Data Descriptor: Comprehensive high-resolution multiple-reaction monitoring mass spectrometry for targeted eicosanoid assays

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Eicosanoids comprise a class of bioactive lipids derived from a unique group of essential fatty acids that mediate a variety of important physiological functions. Owing to the structural diversity of these lipids, their analysis in biological samples is often a major challenge. Advancements in mass spectrometric have been helpful for the characterization and quantification of these molecular lipid species in complex matrices. However, there are technical limitations to this approach, including low-abundant and/or poorly ionizable lipids. Using high-resolution multiple-reaction monitoring (MRMHR), we were able to develop a targeted bioanalytical method for eicosanoid quantification. For this, we optimized the LC-MS/MS conditions and evaluated several parameters, including linearity, limits of quantification, matrix effects and recovery yields. For validation purposes, we looked at the method’s precision and accuracy. A library of high-resolution fragmentation spectra for eicosanoids was developed. Our comprehensive dataset meets benchmark standards for targeted analysis, having been derived using best-practice workflows and rigorous quality assessments. As such, our method has applications for determining complex eicosanoid profiles in the biomedical field.

Design Type(s)
- replicate design
- quality control testing design

Measurement Type(s)
- Calibration
- calibration substance role

Technology Type(s)
- liquid chromatography-tandem mass spectrometry

Factor Type(s)
- Matrix
- Concentration
- pH
- technical replicate

Sample Characteristic(s)

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Background & Summary

Lipidomics involves the identification and quantification of lipids within a given biological system. Recent advances have been enabled by the development of new mass spectrometric tools and protocols for the analysis of molecular lipids in complex matrices. Polyunsaturated fatty acids (PUFAs) are known to mediate some of their bioactivities through the formation of oxygenated metabolites. These bioactive lipids include eicosanoids, a class of lipid mediators derived from arachidonic acid via cyclooxygenase (COX), lipooxygenase (LOX), and cytochrome-P450-catalyzed reactions (CYP450), and through non-enzymatic lipid peroxidation. The oxygenated lipid mediators play prominent roles in the physiological and pathological regulation of many biological processes, including those observed in inflammatory diseases.

Using targeted, as opposed to global, lipid analysis provides an alternative method in lipidomics. The analysis of lipid mediators in biological samples has its own set of challenges. This is because these lipids are often present in extremely low concentration, are transiently formed on demand by cells, and often have limited half-lives. To compound the problems faced by researchers, many isomeric species can be generated, which also have a specific metabolic function. Therefore, highly sensitive and accurate methods are urgently needed for the analysis of eicosanoids.

A wide range of techniques has been used for the separation, detection, and quantification of lipid mediators. Currently, progress in liquid-chromatography–electrospray ionization tandem mass spectrometry (LC–ESI-MS/MS) offers a powerful approach for this purpose. The advantage of mass spectrometry is its capacity to separate and characterize ionized analytes according to their mass-to-charge ratios (m/z). Collision-activated dissociation (CAD) can be used to determine structural information acquired through the lipid’s ion fragmentation. LC-MS/MS methods for targeted lipidomics utilize scheduled multiple-reaction monitoring (MRM) to optimize sensitivity, maximize the number of metabolites that can be analyzed at any given time, and reduce the time required for analysis. The beauty of LC is that it simplifies analyses and offers the greatest chromatographic advantage for separating complex mixtures of lipid mediators because no prior derivatization of the analytes was required. Despite this, chemical similarities and the presence of isomeric species still make separating lipid mediator species via LC rather difficult.

MS detection of the biological molecules was found to be dependent on ESI for ion generation, which produced both positive and negative molecular ion species ([M+H]+ and [M-H]−), even though our

Figure 1. Schematic outline of metabolites derivate from arachidonic acid (AA-20:4) via the lipooxygenase (LOX), cyclooxygenase (COX), CYP450 (CYP) or free radical catalyzed pathways. The enzymes involved in eicosanoid biosynthesis were in green and orange arrows indicated the catabolism pathway. Major metabolites (Blue) derived from those metabolic pathways are included in the MRM quantitation assay. The non-stable intermediated metabolites (Red) were not included in the method.
analysis was conducted in the negative-ion mode. Tandem mass spectrometry (MS/MS) is known to increase the sensitivity and specificity of quantitative lipid analyses. Since the acquisition mode MRM in most lipidomics studies is used to monitor the analytes through the transition of a select precursor ion to a specific product ion, there are several lists which feature characteristic eicosanoids MRM transitions performed on triple-quadrupole instruments.

In order to gain greater confidence in detection and quantification methods, the standard practice follow selecting two or more transitions per precursor ion for analysis. Modern instruments have the required speed to cycle through many precursor ions with multiple transitions within a short time window. Indeed, as an alternative to the classical MRM performed on triple-quadrupole, hybrid systems like the TripleTOF5600, a quadrupole time-of-flight (QTOF) instrument can be used to acquire high-resolution MRM (MRMHR). In a MRMHR assay, the first quadrupole basically selected the precursor ion passing into the collision cell, and the instrument subsequently monitored all transitions of this precursor ion. The TOF analyzer recorded the resultant high-resolution spectrum. If retention times were known prior to analysis, many precursors would be selected during a scheduled MRMHR experiment; providing higher confidence detection, since all possible transitions can be monitored. Indeed, targeted eicosanoid lipidomics could be performed using a comprehensive MRMHR strategy, which included important analytical steps, such as the standardization of the operating procedures for sample collection and preparation, as well as the development of both the LC-MS/MS method and the computational process (Fig. 2).

Methods

Descriptions for the procedures were taken from our previous work and were either completely adapted or supplemented with new details where necessary.
Reagents
All eicosanoids, fatty acids (molecular weight standards, MWS), and deuterated internal standards (IS) were purchased from Cayman Chemical Co (Ann Arbor, MI, USA). The details of all analytical-grade standards are listed in file “Metabolite standard details (docx) and Metabolite standard details (pdf), Data Citation 1”. HPLC-grade solvents acetonitrile (ACN), methanol (MeOH), isopropanol were purchased from Merck (Kenilworth, NJ, USA), and deionized water (H2O) were obtained from Milli-Q integral water purification System for ultrapure water from Merck-Millipore (Kenilworth, NJ, USA). The acetic acid and ammonium hydroxide (NH4OH) used in these experiments came from Sigma-Aldrich (St. Louis, MO, USA).

Animals
Six-week-old male C57BL/6 mice were obtained from Faculdade de Ciências Farmacêuticas de Ribeirão Preto, FCFRP/USP. All animal experiment protocols were in compliance with institutional guidelines for ethics in animal experiments approved by the Animal Care Committee of the Universidade de São Paulo (Permit no. 11.1.468.53.6). Euthanasia protocols were performed under CO2/O2 excess atmosphere. Furthermore, all efforts were made to minimize suffering.

Sample preparation

Standard Solutions and Calibration Curve: Individual stock solutions of lipid mediators (MWS) and IS were prepared at 10 μg.mL−1 in MeOH. A standard working solution (WS) composed of a mixture of all of the lipids (MWS and IS) was prepared via direct infusion of 100 ng.mL−1 in MeOH/H2O (7:3, υ/υ) containing 0.1% NH4OH by volume; this was to establish the MRM parameters, such as the declustering potential (DP) and the collision energy (CE). The WS was prepared without adding any isomer compounds to the same solution. For LC parameter studies, the WS was prepared as described with no addition of NH4OH. In another set of experiments, the calibration curves for the validation assay were obtained by diluting the relevant lipids in MeOH/H2O (7:3, υ/υ) to the following final concentrations: 2.3, 4.6, 9.2, 18.5, 37.0, 74.0, 148.1, 222.2, 333.3, and 500 ng.mL−1. To determine matrix effect and recovery yields, IS solutions were prepared in MeOH at three concentrations 18.5, 148.1, and 500 ng.mL−1. All solutions were stored in amber glass vials at −80°C and under argon atmosphere until needed.

Solid Phase Extraction (SPE): Pooled plasma and lung tissue samples were obtained from C57BL/6 mice. Dulbecco’s Modified Eagle Medium (DMEM, Gibco-Grand Island, NY, USA) was used as the culture medium for the validation assay. Lipid extraction was carried out according to a previously described protocol15,16, but with slight modifications. In brief, each sample was spiked with 10 μL of the IS solution before being extracted for use either in the recovery assay or the matrix effect assay. Lung samples were prepared via homogenization (Mixer Homogenizer-Labortechnik, Staufen, Germany) of 50 mg of tissue per mL of MeOH/H2O (1:1, υ/υ) solution. Plasma samples were made via protein precipitation of 150 μL of the sample in 1.5 mL of MeOH/ACN (1:1, υ/υ) at 4 °C, which was left to denature overnight. Afterward, both plasma and lung samples were centrifuged for 10 min at 4 °C and 800 x g. For culture medium extraction a solution in MeOH was prepared (1:1, υ/υ). Subsequently, the supernatant samples, from lungs, plasma or culture medium were diluted with water to a maximum solvent concentration of 15%.

For the first step in the SPE extraction protocol, the cartridge (Hypersep C18-500 mg, 3 mL, Thermo Scientific-Bellefonte, PA, USA) was washed with 4 mL of MeOH and equilibrated with 4 mL of H2O that had been generated using an extraction manifold (Waters-Milford, MA, USA). After loading the diluted samples, the cartridges were again flushed with 4 mL of H2O to remove hydrophilic impurities. The analytes which had been adsorbed on the SPE sorbent were eluted with 1 mL of MeOH and stored at −80 °C to prevent metabolite degradation. The solvent was removed in vacuo (Concentrator Plus, Eppendorf, Germany) at room temperature and re-dissolved in 50 μL of MeOH/H2O (7:3, υ/υ) for LC-MS/MS analysis.

LC-MS/MS analysis
The Ultra-High-Performance Liquid Chromatography (UHPLC) system (Nexera X2, Shimadzu-Kyoto, HO, Japan) was equipped with a binary pump system, a SIL-30AC autosampler, a DGU-20A degasser and a CBM-20A controller. The conditions used for chromatography are outlined in a previously published work16,17. However, some modifications have been implemented to improve the eicosanoids quantifications. For this propose, the method was optimized using an Ascentis Express C18 column (Supelco - St. Louis, MO, USA) with an i.d. of 100 ×4.6 mm and a particle size of 2.7 μm. Elution was conducted under a binary gradient system which consisted of Phase A, H2O/ACN/acetic acid (69.9:30:0.02, υ/υ/υ) at pH 5.8 (adjusted with NH4OH), and Phase B, an ACN/isopropanol (70:30, υ/υ). Gradient elution was carried out for 25 min at a flow rate of 0.6 mL.min−1. Gradient conditions were as follows: 0 to 2.0 min, 0% B; 2.0 to 5.0 min, 15% B; 5.0 to 8.0 min, 20% B; 8.0 to 11.0 min, 35% B; 11.0 to 15.0 min, 70% B; and 15.0 to 19 min, 100% B. At 19.0 min, the gradient returned to the initial condition of 0% B, and the column was re-equilibrated until 25.0 min. Over the course of the analyses, the column was kept at 25 °C, and the samples were maintained at 4°C in the autosampler. A 10 μL aliquot of each sample was injected onto the column. The pH of the mobile phase A was optimized to achieve the best sensitivity and chromatographic separation of lipid metabolites. As such, the pH values of phase A throughout the experiment were 3.8, 5.8, and 6.4.
The UHPLC system was interfaced with a TripleTOF5600+ Mass Spectrometer (Sciex-Foster, CA, USA) equipped with a Turbo-V IonSpray. An Atmospheric-Pressure Chemical Ionization probe (APCI) was used for external calibrations of the Calibrant Delivery System (CDS). Automatic mass calibration (< 2 ppm) was performed using APCI Negative Calibration Solution (Sciex-Foster, CA, USA) injected via direct infusion at 300 μL.min⁻¹ flow rate; this was done periodically after each of the five sample injections. An Electrospray Ionization (ESI) source in the negative ion mode was utilized for MRMHR scanning. For optimization of MRMHR channel settings, CE and DP values were determined for each analyte via direct infusion of MWS and IS solutions at a flow rate of 10 μL.min⁻¹. Compounds were fragmented via CAD using nitrogen as the collision gas. Additional instrumental parameters were as follows: nebulizer gas (GS1), 50 psi; turbo-gas (GS2), 50 psi; curtain gas (CUR), 25 psi; electrospray

| Common name         | Retention time (min) | r²  | LLOQ* (pg) |
|---------------------|----------------------|-----|------------|
| 20-OH-LTB₄          | 3.06                 | 0.9986 | 0.05 |
| LTC₄                | 5.09                 | 0.9987 | 0.09 |
| PGB₂                | 8.70                 | 0.9993 | 0.02 |
| 15-keto-PGE₂        | 5.99                 | 0.9994 | 0.05 |
| 20-OH-PGE₂          | 1.80                 | 0.9989 | 0.05 |
| TXB₂                | 4.06                 | 0.9987 | 0.05 |
| LXA₄                | 5.93                 | 0.9993 | 0.09 |
| PGD₂                | 5.56                 | 0.9988 | 0.05 |
| 6-keto-PGF₁α        | 2.45                 | 0.9963 | 0.18 |
| PGE₁                | 5.20                 | 0.9993 | 0.02 |
| RvD₂                | 6.23                 | 0.9989 | 0.09 |
| PGE₃α               | 4.63                 | 0.9988 | 0.09 |
| 19-OH-PGR₂          | 2.28                 | 0.9982 | 0.09 |
| PGG₁                | 6.73                 | 0.9919 | 0.37 |
| LTB₅                | 10.54                | 0.9981 | 0.02 |
| LTD₃                | 6.90                 | 0.9993 | 0.05 |
| LTA₄                | 7.90                 | 0.9987 | 0.09 |
| 6-trans-LTB₅        | 10.09                | 0.9977 | 0.05 |
| 11-trans-LTD₄       | 6.90                 | 0.9991 | 0.09 |
| PD₂                 | 10.40                | 0.9991 | 0.05 |
| MaR-1               | 10.21                | 0.9986 | 0.05 |
| PGH₂                | 7.61                 | 0.9966 | 0.09 |
| PGI₂                | 8.65                 | 0.9993 | 0.05 |
| 15-deoxy-8,12,14-PGE₂| 13.23               | 0.9970 | 0.09 |
| 5-HETE              | 14.41                | 0.9989 | 0.05 |
| AA                  | 17.17                | 0.9675 | 0.09 |
| 5-oxo-ETE           | 14.68                | 0.9986 | 0.02 |
| 20-HETE             | 13.35                | 0.9977 | 0.09 |
| 5,6-DHETE           | 12.91                | 0.9992 | 0.02 |
| 12-HETE             | 14.22                | 0.9993 | 0.05 |
| 8-HETE              | 14.22                | 0.9984 | 0.02 |
| 11-HETE             | 14.08                | 0.9972 | 0.05 |
| 12-oxo-ETE          | 14.69                | 0.9980 | 0.05 |
| 15-oxo-ETE          | 13.99                | 0.9969 | 0.05 |
| 11,12-DHETE         | 12.94                | 0.9981 | 0.05 |
| 14,15-DHETE         | 12.73                | 0.9931 | 0.05 |
| EPA                 | 16.71                | 0.9951 | 0.09 |
| 5,6-DHETE           | 13.22                | 0.9974 | 0.09 |
| 15-HETE             | 13.89                | 0.9967 | 0.09 |

Table 2. Retention time, r², and LLOQ for all lipid mediator targets analyzed at pH 5.8. *Lower limit of quantification values were considered for accuracy within 20% of the nominal concentration (CV ≤20%). * Calibration curves include six runs, with ten concentrations.
Data processing
Identification of the lipid species obtained in the LC-MS analysis was performed using PeakView 2.1 (Sciex-Foster, CA, USA). Briefly, this software combines information in three dimensions to give a representation of the retention time, signal intensity, and m/z value for each analyte. PeakView software is a stand-alone application for the qualitative review of LC-MS data files. In addition to its general application in exploring and interpreting qualitative data, the software also provides the tools for analyzing accurate and nominal mass data, structures, and multiple samples at any given time. Data processing usually proceeds through multiple steps, including filtering, feature detection, alignment, and normalization. Feature detection is conducted to identify all signals caused by true ions and to avoid the detection of false positives, thereby interpolating theoretical information from the molecular data file and experimental fragments. In our study, alignment to confirm retention time differences between runs was conducted through a comparison of the IS chromatography peaks. Normalization steps help in removing factors, which cause unwanted systematic bias and can be used to determine extraction efficiency during sample preparation via ion intensities and/or peak area between measurements. This was performed by extrapolation of the area ratio among the IS and biological analyte peaks.

For quantitative analysis, we used MultiQuant. This software uses robust integration algorithms, such as MQ4 and advanced SignalFinder, and is able to generate reliable integration results faster with less user intervention in order to maximize productivity. The use of reference standards for each of the target lipid species or class was important in quantifying the lipids under investigation. In this field, it is common practice to normalize the individual molecular ion-peak intensities using an internal standard for each lipid class. The calculated ratio of analyte to internal standard is then multiplied by the concentration of said internal standard so as to obtain the concentration of a particular analyte present in the sample. To quantify the lipid mediators in our work, MRM\textsuperscript{HR} channels were created during computational data processing. These were instructed to select product ions of higher intensity and selectivity in order to achieve greater analytical sensitivity and avoid cross-identification. The MRM\textsuperscript{HR} optimized parameters used in LC-MS/MS analysis are presented in Table 1 (available online only). The final eicosanoid concentration in biological samples was usually normalized by volume, tissue weight, cell number or protein/DNA concentration.

Validation parameters
The method was partially validated in accordance with Food and Drug Administration (FDA) recommendations\textsuperscript{18}. The parameters of interest were the analyte’s linearity, the lower-limits of quantification (LLOQ), inter- and intraday precision and accuracy, recovery and the matrix effects in the culture medium and both the plasma and lung tissue samples.

Linearity and LLOQ. These were assessed by means of 10-point standard curves prepared for five replicate samples within a linear range (Table 2). Standard curves (linear regression, 1/x weighted) were constructed by plotting the standard area against its nominal concentration (File: ‘Retention time for lipid mediators panel under different UHPLC-MS/MS chromatography conditions (pdf)’, Data Citation 1). The acceptance criterion for the coefficient of variation (CV) was ≤20% for the LLOQ level and ≤15% for levels above the LLOQ. The LLOQ was defined as the lowest concentration level at which the signal-to-noise ratio (S/N) is ≥10 and the CV is ≤20%.

Inter- and Intraday Precision and Accuracy. These were evaluated at three concentration levels for the quality control samples (QC): 0.15 ng.mL\textsuperscript{−1} for low-concentration quality control (LQC) samples; 1.5 ng.mL\textsuperscript{−1} for medium-concentration quality control (MQC) samples; and 5 ng.mL\textsuperscript{−1} for high-concentration quality control (HQC) samples. Precision and accuracy were expressed in percentage as the Relative Standard Deviation (RSD) and the Relative Error (RE), respectively. Accuracy was defined as the difference between the expected concentration (C\textsubscript{ex}) and the average measured concentration (C\textsubscript{av}) as calculated by the formula (C\textsubscript{ex} - C\textsubscript{av}/C\textsubscript{av}) × 100. The precision, RSD, was calculated using the formula (SD/C\textsubscript{av}) × 100. The method was only considered to be precise if the RSD was ≤15% (≤20% for LLOQ) and accurate if the RE of the analyte concentration measured was ≤15% of the expected value (≤20% for LLOQ). The intraday precision (repeatability) and accuracy were established for five replicates for the LQC, MQC, and HQC over the course of the same day using the same calibration curve. For the interday analysis (reproducibility), the assay was conducted for three consecutive days. Interday accuracy and precision were assessed by running analysis for five calibration curves in quintuplicate over consecutive days. According to FDA guidelines, the conventionally accepted range of inter- and intraday accuracy is from 80 to 120%. Accuracy and precision results for the lipid mediators are given in Table 3 (available online only) and respectively full results on file ‘Values obtained for each replicate of quality control (HQC, MQC and LQC) assay (xlsx) and Values obtained for each replicate of quality control (HQC, MQC and LQC) assay (pdf)’, Data Citation 1.
### Table 4. Mean recovery and matrix effect of the lipid mediator targets using SPE extraction in HQC, MQC, and LQC levels of IS in plasma samples.

| Internal Standard | Recovery Rate (% ± SD)\(^a\) | Matrix Effect (% ± SD)\(^b\) |
|-------------------|-----------------------------|-----------------------------|
|                   | HQC\(^c\) | MQC\(^d\) | LQC\(^e\) | HQC\(^c\) | MQC\(^d\) | LQC\(^e\) |
| 15-HETE-\(d_6\)   | 80.5 ± 12.1 | 99.6 ± 11.7 | 135.2 ± 25.4 | 106.4 ± 7.6 | 118.8 ± 20.0 | 139.7 ± 11.6 |
| TXB\(_2\)-\(d_6\)  | 58.9 ± 7.9  | 70.4 ± 6.8  | 75.0 ± 11.1  | 70.9 ± 3.9  | 72.1 ± 9.18 | 70.4 ± 7.1 |
| 6-keto-PGF\(_{1α}\)-\(d_6\) | 136.2 ± 15.1 | 143.8 ± 25.3 | -----        | 145.3 ± 14.9 | 154.4 ± 27.4 | -----        |
| PGJ\(_2\)-\(d_6\)   | 747 ± 8.6   | 917 ± 5.6   | 946 ± 13.2   | 885 ± 6.4   | 1001 ± 15.3 | 937 ± 10.4 |
| PGD\(_2\)-\(d_6\)   | 844 ± 8.2   | 956 ± 4.2   | 890 ± 29.3   | 101.9 ± 9.7 | 106.7 ± 23.7 | 96.3 ± 17.8 |
| LXA\(_4\)-\(d_6\)   | 74.1 ± 7.5  | 84.1 ± 10.9 | -----        | 101.8 ± 9.2 | 96.8 ± 18.8 | 150.1 ± 24.4 |
| RvD\(_1\)-\(d_6\)   | 76.1 ± 9.1  | 92.0 ± 9.4  | 115.1 ± 13.7 | 100.6 ± 10.2 | 107.2 ± 21.4 | 107.0 ± 21.9 |
| PGD\(_2\)-\(d_6\)   | 44.8 ± 3.6  | 48.0 ± 8.2  | 57.6 ± 12.7  | 41.3 ± 3.4  | 48.5 ± 8.5  | 59.5 ± 13.7 |
| 12-deoxy-\(\beta\)-LTB\(_4\)-\(d_6\) | 79.7 ± 3.4 | 98.0 ± 11.2 | 104.2 ± 14.4 | 97.1 ± 6.8 | 109.7 ± 17.4 | 116.3 ± 11.9 |
| LTE\(_4\)-\(d_6\)    | 62.2 ± 7.5  | 69.9 ± 7.9  | -----        | 132.1 ± 6.6 | 141.5 ± 24.9 | 201.8 ± 102.8 |
| AA-\(d_6\)          | 51.3 ± 1.9  | 27.9 ± 4.8  | 21.9 ± 5.1   | 58.7 ± 5.9  | 27.6 ± 6.7  | 26.1 ± 6.4 |
| 15-deoxy-\(\delta\)-12,14-PGJ\(_2\)-\(d_6\) | 69.4 ± 9.5 | 94.0 ± 22.9 | 191.8 ± 29.0 | 109.8 ± 8.2 | 131.1 ± 32.9 | 167.7 ± 30.1 |
| 5-HETE-\(d_6\)      | 76.8 ± 10.4 | 96.7 ± 8.7  | 96.6 ± 13.4  | 99.1 ± 6.5  | 102.2 ± 15.4 | 117.7 ± 7.4 |
| 5-iso-ETE-\(d_6\)   | 81.9 ± 12.9 | 87.9 ± 12.2 | 108.4 ± 25.5 | 99.8 ± 6.8  | 93.5 ± 16.7 | 120.7 ± 24.9 |
| 12-HETE-\(d_6\)     | 78.7 ± 11.9 | 99.2 ± 8.0  | 126.1 ± 14.8 | 104.2 ± 8.5 | 121.7 ± 21.2 | 138.7 ± 14.0 |

\(^a\)Recovery were performed by comparing the extracted samples with unextracted standards that represent 100%.

\(^b\)Matrix effect expressed as the ratio of the mean peak area of an analyte spiked post-extraction to the mean peak area of the same analyte standards multiplied by 100. A value of >100% indicates ionization enhancement, and a value of < 100% indicates ionization suppression.

\(^c\)High-concentration Quality Control.

\(^d\)Medium-concentration Quality Control.

\(^e\)Low-concentration Quality Control. \(n = 6\), injected in triplicate.

### Recovery and Matrix Effects.

To determine the extraction efficiency of SPE for the analytes in the biological matrix studied, recovery was assessed by means of the IS solutions. Plasma and lung tissue samples, as well as the culture medium, were spiked with IS solution before SPE extraction. Recovery was determined by comparing the analytical response of the IS for the extracted samples at three concentrations (LQC, MQC, and HQC) to the IS that had not been extracted and that had been prepared in solvent (MeOH/H\(_2\)O, 7:3, \(v/v\)). Recovery was expressed as the percentage of the expected value, as calculated by the average area ratio between the extraction samples and the standard samples. The extent of recovery of the IS should be consistent, precise, and reproducible. Matrix-dependent recovery was established using different matrices, such as lung tissue homogenate, plasma, and culture medium; these were spiked with IS at three concentrations (LQC, MQC, and HQC) after SPE extraction. The peak areas after extraction were compared with the corresponding peak areas of the pure IS solutions. Samples for matrix effect and recovery studies were prepared in quintuplicate, and a CV of ≤15% was used as the accepted criteria for quality control. The recovery and matrix effect percentages for lung tissue, plasma, and culture cell are shown in Tables 4, 5 and 6, and respectively full results on file. Values obtained for each replicate of matrix effects and recovery samples (lung tissue, plasma and culture medium) (xlsx) and Values obtained for each replicate of matrix effects and recovery samples (lung tissue, plasma and culture medium) (pdf)\(^f\), Data Citation 1.

### Data records

**Data Record 1**–(Metabolite standard details (docx) and Metabolite standard details (pdf), Data Citation 1) gives for each metabolite standard purchase from Cayman Chemical all details, such as molecular weight, molecular formula, CAS number and supplied solution, as well biological information, such as fatty acid precursor, biosynthetic pathway and chemical class. The file include Internal lipid standards, with matching natural lipid (MWS), used in for extraction control, matrix effect and recovery experiments, also standards for calibration curves (MWS) in UHPLC-MS/MS and for quantitative calculation of lipid species.

**Data Record 2**–(Tandem mass spectral libraries in high resolution (HR-MS/MS) for eicosanoid panel (pdf), Data Citation 1), described tandem mass spectral libraries in high resolution (HR-MS/MS) for eicosanoid panel. Also, the file provided information about collision energy, molecular formula and common name, as well theoretical mass-to-charge ratio (m/z) of precursors and fragments ions (illustrated by chemical structures). Experimental spectrum showed numerical values that correspond to the m/z (x-axis) and relative intensity (y-axis) for each detected metabolite. The precursors (red) and fragments (blue) ions used for quantification analysis were highlight in the spectra.
Table 5. Mean recovery and matrix effect of the lipid mediator targets using SPE extraction in HQC, MQC, and LQC levels of IS in lung samples. aRecovery were performed by comparing the extracted samples with unextracted standards that represent 100%. bMatrix effect expressed as the ratio of the mean peak area of an analyte spiked post-extraction to the mean peak area of the same analyte standards multiplied by 100. A value of >100% indicates ionization enhancement, and a value of < 100% indicates ionization suppression. cHigh-concentration Quality Control. dMedium-concentration Quality Control. eLow-concentration Quality Control. *n = 6, injected in triplicate.

| Internal Standard | Recovery Rate (± SD)a | Matrix Effect (± SD)b |
|-------------------|----------------------|----------------------|
|                   | HQCc | MQCc | LQCe | HQCc | MQCc | LQCe |
| 15-HETE-d6        | 70.4 ± 5.7 | 86.8 ± 16.9 | 92.9 ± 11.9 | 94.1 ± 2.7 | 145.1 ± 17.4 | 117.5 ± 13.9 |
| TXB2-d6           | 54.8 ± 3.7 | 67.2 ± 14.6 | 64.1 ± 10.1 | 82.4 ± 3.5 | 108.6 ± 19.8 | 104.4 ± 11.6 |
| 6-keto-PGF1α-d6   | 50.3 ± 3.0 | 68.6 ± 16.2 | 68.0 ± 3.0 | 78.7 ± 6.5 | 128.6 ± 45.3 | 167.7 ± 54.9 |
| PGJ2-d6           | 53.9 ± 5.1 | 64.6 ± 11.8 | 60.5 ± 16.2 | 77.1 ± 4.1 | 105.0 ± 12.6 | 95.8 ± 8.7 |
| PGD2-d6           | 56.3 ± 5.6 | 65.8 ± 14.9 | 40.6 ± 12.9 | 79.1 ± 5.8 | 102.1 ± 16.1 | 82.1 ± 15.3 |
| LXA4-d6           | 57.0 ± 6.3 | 69.2 ± 12.2 | 66.5 ± 12.9 | 74.9 ± 3.2 | 103.7 ± 10.7 | 106.9 ± 16.0 |
| RuD1-d6           | 53.9 ± 6.4 | 63.6 ± 12.5 | 68.4 ± 15.0 | 74.9 ± 3.2 | 103.7 ± 10.7 | 106.9 ± 16.0 |
| PGI2-d6           | 48.7 ± 3.8 | 58.8 ± 13.8 | 88.6 ± 7.0 | 73.5 ± 4.7 | 99.8 ± 12.4 | 135.5 ± 14.4 |
| 12-iso-LTB4-d6    | 53.8 ± 3.8 | 64.1 ± 13.6 | 68.2 ± 15.1 | 72.0 ± 2.9 | 99.9 ± 14.3 | 91.4 ± 12.2 |
| LTE4-d6           | 45.2 ± 3.9 | 59.6 ± 17.2 | 144.2 ± 35.1 | 82.3 ± 6.5 | 118.8 ± 17.9 | 301.8 ± 52.5 |
| AA-d5             | 51.8 ± 16.4 | 17.0 ± 5.8 | 12.7 ± 4.3 | 66.5 ± 12.9 | 24.7 ± 4.9 | 23.7 ± 5.6 |
| 15-deoxy-d-12,14-PGJ2-d6 | 52.1 ± 4.2 | 70.9 ± 10.4 | 79.8 ± 13.2 | 69.2 ± 5.2 | 121.1 ± 10.1 | 105.4 ± 7.4 |
| 5-HETE-d6         | 58.0 ± 5.2 | 65.6 ± 11.3 | 62.4 ± 12.6 | 87.5 ± 6.4 | 104.7 ± 16.2 | 91.4 ± 9.3 |
| 5-iso-ETE-d7      | 67.9 ± 9.0 | 61.5 ± 8.1 | 50.4 ± 26.6 | 89.9 ± 6.6 | 111.0 ± 16.9 | 83.7 ± 12.7 |
| 12-HETE-d4        | 61.7 ± 5.6 | 74.3 ± 13.4 | 75.8 ± 12.0 | 78.9 ± 4.1 | 118.4 ± 10.9 | 94.9 ± 14.0 |

Table 6. Mean recovery and matrix effect of the lipid mediator targets using SPE extraction in HQC, MQC, and LQC levels of IS in cell culture medium. aRecovery were performed by comparing the extracted samples with unextracted standards that represent 100%. bMatrix effect expressed as the ratio of the mean peak area of an analyte spiked post-extraction to the mean peak area of the same analyte standards multiplied by 100. A value of >100% indicates ionization enhancement, and a value of < 100% indicates ionization suppression. cHigh-concentration Quality Control. dMedium-concentration Quality Control. eLow-concentration Quality Control. *n = 6, injected in triplicate.

| Internal Standard | Recovery Rate (± SD)a | Matrix Effect (± SD)b |
|-------------------|----------------------|----------------------|
|                   | HQCc | MQCc | LQCe | HQCc | MQCc | LQCe |
| 15-HETE-d6        | 42.7 ± 5.8 | 56.6 ± 12.8 | 52.3 ± 8.9 | 79.9 ± 12.9 | 114.1 ± 16.0 | 118.9 ± 9.8 |
| TXB2-d6           | 58.3 ± 6.4 | 58.9 ± 2.9 | 58.6 ± 5.2 | 54.2 ± 6.9 | 74.59 ± 4.9 | 62.0 ± 10.8 |
| 6-keto-PGF1α-d6   | 34.4 ± 4.5 | 74.9 ± 17.1 | 57.6 ± 7.1 | 95.3 ± 7.7 | 126.9 ± 33.3 | 120.4 ± 9.9 |
| PGJ2-d6           | 56.7 ± 3.2 | 78.8 ± 5.0 | 71.5 ± 4.9 | 64.2 ± 12.3 | 94.3 ± 3.8 | 80.1 ± 10.2 |
| PGI2-d6           | 73.1 ± 7.1 | 82.5 ± 9.0 | 74.2 ± 19.2 | 78.5 ± 17.1 | 104.2 ± 21.6 | 79.9 ± 20.9 |
| LXA4-d6           | 44.5 ± 5.7 | 51.5 ± 4.3 | 57.6 ± 7.1 | 48.6 ± 6.6 | 61.4 ± 4.9 | 61.4 ± 4.9 |
| RuD1-d6           | 49.9 ± 10.1 | 51.9 ± 3.7 | 57.6 ± 7.1 | 49.2 ± 9.4 | 67.4 ± 5.6 | 63.4 ± 15.4 |
| PGI2-d6           | 49.3 ± 8.5 | 54.8 ± 3.9 | 56.7 ± 10.8 | 47.3 ± 8.7 | 69.4 ± 2.9 | 73.6 ± 9.3 |
| 12-iso-LTB4-d6    | 61.1 ± 8.2 | 59.8 ± 3.2 | 63.0 ± 14.4 | 66.2 ± 11.9 | 87.9 ± 5.4 | 78.4 ± 10.1 |
| LTE4-d6           | 73.3 ± 36.6 | 34.3 ± 5.1 | 80.8 ± 28.1 | 68.5 ± 15.7 | 93.6 ± 8.5 | 114.5 ± 45.2 |
| AA-d5             | 44.3 ± 8.4 | 12.3 ± 7.1 | 67.1 ± 12.5 | 47.2 ± 5.3 | 35.4 ± 3.5 | 38.7 ± 6.5 |
| 15-deoxy-d-12,14-PGJ2-d6 | 40.2 ± 5.3 | 43.8 ± 5.0 | 59.6 ± 5.4 | 54.0 ± 9.5 | 77.3 ± 5.6 | 83.3 ± 5.3 |
| 5-HETE-d6         | 37.1 ± 4.2 | 46.3 ± 12.1 | 35.6 ± 13.5 | 79.1 ± 14.3 | 103.9 ± 5.6 | 94.9 ± 8.3 |
| 5-iso-ETE-d7      | 41.5 ± 9.2 | 36.3 ± 13.2 | 66.6 ± 10.0 | 83.3 ± 16.0 | 96.4 ± 10.2 | 81.2 ± 14.3 |
| 12-HETE-d4        | 43.5 ± 6.8 | 42.4 ± 9.6 | 42.3 ± 15.2 | 71.5 ± 12.5 | 88.9 ± 4.6 | 109.5 ± 9.8 |
Data Record 3—(Retention time for lipid mediators panel under different UHPLC-MS/MS chromatography conditions (pdf), Data Citation 1) demonstrated the retention time for lipid mediators panel under different UHPLC-MS/MS chromatography conditions. The chromatography parameters differ by mobile phase pH (3.8, 5.8 and 6.4). Using a elution method of UHPLC with a reversed-phase column (C18), a mixture of commercially lipid mediators standards elutes according to their polarity, number of double bonds and chain length, allowing the separation of most isomeric and isobaric species (represented by blue and red lines respectively) dependent of pH and according to identifying fragmentation patterns described in file 'Tandem mass spectral libraries in high resolution (HR-MS/MS) for eicosanoid panel (pdf), Data Citation 1). An asterisk represented the non-detect metabolites.

Data Record 4—(Linear regression equation, correlation coefficient (r²) and linear range of concentration for each analyte (xlsx) and Linear regression equation, correlation coefficient (r²) and linear range of concentration for each analyte (pdf), Data Citation 1), showed the linear regression graphics. MS files were processed for each metabolite quantified in MRMHR (HR-MS/MS) method with mobile phase pH 5.8. The graphics describe the linear regression equation, correlation coefficient (r²) and linear range of concentration for each analyte assessed by means of 10-point standard curves prepared for five replicate samples. Also the retention time were described in this file.

Data Record 5—(Values obtained for each replicate of quality control (HQC, MQC and LQC) assay (xlsx) and Values obtained for each replicate of quality control (HQC, MQC and LQC) assay (pdf), Data Citation 1) presented all values obtained for each replicate of quality control (HQC, MQC and LQC) assay. The individual values from quality control assay were used to determine the average (n = 5) of precision and accuracy of the MRMHR method presented in Table 3 (available online only).

Data Record 6—(Values obtained for each replicate of matrix effects and recovery samples (lung tissue, plasma and culture medium) (xlsx) and Values obtained for each replicate of matrix effects and recovery samples (lung tissue, plasma and culture medium) (pdf), Data Citation 1) demonstrated all values obtained for each replicate of matrix effects and recovery samples (lung tissue, plasma and culture medium). In these experiments the analytes corresponded to internal standards (IS- deuterated compounds) for eliminate the interference of matrix endogenous metabolites. The individual values were used to determine the average (n = 6) of matrix effect and recovery of the extraction method presented in Tables 4, 5 and 6.

Data Record 7—All data files from TripleTOF 5600+ (.wiff) have been deposited to the MetaboLights metabolomics repository19 (Data Citation 1). Instrument RAW data files (121 files) contain total ion current, spectra and metadata for each replicate. This data set was used to provide the linear range of concentration, retention time, LLOQ (summarized in file 'Linear regression equation, correlation coefficient (r²) and linear range of concentration for each analyte (xlsx) and Linear regression equation, correlation coefficient (r²) and linear range of concentration for each analyte (pdf), Data Citation 1), intra- and interday precision and accuracy (summarized in file 'Values obtained for each replicate of quality control (HQC, MQC and LQC) assay (xlsx) and Values obtained for each replicate of quality control (HQC, MQC and LQC) assay (pdf)', Data Citation 1). Also, the MTBLS641 archive (.zip) contains others files: 'i_Investigation.txt' that provided summary information and project descriptors; 'a_study_id_metabolite_profiling_mass_spectrometry-1.txt', 'a_study_id_metabolite_profiling_mass_spectrometry-1.txt' (Data Citation 2) which afford the contextual data for the MS analyses from data acquisition through to the multiple processing steps. All raw data files were uploaded using an ISA-tab format.

Technical validation
MRMHR channel determination and selectivity
Technical validation of the UHPLC-MS/MS method for the comprehensive analysis of the lipid mediators on a high-resolution mass spectrometer (TripleTOF5600+ system) in MRMHR mode was carried out before biological sample measurements were taken; this was previously described in more detail14. To this end, the pre-defined MRM transitions (pairs of precursor/fragment ions) in a low-resolution mass spectrometer typically used for routine lipidomics analysis in our lab16,17 were optimized for high-resolution mass spectra (“Tandem mass spectral libraries in high resolution (HR-MS/MS) for eicosanoid panel (pdf)”, Data Citation 1). The processing software, MultiQuant, was capable of generating these MRM-like channels post-acquisition, thus allowing that each transition was create in accordance with the MS/MS data obtained during analysis. This meant that there was no need for pre-selection of the candidate lipid fragments in the mass analyzer. In addition, time-resolved mass measurements enabled proper alignment of the detected fragment ions with their respective precursor ions for accurate identification. Table 1 (available online only) shows the optimized mass spectrometric parameters for the detection of compounds in the MRMHR mode, which were done using the most sensitive and most selective MRMHR transitions. These parameters allowed us to accurately discern between the individual lipid mediators and the background interferences for each analyte.

Influence of pH on elution behavior
Developing an LC-MS elution methodology that would efficiently assay all of the target analytes was filled with challenges, especially when it came to finding the right balance between the method’s sensitivity and chromatographic resolution. To compound matters, chromatographic separation was required for
isomers that were not differentiated through MRMHR transitions (i.e., those which had identical CAD spectra); these included the pairs of lipids PGE2/PGD2, LTD4/11-trans-LTD4, and LTB4/6-trans-LTB4. This observation forced us to look at the effect of pH on the elution behavior of the analytes.

As previously suggested, the elution behavior of a mixture of all compounds was first examined in the mobile phase at pH of 3.8 (ref. 16). However, since most of the standard lipids were differentiated either through MRMHR transition or their respective chromatographic retention times, this analytical method could not be used to quantify lipids like LTC4, AA, and EPA under these conditions. As such, no chromatographic peaks were detected for the aforementioned lipids. Moreover, the shape of the peak obtained for 6-keto-PGF1α was too broad. Given this, a range of pH levels was evaluated using the developed LC method in order to optimize sensitivity and selectivity. At pH 6.4, the analytical response for all lipid species was improved, including those for LTC4, AA, and EPA.

**Figure 3. Representative MRMHR eicosanoid chromatograms.** A mixture of commercially available eicosanoid standards elutes in C18 column according to different experimental conditions of phase A elution: (a) pH 3.8, (b) pH 5.8 and (c) pH 6.4. The chromatograms showing separation of pairs of isobaric species with identical fragmentation patterns (LTD4 and 11-trans-LTD4, PGE2 and PGD2, LTB4 and 6-trans-LTB4), also showing elution behavior of critical metabolites (LTC4, 6-keto-PGF1α, EPA, and AA). The absence of metabolites peaks was represented by an asterisk (*). The retention time of all lipid mediators quantified in MRMHR method, for different pH elution conditions, were available in file ‘Retention time for lipid mediators panel under different UHPLC-MS/MS chromatography conditions (pdf)’, Data Citation 1.
the lipid mediators that contained free carboxyl moieties were abundant and present as the negative [M – H] species. On the other hand, isomers with identical MRMs were co-eluted with the same retention time under these LC conditions. Using a mobile phase with pH of 5.8 could resolve these critical co-elution isomers while still enabling the detection of LTC₄, AA, and EPA. As evidence of this, it was noted that the peak shape of 6-keto-PGF₁α was improved under these conditions (Fig. 3).

Validity of the UHPLC-MS/MS method

Validation assays of the UHPLC-MS/MS method were attained through analysis of non-endogenous lipid mediator standards diluted to various concentrations. Reference standards were quantified as described in the data processing section. UHPLC-MS/MS method validation showed that accuracy values ranged from 0.3 to 15.0% for LOQ, from 0.4 to 13.7% for MOQ, and from 0.2 to 10.1% for HOQ on the calibration curve. Method precision for five technical replicates was between 2.3 and 14.4% for LOQ, 1.2 and 14.8% for MOQ, and between 2.1 and 14.9% for HOQ. These were in accordance with FDA guidelines for targeted LC-MS methods, which proposed a maximum CV of 15%. The dynamic range of the method covered more than four orders of magnitude and was between 0.02 and 500 ng.mL⁻¹. Linear fits for the calibration curves obtained in this concentration range yielded correlation coefficients (r²) of up to 0.99 (Table 2), which was acceptable for a targeted lipidomics method. Furthermore, there was a measure of comparability in the IS across the whole range of retention times for equivalent mixtures of the IS and MWS. The retention time of the IS and their correspondent non-deuterated lipid standards matched exactly during the course of the LC runs. Table 2 shows the linearity and LLOQ for each lipid mediator as determined at pH 5.8 alongside their corresponding chromatographic retention time. Precision and accuracy values are shown in Table 3 (available online only). Matrix effect and recoveries were determined for the deuterated analytes since it was thought that the biological samples studied contained endogenous lipid mediators that would interfere with the experiment. The recoveries and matrix effect data obtained for plasma, lung tissue and culture medium samples are given in Tables 4, 5 and 6, respectively. For PGE₂-d₆, the recovery of extraction process in plasma was 74.7% when the HQC concentration was used for spiking the biological fluid; 91.7% for MHC and 94.6% for MQC. For lung tissue we observed a recovery of 53.9% for HQC, 64.6% for MQC and 60.5% for LQC. In cell culture the recovery was 56.6% for HQC, 78.8% for MQC and 70.5% for LC. Thus, considering all analytes the recoveries ranged from 27 to 136% for the plasma samples, from 12 to 92% for the lung tissues samples, and 34 to 79% for the culture medium, approximately. The matrix effect results observed for the samples of plasma, lung tissue and culture medium ranged, approximately, from 26 to 138%, from 23 to 145%, and from 38 to 126%, respectively.

Usage Notes

The eicosanoids data file acquired by the ‘TripleTOF5600+’ instrument are publicly available in ‘.wiff’ file format at European Bioinformatics Institute’s MetaboLights repository (Data Citation 2). After download, the data can be available for analysis by different software packages. For this, Sciex (Foster, CA, USA) offer a tool called MS Data Converter, free download at https://sciex.com/software-support/software-downloads. Output can be a simple MGF peak list or an mzML file containing a fairly complete representation of the raw data. Choosing an exact translation of the instrument recorded or converts to a processed version, reducing the data down to peak lists. The mzML format is the single XML standard mass spectrometry format that was created by the merger of the older mzXML and mzData formats. However, the conversion of data in mzML output files result in larger files than the original raw data. To require mzXML format than mzML, it was recommended to first convert to mzML using the MS Data Converter and then use a public tool such as msconvert in the ProteoWizard package to convert mzML to mzXML, available in http://proteowizard.sourceforge.net/downloads.shtml. Informatics challenge associated with analysing MS data required a wide variety of data file formats to encode the complex data types associated with MS workflows. Each company has developed individual formats of its own and continually extends them, as emerging instrumentation requires new features. Sciex (Foster, CA, USA) instruments are saved as files with the ‘.wiff’ extension. These files might sometimes contain all information in a run or alternatively might contain only metadata and be paired with a file having ‘.wiffscan’ extension that contains the spectra. To solution this, the new format mzML were created that would include the best features from both mzXML and mzData and eventually replace them. Likely the most common text format is the Mascot Generic Format (MGF) file. It is still common convert the binary mass spectrometer output files into simple text files containing only the MS/MS spectra. Finally, targeted and quantitative data obtained by UHPLC-MS/MS method technology in this data descriptor can be automatically processed and rapid evaluation. Also, the data sets documented her will allow draught subsets of lipid species suggestive for a physiological and metabolic condition.

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**Data Citations**

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**Author Contributions**

C.A.S. and A.P.F.P. conducted mass spectrometric assays, bioinformatic analysis, data evaluation, and validation, prepared the schematic figures, raw data storage in repository and wrote the manuscript; T.P. conducted data analysis; A.F.G.M. and C.F. followed up on the experimental results; L.A.B.M. organized and structured the datasets, as well as revised the manuscript and L.H.F. supervised the work and revised the manuscript and data repository storage. All co-authors contributed to the final manuscript.

**Additional Information**

Tables 1 and 3 are available only in the online version of this paper.

**Competing interests:** The authors declare no competing interests.

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