        | —      | 8.44 ± 0.82| —      | 0.72 ± 0.13 |
| C18:3n3 or n6     | —      | 2.1        | —      | 0.68 ± 0.10| —      | 0.23 ± 0.04 |
| C20:4             | —      | 1.4        | —      | 0.91 ± 0.11| —      | 1.10 ± 0.15 |
| C20:0             | —      | —          | —      | —          | —      | —          |
| C20:1Δ11          | —      | 3.9        | —      | 1.13 ± 0.09| —      | —          |
| C20:2Δ11,14       | —      | —          | —      | 1.54 ± 0.21| —      | —          |
| C20:3n3 or n6     | 2.2    | 2.6        | 0.26 ± 0.01| 10.5 ± 0.94| 0.01 ± 0.00| 0.19 ± 0.04 |
| C20:4n6           | 1.9    | 2.3        | 22.2 ± 1.42| 21.5 ± 2.39| 1.03 ± 0.01| 0.15 ± 0.04 |
| C20:5n3           | 1.7    | 1.8        | 5.43 ± 0.25| 2.26 ± 0.18| 0.16 ± 0.01| 23.0 ± 3.07 |
| C22:0             | —      | —          | —      | —          | —      | —          |
| C22:1n9           | —      | —          | —      | —          | —      | —          |
| C22:2Δ13,16 or n6 | —      | 4          | —      | 0.14 ± 0.03| —      | —          |
| C22:3             | —      | —          | —      | —          | —      | —          |
| C22:4n6           | 2.2    | 2.7        | 0.19 ± 0.02| 1.54 ± 0.20| 0.02 ± 0.00| 0.20 ± 0.00 |
| C22:5n3           | 1.9    | 2.4        | 3.40 ± 0.22| 4.28 ± 0.44| 0.30 ± 0.01| 6.26 ± 1.10 |
| C22:6n3           | 1.7    | 2          | 32.7 ± 1.98| 21.7 ± 2.32| 2.59 ± 0.11| 1.55 ± 0.24 |
| C24:0             | —      | —          | —      | —          | —      | —          |
| C24:1Δ15 or n9    | —      | —          | —      | —          | —      | —          |

\(^a\) Average of retention time. \(^b\) Average of relative peak area (compound/internal standard) ± S.D \((n = 4)\). \(^c\) The values of percentages are in mean ± SD \((n = 4)\).

Fig. 5 Tandem mass spectra of representative methylated C18:1 FFA (the \( m/z \) 297 \([M + H]^+\) ion corresponds to methylated C18:1 FFA (oleic acid) in positive ion mode).
Conclusions

In this study, the LC/MS/MS with simple and rapid methylation method was used for the profiling of FFAs. The proposed method is simple for profiling of methylated and various other FFA species within a short analysis time (5 min) and derivatization time (10 min). We observed that our method has high derivatization efficiency after the optimization of TMSD methylation by using FFA standards (palmitic acid-saturated FFA, oleic acid-monounsaturated FFA, arachidonic acid-polyunsaturated FFA). The results also revealed that methylated FFAs showed higher detection sensitivity, reproducibility and reliability in the positive ion mode than intact FFAs in the negative ion mode. The data showed that our methylation method provided in detail information of FFAs with higher intensity and more number of identifications than the nonmethylation. Besides, we also applied the developed method for profiling of FFAs in animal models. The utility of FFA profiling with methylation was also proved by PCA and HCA of HCV infected and control liver tissues. In the lipidomics, this method can be used for analysing even low abundant FFA species qualitatively and quantitatively. This method can also be applied to many biological and clinical samples to discover FFA biomarker. Furthermore, this study also explained the utility and effectiveness of methylation derivatization in profiling of FFAs in biological samples (mice liver and plasma).

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