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Oxylipins represent a large network of bioactive lipid mediators (Figure S1), possessing both physiological and pathological roles. These lipids derive from oxidation of polyunsaturated fatty acids (PUFA), either enzymatically by cyclooxygenases (COXs), lipoxygenases (LOXs) and cytochrome P450s, or non-enzymatically via free-radical peroxidation processes.\textsuperscript{1,2} Within the class of oxylipins, several isomers can be found which exert distinct biological functions and have a different biochemical origin. For example, the six isoforms of HETEs can derive from oxidation of arachidonic acid either by LOXs (5-, 12- and 15-HETE) or COXs (11- and 15-HETE), while 8- and 9-HETE derive mostly from non-enzymatic autoxidation. Hence, identification of the correct isomers is fundamental to fully understand both their path of generation and function. Oxidation products of C20 PUFA, such as arachidonic acid are commonly known as eicosanoids, which are generated in cells and tissues in response to physiological and pathological changes\textsuperscript{2} and play crucial roles in mediating inflammation and homeostasis as well as diseases such as cancer, atherosclerosis and Alzheimer’s.\textsuperscript{3,5} Oxylipins are either released by cells/tissues to act as free mediators, or re-esterified into phospholipids and integrated into the cell membrane of platelets and leukocytes where they can interact with coagulation factors or act pro- or anti-inflammatory.\textsuperscript{6} Oxylipins are found in low nanomolar concentrations in biological samples, therefore their analysis requires highly selective and sensitive methods. To achieve this, ultra-high performance liquid chromatography (UHPLC) using reversed-phase columns in combination with triple quadrupole mass spectrometers coupled with negative electrospray ionization are commonly used.\textsuperscript{7,8} Although this methodology allows both sensitive and selective analysis, targeted approaches are time-consuming in their development and will miss lipids that are not specifically being analyzed. Ion mobility coupled with high-resolution mass spectrometry (IM-MS) can be used to achieve more specificity in untargeted approaches, allowing measurements of both accurate mass and collision cross-section (CCS) values of the ions to provide greater certainty in terms of identification. In recent years, IM-MS has developed as an emerging technique in lipidomics,\textsuperscript{9,10} however, oxylipins analysis by IM-MS has been limited to some eicosanoids using travelling-wave IM.\textsuperscript{11,12} Leukotrienes and protecins using differential IM,\textsuperscript{13} and some oxylipins using drift tube IM (DTIM-MS).\textsuperscript{14} In this study we describe a comprehensive analytical and computational workflow for the analysis of oxylipins and fatty acids (FAs) using UHPLC-DTIM-QTOF, in which size and shape of the ions affect their mobility when travelling through a drift tube filled
with low pressure nitrogen gas, allowing their separation and the calculation of their CCS values for nitrogen ($^{19}$CCS$_{N2}$). Annotation and identification of oxylipins and FAs was enhanced through an extension of the KniMet pipeline for the processing of metabolomics data with a combined annotation method comprising the LIPID MAPS Structure Database (LMSD) and an internally generated library containing 47 unique lipids. The UHPLC-DTIM-MS method presented in this study also allows quantification of oxylipins in biological samples, and its combination with data-independent acquisition enables lipid identification based on drift time alignment between diagnostic product ions and their precursors. The computational pipeline as well as the data obtained for the 47 lipid standards is publicly available at https://github.com/sonial/KniMet and could contribute to the creation and expansion of CCS databases. Together, this method can be used for both discovery and quantification of oxylipins and FAs in biological samples. The approach is here demonstrated on human thrombin activated platelets and Salmonella enterica serovar Typhimurium infected murine bone-marrow derived macrophages (BMDMs).

**MATERIALS AND METHODS**

**Materials.** Lipid standards were purchased from Santa Cruz Biotechnology (Heidelberg, Germany) or Cayman Chemical (supplied by Cambridge Bioscience, Cambridge, UK). All solvents were Chromasolv LC-MS grade from Honeywell. Ultrapure LPS from Escherichia coli O111:B4 was from InvivoGen. All other reagents were purchased from Sigma Aldrich (Merck KGaA, Darmstadt, Germany) unless otherwise stated.

**Mice.** Wild type C57BL/6 mice were obtained from Charles River, UK. All animals were housed under specific pathogen-free conditions and all work involving live animals complied with the University of Cambridge Ethics Committee regulations under Home Office Project License number 80/2572.

**Cell isolation and culture.** BMDMs were retrieved from the bone marrow of legs of wild type C57BL/6 mice and cultured in BMDM growth media (Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum, 20% L929 conditioned media and 5 mM L-glutamine). After centrifugation, both isolated cells and medium were divided into 2 groups to subject only one of them to infection with a solution of $S$. Typhimurium strain SL1344 in BMDM growth media with conditioned media and 5 mM L-glutamine. After a 5 min at 4°C, the column was re-equilibrated for 2.5 min at 30% solvent B. A flow rate of 0.3 mL/min (50% solvent B) was used for flow injection experiments. Lipids were analyzed in negative IM-QTOF mode in the $m/z$ range of 100-1000 and maximum drift time (DT) of 60 ms using nitrogen as drift gas, a trap fill time of 25 ms, trap release time of 0.5 ms and acquisition rate of 1 IM frame/s. An Agilent tuning solution was injected both before the analysis to tune the instrument in the $m/z$ range 100-1700, and every 10 samples to perform $^{19}$CCS$_{N2}$ re-calibration. An Agilent reference mix for mass re-calibration was continuously injected alongside the samples using the secondary sprayer. The Dual Agilent Jet Stream electrospray ionization source was operated at gas temperature of 325°C, drying gas 5 L/min, sheath gas temperature of 275°C and a sheath gas flow of 12 L/min. A drift gas upgrade kit maintained both drift tube and trap funnel pressure constant at 3.9 and 3.8 Torr respectively, while drift tube temperature was found to be stable at 23.5°C ± 0.3°C across all acquisition runs. Fragmentation with alternating frames was used for data-independent acquisition (IM-MS/MS) of pooled samples (QCs), with the collision energy of frame two fixed at 30V.

**Creation of an Internal Database of Oxylipins.** An internal database containing accurate mass, retention time (RT) and $^{19}$CCS$_{N2}$ values was obtained for 47 lipid standards previously acquired in triplicates across different days using the instrumental, analytical and pre-processing method described. This data was further processed with an in-house pipeline based on the KNIME Analytics Platform to build a database containing [M-H], [M-H+CH$_3$CO$_2$Na]$^-$, [2M-2H+Na]$^-$ and [2M-H]$^-$ species (Table S1 and https://github.com/sonial/KniMet). First, a list containing lipid standard names, exact molecular weight, expected RT (manually inferred from previous acquisitions), SMILES, formula and InChI was created with the aid of the ChemAxon and RDKit KNIME nodes. This list was used to annotate the individual acquisitions based on accurate mass and RT match (error ranges 10 ppm and 0.15 min, respectively) with an adapted version of the KniMet functionality “Feature Annotation”, which was modified to take into account also RT range.
A subsequent cleaning of the annotations consisted of keeping only the features annotated as different adducts of the same molecule within a RT range of ± 0.01 min from a [M-H]+ conformer. This allows one to keep only co-eluting adducts and eliminate erroneous annotations derived from the large RT range required for the annotation step. The results obtained for each different acquisition were then merged in the database if they were present in at least two out of three injections with a relative standard deviation (RSD) smaller than 1% for m/z and 1.5% for both RT and DTCCS\(_N2\). This results in five peaks on average per lipid, with some of them deriving from the same adduct suggesting either different conformations or more complex ion arrangements (discussed in the section “Ion mobility allows identification of lipid clusters”).

**Data processing.** Data pre-processing, including mass and DTCCS\(_N2\) re-calibration and feature finding, was carried out using the packages IM-MS Reprocessor, IM-MS Browser and Mass Profiler from the MassHunter Suite (version B.08.00, Agilent Technologies, Santa Clara, USA). In particular, CCS calibration was performed in batches containing 10 samples and one calibrant, where the slope and intercept (β and TFIx in IM-MS Browser) values obtained for the DT correction of the calibrant were applied to samples for calculation of DTCCS\(_N2\) (please refer to \(^{24}\) and \(^{25}\) for more details of CCS calibration in DTIM). Stability of the calibration was assessed by calculating the CCS for a dummy feature (DT = 28 and m/z = 313, which are the average values found in our internal database) using the β and TFIx values obtained from the several calibrant infusions, with RSD equal 0.097% for the 4 calibrants infused in BMDM and 0.103% for the 5 calibrants infused in the platelets study. Feature finding was performed for signals having a minimum ion count of 100 using the maximum ion volume as a measure of abundance and an isotopic model designed for common organic molecules without halogen atoms. Moreover, only species with charge state = 1-2 were extracted, including single ion features with charge state = 1. Features were aligned if they fell within the confidence intervals of 10 ppm ± 3 mDa in m/z and in 0.3 min in RT. Furthermore, an m/z range (199-1000) was selected to include monomers, dimers and adducts. A final filter obtained from Mass Profiler, provides an indication about data based on Q-Score (greater than 60 on a scale of 0-100; value selected to include monomers, dimers and adducts. A final filter was performed using the Agilent MassHunter Quantitative Processing software. Although the input files contained IM data, the software simply ignores the IM dimension and quantities based on the chromatographic peak. Lipid levels were normalized to internal standard levels. On the other hand, an adaptation of the KniMet pipeline for MS-based metabolomics data processing\(^{27}\) was used to perform profiling of oxylipins. More specifically, features were retained if they were found in at least 70% of the samples belonging to a class with an average abundance higher than 5000. This first filtering of noise was finalized by the removal of all features whose average intensity in samples was less than twice the average intensity in uninfected medium samples in the BMDM experiment, while periodic injections of a dilution series of quality control samples (QCs, obtained pooling all samples analyzed) was used in the platelets experiment to remove features whose intensity does not vary linearly with concentration of the sample (node “QC dilution series-based Features Filtering” added to KniMet). The remaining features were then subjected to missing value imputation using k-nearest neighbor for multivariate statistical analysis, while replacement of missing values with half of the minimum value found in each sample was used for univariate analysis to avoid a possible change of the data distribution.\(^{26}\) Finally, signals were normalized using probabilistic quotient normalization based on all samples\(^{27}\) and logarithmic transformed. Identification of oxylipins and FAs was performed by accurate mass, RT and DTCCS\(_N2\) match with the internal database, with error ranges of 10 ppm for mass accuracy, 0.15 min for RT difference and 2% RSD for DTCCS\(_N2\). Moreover, lipid-like flags were added: features presenting m/z and RT within the aforementioned ranges, but different DTCCS\(_N2\) values from those present in the database, were flagged as possible new conformers, while features with m/z and DTCCS\(_N2\) within these ranges but different RT were flagged as possible new compounds. Annotation of lipids not present in our internal database was performed by comparison with a subset of the LMSD (version updated in 12/2016)\(^{16}\) containing only lipids belonging to the fatty acyl class (to which oxylipins belong).\(^{28}\) In both annotation and identification steps, species whose RT is further than 0.01 min from the relative [M-H]+ were removed to eliminate wrongly annotated dimers and adducts. Both annotations were performed by adapting the “Feature Annotation” functionality of KniMet. In particular, “Feature Annotation from internal IMS DB” was added to KniMet to allow annotation based also on DTCCS\(_N2\) match and, if available, RT, to flag lipid-like compounds, and to remove dimers with no monomers in a given RT range. Similarly, the “Feature Annotation - Internal2external” functionality performs annotation on a specific class of LIPID MAPS and removes dimers with no monomers in a given RT range (all created KniMet nodes available at https://github.com/sonial/KniMet).

Principal Component Analysis (PCA) was performed using SIMCA (Umetrics, Sweden), while heatmap representations were created using the function heatmap2, part of the gplots\(^{29}\) library of R,\(^{30}\) with default parameters for both hierarchical clustering and dissimilarity calculation.

**RESULTS AND DISCUSSION**

**Acquisition and data processing workflow for oxylipin and fatty acid analysis.** Oxylipins are a class of lipids presenting strong structural similarities, but their polarity can vary depending on number of oxygen atoms, position of hydroxyl groups or number of double bonds. Therefore, a reversed-phase gradient method\(^{28}\) was used to separate oxylipins and FAs in the first dimension (Figure S2) prior to further separation by IM and accurate mass detection by QTOF MS. To profile the lipid content, features picked by Mass Profiler were processed using KniMet to perform data normalization, feature filtering, identification and annotation. For feature identification, an in-house lipid library (Table S1) was used. Features not present in the library were putatively annotated based on accurate mass using LMSD. In addition to lipid discovery, lipids were quantified using the Quantitative Analysis software if a standard curve was analyzed alongside. To confirm lipid identities, either a QC or a known sample containing oxylipins (if concentration in the QC was too low) was analyzed using data-independent acquisition, which does not require the long development time of targeted methods. Due to fragmentation taking place after the drift tube, product ions time-align to their precursors, allowing identification of diagnostic product ions\(^{9}\) and conferring more confidence in oxylipin identification.
Ion mobility allows identification of lipid clusters. DTIM separation can be visualized in drift spectra, in which counts are plotted versus DT (ms), and driftograms, where instead DT is plotted against m/z. Multiple drift peaks were observed for all lipids analyzed in this study, with one of these peaks being dominant in intensity and hence corresponding to the most stable conformer in the gas phase. DTIM-MS analysis of the most abundant [M-H] (referred to as species 1) conformer of oxylipins and FAs shows that they occupy both tight $^{17}$CNS$_2$ and mass ranges ($CCS = 174-221$ Å$^2$, m/z = 296-496 for the former, $CCS = 158-185$ Å$^2$, m/z = 200-328 for the latter). However, the driftograms produced by most lipids suggest the presence of multiple conformations, with several features having identical RT and m/z but different DT. Among these, low abundant species drifting in close proximity to the dominant peak were found, which could derive from either different isomers (geometric- or stereo-isomers) or different conformers of species 1. In contrast, the second most intense signal corresponds to a feature (species 2) drifting about 10 ms later than species 1. Although differences in $^{17}$CNS$_2$ values, corresponding to different conformations, have been previously described for FAs, the DT differences observed between species 1 and 2 (Table S1) may be too great to correspond to different conformers. The appearance of species 2 depends on the concentration of both lipid and organic solvent used, which was tested under six different conditions for PGE$_2$ (Figure 1A), arachidonic acid, 12-HETE and 5(6)-EET (Figure S3). Although the intensity of species 2 increased with an increase in the organic solvent content, the relationship is not linear and may also depend on the polarity and solubility of the individual lipid (Figure S4). Along with [M-H] of species 2, co-drifting (± 0.5 ms) [2M-H] and [2M-2H+Na] ions were detected as well (Figure 1B). To identify both dimers and species 2, we undertook targeted fragmentation of all species, namely [M-H], [2M-H] (Figure 1C and D), [M-H+CH$_3$CO$_2$Na] [2M-2H+Na], [M-H-H$_2$O], [M-H-2H$_2$O]. This led to diagnostic product ions m/z 271.2064 and m/z 189.1289 (Figure S5), which DT-align (± 0.5 ms) with their precursor ions (Figure 1C-D). Comparable results were also obtained when applying data-independent fragmentation (Figure 1E-F), demonstrating that diagnostic product ions derive from monomers, adducts, and dimers. Therefore, we propose that species 2 are product ions derived from lipid dimers, which are formed before IM separation and partly dissociate after the IM cell prior to mass detection. This would explain their detection as m/z corresponding to [M-H]: species but co-drifting with lipid dimers. Indeed, using the Agilent 6560 DTIM-MS, ions are focussed by a high-pressure ion funnel and accumulated by an ion trap prior to be released into the drift tube. When ions leave the drift tube, they are refocussed by a rear ion funnel, followed by m/z detection in the QTOF. These focussing and trapping procedures could result in the formation of unstable ion clusters such as dimers and solvent clusters, which partly fall apart before m/z detection and cause the aforementioned peaks with different DTs. We tested this hypothesis by altering trap fill time (20 or 30 ms) and lowering trap funnel RF voltages (-150V, -100V, -76V), but the signals remained the same (Figure S6), suggesting that more complex ion arrangements may occur during analysis. Similarly, the described ion species were present using different source conditions, including higher temperatures (data not shown). In addition, [M-H+CH$_3$CO$_2$Na] species were observed, but these species drift independently from monomers and dimers (Figure 1E, Figure S3 B, D and F, red) due
to their different m/z. Moreover, using altering high/low fragmentation with DTIM it was observed that the [M-H-H₂O]⁻ and [M-H-2H₂O]⁻ species are not only products but also precursors ions, suggesting that some [M-H]⁺ precursors lose a water molecule before fragmentation (Figure 1G). Note, this may not be an in-source fragmentation as modifying source parameters did not remove these species (data not shown). Further discussions to identification of unspecific losses using DTIM can be found in Supporting Discussion. On the other hand, formation of sodium acetate adducts increases with increasing organic solvent content (Figure S4 C), as expected given the enhanced concentration of acetic acid in the solution. However, this does not apply to dimers (Figure S4 B and D), indicating that dimer formation occurs following ionization.

In addition to our observation using lipid standards and flow injection DTIM-MS, we also observed lipid dimers and solvent adducts using both LC-DTIM-MS analysis of lipid standards and biological samples and flow injection MS analysis of lipid standards using a triple quadrupole instrument in Q1 scan mode (Figure S7). Therefore, using LC-DTIM-MS allowed additional information regarding lipid cluster formation, which may also take place on other instruments but could potentially be overlooked.

The level of unsaturation changes the drift peak shape of fatty acids. In addition to oxylipins, free FAs can be analyzed using our method. As previously shown, 9,31 DT increases with both chain length and degree of saturation (Figure S8). The shape of the drift peak changes with the degree of saturation, as shown for a series of FAs with 18 carbon atoms in their chain and an increasing number of double bonds (Figure 2). The drift spectra of saturated C18:0, as well as the unsaturated species C18:1 C18:2, contain two peaks. In contrast, the drift spectra of poly-unsaturated C18:3 and C18:4 contain one tailing peak which possibly covers the equivalent to the second peak detected in C18:0, C18:1 and C18:2. Drift peak tailing suggests multiple molecular conformations of the ions, possibly due to the higher degree of unsaturation. This hypothesis finds confirmation in the recent work of Stow and co-workers, who described an increasing number of theoretical DTCCS₉₂ values for fatty acid conformers with different levels of unsaturation. 24 Presence of multiple conformers does not affect comparability of DTCCS₉₂ values between different studies and laboratories, as the most abundant ones (and therefore the most stable conformers) can be used for comparison. However, as isomers are expected to drift close to the original conformer, 31 peak tailing should be taken into consideration when investigating isomer formation, for example under certain biological conditions such as oxidative stress where non-enzymatic autoxidation may lead to the formation of isomers not normally observed.

Figure 2: Drift spectra of C18 fatty acids differing in the number of double bonds: the second peak gets included in the tail of the first (most abundant) peak with increasing number of double bonds in the acyl chain.

HETE-esterified phosphatidylcholines show a distinct mobility behavior. Eicosanoids not only act as free lipid mediators released by cells, but can also be re-esterified into phospholipids. 6 To explore the DTIM behavior of eicosanoid-esterified phospholipids, we analyzed 8-, 9-, 11-, 12-, and 15-HETE-esterified phosphatidylcholines (PC) using flow injection DTIM-MS. Separation of the different HETE-PCs by DTIM depends on the ion type analyzed, with [M+OAc]⁺ species presenting the greatest separation between structural isomers, followed by sodiated species, and finally protonated ions with the lowest drift separation (Error! Reference source not found., Figure 3). In addition, drift separation of acetate adducts demonstrated that a hydroxyl group close to the PC headgroup (such as in 8- and 9-HETE) results in faster mobility perhaps due to a smaller molecular size than that conferred by hydroxyl groups further away from the headgroup (12- and 15-HETE). The same difference in mobility, but to a smaller extent, can also be observed for the sodiated adducts. Therefore, we propose that both acetate and sodium interact with the hydroxyl group of the esterified HETE, resulting in a rather rigid structure of the molecule and distinct mobility behaviors. Although these phospholipids separate in LC, 32 and baseline separation of different isomers by DTIM could not be achieved, our findings are in agreement with previous observations 9 and show the potential to identify these isomeric lipids in complex samples based on DTCCS₉₂ values using a higher IM resolving power.

Quantification of oxylipins and fatty acids in S. Typhimurium activated murine BMDMs and thrombin activated human platelets. Eicosanoids are one of the least abundant lipid species found in cells and tissues, however, they are generated rapidly in response to acute cell activation during inflammatory processes or thrombus formation. As absolute quantities

Figure 3: Separation of eicosanoid-esterified phosphatidylcholines by DTIM-MS depends on both position of the hydroxyl group and adduct type. DTIM-MS analysis in both positive and negative mode of 8-, 9-, 11-, 12-, and 15-HETE esterified PCs (structures shown in D) shows enhanced separation between these isomers for [M+OAc]⁺ conformers (A) compared to sodium adducts (C), while [M+H]⁺ species (B) are not distinguishable in drift space.
of oxylipins is important in many studies, we developed a workflow which can be used for both profiling and quantification. A calibration curve containing increasing amounts of 16 lipid standards as well as steady amounts of 6 deuterated analogues (Table S2) was generated. The lower limit of quantification (LLOQ) was defined as the amount of lipid required to achieve a signal-to-noise ratio greater than 3. Nine deuterated standards (PGE$_2$-d$_4$, arachidonic acid-d$_4$, and 12-HETE-d$_3$) covering both oxylipins and FAs as well as different polarities, were spiked into platelet samples before lipid extraction to determine complete recovery rates, which were found to be 74% for PGE$_2$-d$_4$, 87% for arachidonic acid-d$_4$ and 91% for 12-HETE-d$_3$. This indicates low matrix and ionization effects on our approach. Lipid concentrations in samples were calculated using an external calibration curve and internal standards spiked into the samples before lipid extraction to account for both matrix effects and losses during the extraction procedure. The LLOQs ranged between 2 and 20 pg on column, equivalent to 0.6 and 11.3 nM, and regression coefficients $R^2 > 0.98$ were achieved (Figure S9, Table S2). To confirm lipid annotation from KniMet, both RTs and product ion spectra were compared to either lipid standards or spectra published on LIPID MAPS. Among the 16 compounds included in the calibration curve, seven were identified and quantified in BMDMs, with their concentration levels found to be in the low nanogram range per $5 \times 10^5$ cells (Figure 4A, Table S3) as previously reported upon S. Typhimurium infection. Similarly, quantification of PGE$_2$, 12-HETE and arachidonic acid in thrombin activated human platelets (Figure 4B) agrees with the results of similar studies. Moreover, simultaneous treatment of platelets with thrombin and the COX inhibitor aspirin resulted in decreased prostanoid levels. In contrast, 12-HETE, which is the most abundant oxylipin amongst those quantified, was increased following both treatment with thrombin and a combination of thrombin and aspirin. This is possibly due to more arachidonic acid being available and therefore a higher turnover rate of 12-LOX. Indeed, aspirin treatment resulted in enrichment of arachidonic acid in the platelets, as a consequence of one oxylipin pathway (COX) being inhibited. Therefore, these results demonstrate that our quantification method not only reaches low oxylipin levels, but also covers the wide dynamic range required for oxylipin analysis in biological samples.

Profiling of oxylipins in biological samples. Although research around oxylipins, especially eicosanoids, emerged in the 1960 after the discovery of their generating enzymes, several members of this lipid class were only characterized in more recent years with many oxylipins possibly still remaining unknown. Hence, we developed a profiling UHPLC-DTIM-MS workflow to investigate oxylipin generation, exemplarily shown on S. Typhimurium infected murine BMDMs and thrombin activated human platelets. A three-level annotation was implemented, with features (i) identified by accurate mass, RT and CCS match with the internal database, (ii) annotated based on accurate mass match with LIMSD, or (iii) flagged as “lipid-like” features. Features were classified as lipid-like if presenting identical accurate mass but differing on either RT or CCS values from the compounds in the database. More specifically, lipid-like features can be oxylipin category-like (e.g. prostaglandin (PG)-like) if their accurate mass and CCS values fall within the annotation thresholds but their RT does not, hence indicating a possible new compound with different polarity within the lipid class. This is illustrated in thrombin activated platelets on the extracted ion chromatogram for m/z 351.2177, which corresponds to a number of different oxylipins, including PGE$_2$ (Figure 5 F, G). On the other hand, if a compound is detected with accurate mass and RT comparable to those of the lipid standard but different DT values, it is flagged as specific lipid-like (e.g. PGF$_2$-α-like). The latter is based on the assumption that structurally similar lipids have a similar “drift-fingerprint”, i.e. a similar distribution of species based on the assumption that structurally similar lipids have a similar “drift-fingerprint”, i.e. a similar distribution of species with identical m/z and RT, but different DT. Hence, the discovery of a feature in a biological sample whose m/z and RT fall within the ranges of a standard, but DT values does not, might represent a new isomer of the molecule. These oxylipin category-like or specific lipid-like flags give one more confidence in identifying new compounds or new conformers not present in the internal database than using solely accurate mass match with an external library, which commonly results in multiple putative annotations for the same feature.

Using this protocol on the BMDMs data, out of 3339 features we standard-identified 85 species, flagged 146 as lipid-like, and putatively annotated with LIPID MAPS 624 features, while 2484 features remained unknown (Table S4 and Figure S10 A).
Figure 5: The changes in the oxylipin profile induced by thrombin treatment of human isolated platelets can be identified using MS/MS spectra and DT alignment of product and precursor ions. (A) Score plot of the principal component analysis (first versus second PC) of lipid extracts of isolated human platelets analyzed by LC/DTIM-MS and processed with Knimet. The four treatment groups are: thrombin plus aspirin (green), thrombin (blue), thrombin plus ethanol (yellow) and untreated (red). (B) Heatmap representation and hierarchical clustering of the distance matrix between samples in the four treatments. Separation between treatment groups can be observed, with three distinct clusters (aspirin, control and ethanol/thrombin) in both PCA and heatmap. (C) Full scan at 3.3 min shows m/z 369.2274, which is putatively annotated as TXB2 and other oxylipins. (D) Product ion spectrum at 3.3 min shows the diagnostic product ions m/z 195.1018 and m/z 169.0864 for TXB2. (E) Precursor (black) and product ions (m/z 195.1018 dashed red; m/z 169.0864, dashed green) align in DT, confirming the identification of the feature as TXB2. Unknown species 1, 2, 3 present identical m/z, different RT (F) but very similar DT (G) of TXB2, hence they are flagged as “lipid-like” features during data annotation.

Our findings agree with previous results showing that infection with S. Typhimurium induces eicosanoid production in BMDMs. Among the identified lipids differentiating between infected and uninfected samples were known mediators of inflammation, such as PGE2 and 5-HETE, (Figure S11, Table S5). The high number of unknown features is due to (i) the accurate mass range of m/z 199-1000 used for feature extraction, which was chosen to include lipid dimers and adducts but might have led to the addition of features not corresponding to oxylipins; (ii) the limited size of our internal database, as well as the relatively low number of oxylipins present in the LMSD. Nonetheless, it has to be taken into consideration that each lipid characterized contains on average 5 drift peaks (discussed above), which play an important role on the overall high number of features. Considering a tighter m/z range comprising only the [M+H]+ species of FAs and oxylipins present in our database (200-500), the number of features reduces to 2595 with 1775 unknowns. However, information about dimers and cluster formation would be lost using this mass range. On the other hand, data processing ignoring the mobility dimension yields 580 features, out of which only 177 are annotated with the fatty acyl portion of LMSD (Supporting Discussion, Figure S12). This noticeable reduction of annotated features further highlights the importance of IM in gaining new insights into LC-MS analysis of lipids. The high abundance of features found using IM might raise concerns about ion formation in the source, ion movement in the drift tube or feature finding algorithms. As described above, we tested the consistency of features found across multiple injections, as well as their changing abundance dependent on either sample concentration or solvent composition, and the results suggest that their presence is not triggered by instrument settings or data processing algorithms but that the features detected by DTIM are present in the sample.

Our oxylipins and FAs profiling method was further tested on human platelets isolated from three donors treated with the pro-coagulant thrombin to trigger oxylipin generation. Statistical analysis of the data processed with Knimet revealed differences between treatment groups (Figure 5, Figure S13 and Table S6), which are in agreement with previous findings. Out of the 2591 features remaining after filtering, 57 were identified by the internal library of standards, 109 were flagged as lipid-like, and 433 could only be annotated with LMSD, while 1992 remained unidentified (Table S7, Figure S10 B). Although putative annotation results in multiple possible identities, the combination of lipid-like flagging, annotation from the external database and finally data-independent acquisition in IM-MS/MS mode allows lipid identification, as exemplified by the identification of Thromboxane B2 (TXB2: Figure 5 C-E): the feature presenting m/z = 369.2270, RT = 3.33 min and DT = 23.89 ms was flagged as PG-like [M-H]- by the internal annotation system and putatively annotated by LMSD as 10,11-dihydro-20-dihydroxy-LTB4, 19-hydroxy-PGE2, 19-hydroxy-PGFα, 20-hydroxy-PGE2α, TxB2 (Table S7). Alignment in DT between precursor and product ions allowed identification as TxB2 (Figure 5 E) and confirmed our strategy of flagging compounds as structural similar to those in the internal database based on their chromatography and mobility characteristics. On the other hand, features flagged as new compounds are identical in m/z and similar in ΔT/CCS but differ in RT, as illustrated on m/z 351.2177 (Figure 5 F, G).

Conclusion. In this study we present a comprehensive LC-DTIM-MS method for both profiling and quantification of oxylipins and their precursors FAs in biological samples. Using replicate acquisitions of lipid standards across different days, an average RSD of 0.19%, 0.21%, 0.20% and 0.18% for the ΔT/CCS of the species [M-H]-, [M-H+CH3CO2Na]+, [2M-H]- and [2M-H+Na]+ respectively, was obtained (Table S1). Further confirmation of the validity of our measurements comes from comparison with CCS values in the literature. This results in an average RSD below 2% (Table S8), which is the accepted error threshold for intra-lab CCS measurements. However, several species are found above this value, with the strongest outlier being lauric acid (RSD = 5.59%). Nonetheless, in-house triplicate acquisitions of lauric acid resulted in highly repeatable measurements (RSD = 0.013-0.466, table S1). This high variability with the CCS reported in the literature could derive from a number of sources of uncertainty, including differences in the analytical details, which must be taken into consideration when building CCS libraries on standards and/or literature values if not enough experimental or computationally generated CCS information exists for comparison. However, the low internal RSDs allowed us to use ΔT/CCS in addition to
RTs and accurate mass to identify oxylipins and FAs in two sets of biological samples. As the CCS value of a compound is a physical-chemical parameter and therefore instrument-independent, it can be used by others as additional lipid identifiers. We showed how CCS values in combination with m/z can improve the identification process of oxylipins and FAs and allows to flag potential new members of this family. Moreover, addition of IM to the workflow allowed separation and identification of different lipid conformers, including dimers and solvent clusters, based on DT-alignment of product ions to their precursors. In addition, our study highlights the complexity of lipid analysis by LC-MS and the necessity of additional ion separation level to gain further insights into ionization, adduct formation and fragmentation mechanisms. Despite a higher number of detected features by LC-DT-MS compared to LC-MS alone, the results presented herein show the potential of this technique in simplifying lipid identification by using CCS and DT-alignment of product and precursor ions as additional identifiers. To further automate the identification process, diagnostic product ions could be added to the library. Moreover, the constant improvement of in-silico tools for DTIM data analysis will allow the combination of our methodology with these computational techniques once oxylipins have been integrated in their training set.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: XXXXXX.

Oxylipin and FAs library; Features found in BMDMs and platelets samples; Features driving PCA separation between BMDMs and platelets samples (xlxs). Supporting Materials and Supporting Discussion; upper and lower limits of quantification for standards; absolute concentrations of species in BMDMs; Comparison between measured CCSs; and literature; figures regarding features found in the BMDM and platelet samples and the advantages of DTIM in oxylipins and FAs identification (PDF).

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