Mass spectrometry images acylcarnitines, phosphatidylcholines, and sphingomyelin in MDA-MB-231 breast tumor models

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Abstract The lipid compositions of different breast tumor microenvironments are largely unknown due to limitations in lipid imaging techniques. Imaging lipid distributions would enhance our understanding of processes occurring inside growing tumors, such as cancer cell proliferation, invasion, and metastasis. Recent developments in MALDI mass spectrometry imaging (MSI) enable rapid and specific detection of lipids directly from thin tissue sections. In this study, we performed multimodal imaging of acylcarnitines, phosphatidylcholines (PC), a lysophosphatidylcholine (LPC), and a sphingomyelin (SM) from different microenvironments of breast tumor xenograft models, which carried tdTomato red fluorescent protein as a hypoxia-response element-driven reporter gene. The MSI molecular lipid images revealed spatially heterogeneous lipid distributions within tumor tissue. Four of the most-abundant lipid species, namely PC(16:0/16:0), PC(16:0/18:1), PC(18:1/18:1), and PC(18:0/18:1), were localized in viable tumor regions, whereas LPC(16:0/0:0) was detected in necrotic tumor regions. We identified a heterogeneous distribution of palmitoylcarnitine, stearoylcarnitine, PC(16:0/22:1), and SM(d18:1/16:0) sodium adduct, which colocalized primarily with hypoxic tumor regions. For the first time, we have applied a multimodal imaging approach that has combined optical imaging and MALDI-MSI with ion mobility separation to spatially localize and structurally identify acylcarnitines and a variety of lipid species present in breast tumor xenograft models.—Chughtai, K., L. Jiang, T. R. Greenwood, K. Glunde, and R. M. A. Heeren. Mass spectrometry images acylcarnitines, phosphatidylcholines, and sphingomyelin in MDA-MB-231 breast tumor models. J. Lipid Res. 2013. 54: 333–344.

Supplementary key words multimodal imaging • phospholipids • matrix-assisted laser desorption/ionization • mass spectrometry

MALDI mass spectrometry imaging (MSI) can detect, localize, and identify multiple biologically relevant molecules directly from thin tissue sections without the necessity for any labeling (1). The sample preparation of thin tissue sections for MALDI-MSI requires the application of laser absorbing matrix crystals on the tissue surface. During an MSI experiment, the laser beam ablates the matrix coated tissue sample surface while the mass spectrometer acquires a collection of spectra that contain comprehensive information about the local molecular composition of the sample. Biomolecules present in the tissue are desorbed and separated by a mass spectrometer according to their m/z ratios. Each m/z value present in this spectral collection can be converted to an image by using dedicated imaging software. The chemical structure of each ion detected from the tissue surface can be identified after its isolation and fragmentation inside a mass spectrometer.

An additional advantage of MSI is its compatibility with other imaging techniques such as optical bright field and fluorescence microscopy. Such a multimodal approach is particularly useful for analyzing the molecular composition of complex, heterogeneous tumor tissue, which comprises several distinct tumor microenvironments. One example...
of such heterogeneity is tissue hypoxia, a state of low oxygen tension (pO_2 values \( \leq 2.5 \) mmHg) observed in many solid tumors (2). Hypoxia increases overall tumor aggressiveness, induces radio and drug resistance, triggers resistance to apoptosis, and enhances cellular migration and invasiveness, eventually leading to cancer metastasis (3). At the molecular level, hypoxia triggers many known and unknown metabolome, lipidome, and proteome changes that modulate tumor vascular expansion and provide an overall advantage for malignant growth, causing rapid disease progression (4). Severe, prolonged hypoxia can lead to cell death and tissue necrosis (4). Tumor expansion results in the formation of metabolically diverse, viable and necrotic, normoxic and hypoxic, vascularized and avascular tissue regions, which, to date, have not been imaged in a single experiment or by a single imaging technique. We hypothesize that these different tumor tissue regions have different distributions of metabolites and lipids. MSI has the capability of parallel detection and visualization of biomolecules present in these specific tissue microenvironments, which allow for a more-comprehensive analysis of tumor biology.

Unlike cancer cell cultures, human tumor xenograft models in mice provide a three-dimensional, native-like environment for studying different aspects of tumor growth (5). Additionally, tumor xenograft models allow for genetic modifications such as incorporation of genes encoding fluorescent proteins for visualizing and monitoring selected processes such as the influence of tissue hypoxia on tumor extracellular matrix (6) or cancer metastasis (7).

MALDI-MSI is well-suited for ex vivo detection and localization of different lipid categories and classes from samples of various origins and different sizes. MALDI-MSI has recently been applied in lipidome studies of human skin (8), veins (9), and ovary (10); mouse embryo implantation sites (11), tongue (12), and leg muscle (13, 14); and rat spinal cord (15), cardiac tissue (16), and liver (17). Mammalian brain tissue, due to its high lipid content, has also been extensively studied by MSI. Ion images of lipid distributions have been obtained for rat (17–22), mouse (23, 24), and human (25, 26) brain sections. Lipid MSI has also been performed for various types of tumor tissue, such as human brain tumors (27), seminoma (28), ovarian cancer (29, 30), myoid liposarcomas (31), human astrocytoma (32), human prostate cancer tissue (33), colon cancer liver metastasis (34), and human liver adenocarcinoma (35).

Lipid-rich tumors have been associated with increased tumor aggressiveness and metastasis (36–38). To develop effective cancer treatment strategies, it is crucial not only to identify but also to localize the lipid molecules involved in cancer progression. To date, lipid distributions in breast tumors remain largely unknown due to limitations in lipid imaging methods, which often require incorporation of fluorescent tags and provide limited specificity (39–41). Here, we have adopted a novel multimodal imaging approach, which integrates microscopy and MSI, for a label-free, ex vivo visualization of different phospholipids directly from breast tumor xenograft tissue sections. We present spatial distributions of sphingomyelin (SM) and multiple phosphatidylcholine (PC) species, which were colocalized with normoxic, hypoxic, or necrotic tumor regions. MSI also localized the distribution of two acylcarnitine molecular ions, which were identified from tumor tissue by ion mobility separation (IMS) followed by tandem mass spectrometry. We have demonstrated that the incorporation of MSI into a multimodal imaging approach revealed the molecular complexity of tumor tissue at an unprecedented level.

### MATERIALS AND METHODS

#### Chemicals and materials

The matrix α-cyano-4-hydroxycinnamic acid (CHCA) was purchased from Sigma-Aldrich (Germany). Water, acetonitrile (ACN), and trifluoroacetic acid (TFA) were purchased from Biowise (The Netherlands). Cresyl violet acetate, Ponceau S, and paraformaldehyde powder were purchased from Sigma-Aldrich.

Gelatin Type A was purchased from Sigma. Mayer’s hematoxylin was purchased from Sigma, and aqueous Eosin Y from EMD Chemicals, Inc. Cytoseal 60 Mounting Medium, Richard-Allan Scientific, was purchased from Thermo Scientific.

#### Breast tumor xenograft model

The MDA-MB-231 human breast cancer cell line was purchased from the American Type Culture Collection (ATCC) and used within 6 months of obtaining from ATCC. This cell line was tested and authenticated by ATCC using two independent methods: the ATCC cytrome C oxidase I PCR assay, and short tandem-repeat profiling using multiplex PCR. The MDA-MB-231 cell line was genetically modified to express tdTomato red fluorescent protein under the control of hypoxia response elements (MDA-MB-231-HRE-tdTomato) as previously described (42). Cells were injected into the upper thoracic mammary fat pad of athymic nude mice (2 × 10^6 cells/injection), and tumor growth was monitored using standard calipers. The research was conducted in conformity with the Public Health Service policy on Humane Care and Use of Laboratory Animals. All experimental animal protocols were approved by the Institutional Animal Care and Use Committee of the Johns Hopkins University School of Medicine.

#### Tumor dissection and optical imaging

When tumors reached a volume of approximately 500 mm³, mice were euthanized and tumors were removed. Three individual tumors were embedded into 10% gelatin blocks prepared using 15 mm × 15 mm × 5 mm cryomolds (Sakura Finetek). Cresyl violet acetate and Ponceau S were prepared in 10% warm (37°C) gelatin at a concentration of 10 mg/ml, mixed in a 1:1 ratio, and injected into 10% gelatin blocks next to the tumor tissue. Each block was sectioned into serial 2 mm-thick fresh tumor sections using an acrylic adjustable tissue slicer (12 mm depth up to 25 mm width, Braintree Scientific, Inc.; Braintree, MA), and tissue slicer blades (Braintree Scientific, Inc.) as previously described (43, 44). These serial fresh tumor xenograft sections were each placed on individual microscope slides (Fisherbrand catalog number 12-550-34, Fisher Scientific; Pittsburgh, PA), and stored in an ice box containing ice on the bottom, with the slides located on a perforated plate at approximately 1-cm above the ice to minimize tissue degradation (43, 44). These fresh sections were imaged by...
bright-field and fluorescence microscopy with a 1× objective attached to a Nikon inverted microscope, equipped with a filter set for 528 to 553 nm excitation and 600 to 660 nm emission, and a Nikon Coolpix digital camera (Nikon Instruments, Inc.; Melville, NY). Bright-field imaging captured the position of the fiducial markers present inside the gelatin block, as well as the shape of the tumor tissue. The fluorescence from tdTomato expression in hypoxic regions of these tumor sections was detected by fluorescence microscopy. The GNU Image Manipulation Program (GIMP 2.6) was used for two-dimensional (2D) coregistration and overlay of bright-field and fluorescence images of 2 mm-thick tumor sections. All 2 mm-thick sections were snap-frozen immediately after microscopic imaging and stored at −80°C until further analysis.

Tumor sectioning

From each 2 mm-thick section, 10 µm-thick sections were cut at −16°C for MSI using a Microm HM550 cryo-micromtome (Microm International GmbH; Walldorf, Germany) along with adjacent 10 µm-thick sections for histological staining. Tissue sections for MSI analysis were mounted onto 25 mm × 50 mm × 1.1 mm, R = 4–8 Ω Indium Tin Oxide-coated slides (Delta Technologies) and for histological staining onto Superfrost Slides (VWR International; Cat. 48311-600).

H and E staining method

Tissue sections were stained using a modified H and E staining protocol as previously described (43–45). Briefly, 10 µm sections attached to Superfrost Slides (VWR International, Cat. 48311-600) were washed with PBS, fixed with freshly depolymerized 3% paraformaldehyde solution for 30 min, washed with distilled water (dH2O), and treated with Mayer’s hematoxylin for 30 min at room temperature, followed by five washes with dH2O. Sections were immediately immersed in aqueous Eosin Y for 30 min, followed by five washes with dH2O, mounting with aqueous mounting medium, and attaching of a coverslip. Bright-field images of H and E-stained sections were acquired using 1× or 20× objectives attached to a Nikon microscope equipped with a Nikon Coolpix digital camera (Nikon Instruments, Inc.; Melville, NY).

MALDI-IMS-MSI

Six tissue sections obtained from three individual tumors were subjected to MSI analysis. Before MSI analysis, tissue sections were placed inside a vacuum desiccator for 30 min. CHCA matrix was prepared at a concentration of 10 mg/ml in 1:1 ACN:H2O/0.1% TFA and was applied by an ImagePrep (Bruker; Germany) application system. Tissue sections were analyzed on a MALDI-Q-TOF (Synapt HDMS; Waters, UK) instrument in IMS mode detecting positive ions. Images were acquired with 150 µm × 150 µm spatial resolution and a 100 µm laser beam spot size. For better detection of masses above m/z 700, the instrument was set to suppress all masses below m/z 700. For 2D MSI analysis and overlay of images, data were visualized using BioMap software (Novartis; Basel, Switzerland). Ion mobility data analysis and visualization were performed by using Driftscope software (version 2.1; Waters, UK) as well as high-definition imaging (HDI) software (version 31; Waters, UK). The LIPID Metabolites And Pathways Strategy database (LIPID MAPS, www.lipidmaps.org) was used to search for possible lipid structures (46, 47).

Lipid identification by MS/MS analysis

The identification of selected lipid species was performed on tissue in positive-ion mode using a MALDI-Q-TOF instrument (Synapt; Waters, UK) after completing MSI experiments. Lipid ion precursors were selected within a 2 Da selection window and fragmented using 20–30 V collision energy applied in the trap cell (in TOF mode) or 30–40 V applied in the transfer cell (in IMS mode). Spectra were analyzed using MassLynx software (Waters, UK). Ion mobility-separated ions were visualized by Driftscope software (Waters, UK), and selected drift time windows were exported to MassLynx for spectral analysis. FA chains of lipids of interest were identified from a combination of the mass of the lipid and the MS/MS fragmentation pattern obtained with a mass accuracy of 0.1 Da at m/z 1,000. The LIPID MAPS, Human Metabolome Database version 2.5 (www.hmdb.ca) and the METLIN Metabolite Database (metlin.scripps.edu) were used to search for possible lipid structures and MS/MS fragmentation spectra.

Overlap analysis

We analyzed three MDA-MB-231-HRE-tdTomato breast tumor xenografts that were imaged by the multimodal approach described above. Necrotic tumor regions were outlined in H and E-stained 10 µm-thick sections. Hypoxic tumor regions were detected from the fluorescence images of tdTomato protein expression in 2 mm-thick sections. The remaining tumor regions were normoxic. Ion images of each individual m/z were overlaid with the corresponding optical images and coregistered by using the positions of fiducial markers and tumor boundaries as previously described (43, 44). The tumor region was masked by applying an active-shape model to the registered optical bright-field images (48). The highest 5% of the area under the histogram of images was automatically thresholded as the hypoxic region inside the tumor, respectively. The volume percentage of ion overlap with normoxic, hypoxic, and necrotic regions inside the tumor was calculated to quantify their colocalizations as previously described (43, 44). Volume percentage of hypoxia was defined as the number of hypoxic voxels that overlap with a given ion normalized to total hypoxic voxels in the tumor. Similarly, volume percentage of ions with normoxic voxels and necrotic voxels was measured. The image registration and colocalization analysis program was written in Matlab 2012b (Mathworks, Inc.; Natick, MA). Mean and standard error of overlap volume percentage from tumor center sections (n = 5) were calculated for each m/z and tumor region using MS Excel 2003.

RESULTS

Multimodal imaging approach

We have employed a multimodal imaging approach for investigating the effects of tumor hypoxia and tumor necrosis on the regionally distinct, spatially heterogeneous lipid composition in a breast tumor model. In this approach, we have integrated optical microscopy with MSI. Bright-field microscopy visualized the shape of the tumor, fluorescence microscopy outlined the position of hypoxic tumor regions, and histochemical staining revealed the localization of necrotic tumor tissue. MSI employing MALDI-MSI visualized the distribution of intact phospholipid molecular ions. All modalities were combined through the presence of fiducial markers visible in all imaging modalities (45). Figure 1 presents results obtained from a representative breast tumor xenograft using our multimodal imaging approach. Figure 1A shows a bright-field image of a 2 mm-thick tumor section obtained from the tumor center, which was embedded in a gelatin block containing the
shows an adjacent 10 µm-thick section stained with hematoxylin and eosin (H and E) for visualization of the necrotic tumor regions. Figure 1H presents the shape of the necrotic regions inside the tumor, outlined by a dotted line. Figure 1I shows the coregistration of the necrotic tumor regions obtained from the H and E image (Fig. 1H) with the hypoxic tumor regions from the fluorescence image (Fig. 1B). The fiducial markers were not clearly visible in some H and E images, in which case, the coregistration was performed based on the tumor shape by using the ImageJ-TurboReg plug-in (http://rsb.info.nih.gov/ij). Magnified images of viable and necrotic tumor regions, as indicated in Fig. 1G, are shown in Fig. 1J, K, respectively.

Mass spectrometric analysis of lipids

Mass spectrometric imaging was employed directly on thin tumor tissue sections for detection, visualization, and identification of a number of lipid biomolecules. Figure 2 presents selected molecular structures of acylcarnitine and lipids detected by MSI. These biomolecules all have a quaternary amine group, which can attain a positive charge, resulting in good ionization properties.

Figure 3 shows a representative combined mass spectrum of positively charged ions detected by MALDI-MSI from MDA-MB-231-HRE-tdTomato breast tumor xeno-graft tissue. In the low-mass region, we observed phosphate-related metabolite ions at \( m/z \) 125.0 [M+H]+, \( m/z \) 146.9 [M+Na]+, and \( m/z \) 162.9 [M+K]+, the phosphocholine at \( m/z \) 184.0 [M+H]+ as well as a phosphocholine-realted ion at \( m/z \) 198.0 [M+H]+. Matrix-related ions were detected at \( m/z \) 172.0 [M-H2O]+, \( m/z \) 190.0 [M+H]+, and \( m/z \) 379.0 [2M+H]+. The fiducial marker cresyl violet was observed at \( m/z \) 262.0. Two acylcarnitines, palmitoylcarnitine and stearoylcarnitine, were observed at \( m/z \) 400.3 [M+H]+ and at \( m/z \) 428.5 [M+H]+, respectively. Lysolipid fiducial markers, as described in Materials and Methods.

Figure 1B presents the fluorescence image of tdTomato red fluorescent protein exclusively expressed in the hypoxic tissue regions of this genetically modified tumor model (42). The coregistration of the bright-field image (Fig. 1A) with the tdTomato fluorescence image (Fig. 1B) was performed based on the position of the fiducial markers, which were visible in both images. The red color of the tdTomato signal was changed to black prior to generating this overlay for better visualization of the hypoxic tumor regions. D: Corresponding optical image of a thin tissue section used for MSI. E: Corresponding MSI image of the fiducial markers. F: Coregistration of fiducial markers imaged by MSI and the hypoxic regions imaged by fluorescence microscopy. G: Image of the adjacent H and E-stained tissue section. H: Image of the outlined necrotic tissue present inside the tumor. I: Coregistration of the necrotic region visualized by H and E staining and the hypoxic region imaged by fluorescence microscopy. J: Magnified image of viable tumor region as indicated in G. K: Magnified image of necrotic tumor region as indicated in G. Scale bar, 5 mm A–I; 250 µm J–K.

Fig. 1. Co-registration of optical microscopy and mass spectrometric imaging. A: Representative bright-field image of a 2 mm-thick fresh breast tumor tissue section embedded in gelatin containing fiducial markers. B: Corresponding fluorescence image of tdTomato expression in the hypoxic tumor regions. C: Coregistration of the bright-field image shown in A and the fluorescence image shown in B based on the position of the fiducial markers present in both images. The red color of the tdTomato signal was changed to black prior to generating this overlay for better visualization of the hypoxic tumor regions. D: Corresponding optical image of a thin tissue section used for MSI. E: Corresponding MSI image of the fiducial markers. F: Coregistration of fiducial markers imaged by MSI and the hypoxic regions imaged by fluorescence microscopy. G: Image of the adjacent H and E-stained tissue section. H: Image of the outlined necrotic tissue present inside the tumor. I: Coregistration of the necrotic region visualized by H and E staining and the hypoxic region imaged by fluorescence microscopy. J: Magnified image of viable tumor region as indicated in G. K: Magnified image of necrotic tumor region as indicated in G. Scale bar, 5 mm A–I; 250 µm J–K.

Fig. 2. Representative examples of acylcarnitine and lipid molecular species such as stearoylcarnitine, LPC, SM, and PC detected from tumor tissue by MSI. The molecular structures were obtained from the LipidMaps web site.
Mass spectrometry imaging of breast cancer

species were detected in the mass range from $m/z$ 450 to $m/z$ 550. Heme was observed at $m/z$ 616.1 [$M^+$] as a singly charged ion. Intact lipid molecular ions were detected as protonated ions as well as sodium and potassium salt adducts in the mass range from $m/z$ 700 to $m/z$ 860.

Figure 4 presents an expanded view of the spectrum shown in Fig. 3 showing multiple sphingomyelin and phosphatidylcholine species detected as proton, sodium, and potassium adducts. We detected and identified protonated species of SM(d18:1/16:0) at $m/z$ 703.5, PC(16:0/16:0) at $m/z$ 734.5, PC(16:0/18:1) at $m/z$ 760.5, PC(18:1/18:1) at $m/z$ 786.6, and PC(18:0/18:1) at $m/z$ 788.6. The most-abundant SM and PC lipid peaks detected from MDA-MB-231-HRE-tdTomato breast tumor xenograft sections by MALDI-MS and MSI acquisitions as sodium and potassium adducts of lipids are listed in Table 1.

The structures of all described lipid species were confirmed by MALDI-MS/MS profiling experiments performed directly on tumor tissue sections. Figure 5 shows a typical MS/MS spectrum of protonated PC(16:0/18:1), which has a highly abundant fragment signal at $m/z$ 184.0 arising from the protonated phosphocholine ion, a signature peak in the MS/MS of protonated lysophosphatidylcholine (LPC), PC, and SM species. A less-abundant signal at $m/z$ 86 is characteristic of the choline head group. The loss of trimethylamine from the phosphocholine head group leads to the fragment ion [M+H-59]+ at $m/z$ 701.5, whereas the loss of the entire phosphocholine head group leads to the formation of the fragment ion [M+H-183]+ at $m/z$ 577.5. Additionally, the peak at $m/z$ 522.3 corresponds to the protonated LPC(16:0/0:0). These four distinct fragments confirm the identity of $m/z$ 760.5 as PC(16:0/18:1).

Mass spectrometric imaging of lipids

MALDI imaging. Images obtained from optical imaging and MALDI-MSI were coregistered as previously described based on the position of the fiducial markers.
The mass spectra of the lipid species were detected by MALDI-MSI with ion drift times (bins) detected by MSI. Table 1 shows the measured m/z values, maximum peak intensities per pixel, and ion drift times (bins) detected by MSI. Values are listed for protonated ions [M+H]^+ and sodium adducts [M+Na]^+ of lipids.

**TABLE 1.** Most-abundant PC and SM species with their observed m/z values, maximum peak intensities per pixel, and ion drift times (bins) detected by MSI.

<table>
<thead>
<tr>
<th>Measured m/z values</th>
<th>Max peak-intensity/pixel</th>
<th>Drift time (bins)</th>
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<tbody>
<tr>
<td><strong>Phospholipid</strong></td>
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<tr>
<td>SM(d18:1/16:0)</td>
<td>734.56</td>
<td>756.54</td>
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<tr>
<td>PC(16:0/16:0)</td>
<td>734.56</td>
<td>756.54</td>
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<tr>
<td>PC(16:0/18:1)</td>
<td>760.58</td>
<td>782.56</td>
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<tr>
<td>PC(18:1/18:1)</td>
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<td>PC(18:0/18:1)</td>
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<td>[M+H]^+</td>
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m/z 760.5 [M+H]⁺ PC(16:0/18:1)

TOF MSMS 760 50LD+
2.90e3

LPC(16:0/0:0) was highly enriched in necrotic tumor regions. The remaining PCs, SM, and acylcarnitines displayed the highest percentage of overlap with hypoxic tumor regions.

**Imaging of isobaric species in IMS mode.** IMS enables the separation of ions on the basis of their collisional cross-section, which depends on their shape and charge. In our study, IMS separated the background ions (chemical noise, matrix clusters, etc.) from the biomolecular lipid species. **Figure 8A** presents a spectrum of peak-picked data acquired during an IMS imaging experiment, and Fig. 8B shows a drift plot of the separated ions. In Figs. 8A and 8B, background ions are highlighted in blue, whereas biomolecular lipid ions are highlighted in red. Ion image 1 in Fig. 8B shows the distribution of a background matrix ion, which was separated from two lipid ions, shown in ion images 2 and 3.

IMS not only improved the MSI data obtained from complex tumor tissue, but also enabled the identification of several biomolecules, which otherwise could not have been separated from other ions of similar mass. **Figure 9** shows an example of two ions at m/z 428.25 and m/z 428.38, which were analyzed in IMS mode. Separated precursor ions were fragmented, which resulted in the identification of m/z 428.3 as stearoylcarnitine. NL of the entire phosphocholine head group leads to the formation of the ion [M+H-183]⁺ at m/z 577.5, whereas the loss of trimethylamine from the phosphocholine head group generates the fragment ion [M+H-59]⁺ at m/z 701.5.

**Fig. 5.** MS/MS spectrum and structure of PC(16:0/18:1). The ion at m/z 86 is a characteristic fragment of the choline head group, which ionizes as an intact molecular ion at m/z 184.0. The fragment ion at m/z 478.3 is generated from loss of the 18:1 fatty acyl chain from the intact lipid molecule, whereas the m/z 504.3 fragment forms from loss of the 16:0 fatty acyl chain from the PC precursor. The protonated LPC(16:0/0:0) is formed as a fragment at m/z 496.3. Similarly, the fragment at m/z 522.3 corresponds to the protonated LPC(0:0/18:1). NL of the entire phosphocholine head group leads to the formation of the ion [M+H-183]⁺ at m/z 577.5, whereas the loss of trimethylamine from the phosphocholine head group generates the fragment ion [M+H-59]⁺ at m/z 701.5.

LPC(16:0/0:0) was highly enriched in necrotic tumor regions. The remaining PCs, SM, and acylcarnitines displayed the highest percentage of overlap with hypoxic tumor regions.
metabolites and phospholipid species in different tumor regions. Unlike classical MALDI MS, which is widely used to identify lipids, MSI does not require extraction of the lipids prior to analysis, but rather visualizes their localization directly in the analyzed tissue. MALDI-MSI does not require any labeling of the lipids. Lipid identification is

**DISCUSSION**

**Advantages of MSI-IMS over other techniques used in lipidomics studies**

In the present study, we analyzed human breast tumor xenograft tissue using MSI-IMS, and localized multiple features in the tissue. These features included viable, necrotic, and hypoxic regions, which were identified using fluorescence microscopy and H and E staining. MSI-IMS was able to visualize the localization of lipids in these different regions, providing a more comprehensive understanding of the lipidome.

**Fig. 6.** Multimodal imaging combining optical microscopy to visualize different tumor microenvironments and mass spectrometric imaging to localize multiple lipid molecular ions. Optical images: A: Diagram showing the localization of viable (in green), necrotic (in blue) and hypoxic regions (in red) in a representative MDA-MB-231-HRE-tdTomato breast tumor section. B: Bright-field image of the corresponding fresh tumor section and fiducial markers visible as red points. C: Image of the hypoxic region obtained by fluorescence microscopy of tdTomato. The black background has been removed for better coregistration with MSI. D: H and E stained image of a tumor section obtained from the fresh section shown in B. The necrotic region present in the tumor center has been outlined. The empty hole within this section originated from loss of tissue integrity in a small part of the necrotic tumor region. Mass spectrometric images from left to right: [M-CH$_3$COO$^-$]/H$_{11002}$ of fiducial markers at m/z 262.0, [M+H]$^+$ of palmitoylcarnitine at m/z 400.3, [M+H]$^+$ of stearoylcarnitine at m/z 428.3, [M+K]$^+$ of LPC(16:0/0:0) at m/z 534.3, [M+Na]$^+$ of SM(d18:1/16:0) at m/z 725.5, [M+H]$^+$ of PC(16:0/16:0) at m/z 734.5, [M+H]$^+$ of PC(16:0/18:1) at m/z 745.5, LPC(16:0/18:1) at m/z 760.5, [M+H]$^+$ of PC(18:1/18:1) at m/z 786.6, [M+H]$^+$ of PC(18:1/18:1) at m/z 788.6, and [M+H]$^+$ of PC(16:0/22:1) at m/z 816.6. Coregistered images: The mass spectrometric images in the top row were coregistered with the optical images in the left column based on the position of the fiducial markers. Scale bar, 5 mm.

**Fig. 7.** Overlap analysis between the MSI-detected molecules and different tumor regions. The necrotic region was separated from the viable region. The latter was further divided into hypoxic and normoxic tumor regions. The necrotic, hypoxic, and normoxic regions together make up the entire tumor tissue. Unlike classical MALDI MS, which is widely used to identify lipids, MSI does not require extraction of the lipids prior to analysis, but rather visualizes their localization directly in the analyzed tissue. MALDI-MSI does not require any labeling of the lipids. Lipid identification is...
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achieved on the basis of their molecular mass and characteristic fragmentation patterns. Such an imaging approach requires very little sample modification prior to imaging, and introduces minimal alterations in lipid localizations and structures. Phospholipid ion formation by MALDI-MSI, when performed directly on a tissue section, generates abundant alkali metal adduct ions, a phenomenon not observed when MALDI is performed on pure lipid extracts. Here, we have employed IMS, because tissue washing removes certain metabolites and lipids from the tissue or leads to their diffusion. Hence, we decided to follow a sample preparation procedure that introduces minimal changes to the composition of the tissue. IMS provided a separation step for our imaging experiments that is usually performed on lipid extracts by HPLC or TLC prior to mass analysis (49, 50). IMS, in which separation occurs in milliseconds, can be easily coupled with MSI and helps obtain ion images of individual lipid species without requiring labor-intensive sample preparation procedures or long acquisition times. The development of new HDI software (Waters, UK) incorporating drift plots and ion drift time made our data analysis fast and simple. The effects of acyl chain length, degree of unsaturation, lipid class, head group, and cation formation on an ion’s collision cross-section in the gas phase and on its ionization efficiency have been studied for direct tissue profiling (51). Such information greatly improves our understanding of ion behaviors detected from a complex tissue sample during MSI-IMS.

Fig. 8. IMS of biomolecular ions detected from thin tumor tissue sections. A: Representative spectrum of peak-picked data acquired during an IMS imaging experiment. The lipid peaks are highlighted in red, whereas matrix-related ions are shown in blue. B: Drift plot of the separated ions. Ion image 1 shows the distribution of a background matrix ion. Ion images 2 (m/z 746.5) and 3 (m/z 746.6) show the distribution of different lipids in the tumor tissue. Lipid-related ions (highlighted in red) were separated from background ions (blue).

Fig. 9. Imaging ion mobility mass spectrometry of molecules present in the MDA-MB-231-HRE-tDTomato breast tumor xenograft model. A: 2D map (m/z vs. drift time) showing ions at m/z 428.2 (red) and m/z 428.3 (green) separated and fragmented during an MS/MS experiment. B: Drift time plot showing drift times of the two clearly separated ions shown in A. C: MS/MS spectrum of fragments detected from both ions without IMS. D: MS/MS spectrum of fragments detected only from the ion at m/z 428.2, whose separation was achieved by IMS. E: MS/MS spectrum of fragments detected only from the stearoylcarnitine ion [M+H]+ at m/z 428.3, whose separation was achieved by IMS.
To date, IMS has been applied to map the distribution of PC and ceramide species from coronal rat brain sections (52) and for imaging of glucosylerceramide in spleen sections from a mouse model of Gaucher disease (53).

Identification of lipids and their biological role

Most of the lipids identified in this study were SM and PC species detected in the mass range from m/z 700 to m/z 850. The presence of a positively charged quaternary ammonium group in both PC and SM species aids in the detection of these two classes of phospholipids in positive-ion mode (51). Previous MALDI-MS studies of phospholipid mixtures have demonstrated that in positive-ion mode, PC and SM species can inhibit the detection of other classes of phospholipids, such as phosphatidylethanolamines, present in the tissue at similar concentrations (54, 55). In this study, we focused on identification and imaging of phosphocholine-containing lipids present in MDA-MB-231-HRE-ttdTomato breast tumor xenograft models.

Analysis of MALDI mass spectra directly obtained from MDA-MB-231-HRE-ttdTomato breast tumor sections revealed multiple PC species with FA chains ranging between 16 and 22 carbons and containing 0 to 2 unsaturated bonds (Fig. 4). These PC species were detected primarily in viable regions of MDA-MB-231-HRE-ttdTomato breast tumors, with PC(16:0/16:0) predominantly localized to normoxic tumor regions and all other PC species localized primarily to hypoxic tumor regions (Fig. 7). The identification of all lipid species involved collisional activation (see Fig. 5 and supplementary Fig. I–V). The protonated LPCs and PCs yielded a major product ion at m/z 184.0, corresponding to the phosphocholine headgroup, as well as a phosphate-related ion at m/z 125.0 [M+H]+, whereas the alkali-cationized molecular species produced characteristic ions at m/z 146.9 [M+Na]+ and m/z 162.9 [M+K]+ (24) instead of m/z 125.0.

Our MSI and MS/MS analyses of MDA-MB-231-HRE-ttdTomato breast tumor tissue revealed an abundant presence of LPC(16:0/9:0) in necrotic tumor regions, which could be due to breakdown of intact PC species. LPC species are formed during inflammation, in cases of chemical exposure (56), and following heart (16) and brain (19, 57) injury. Multiple different bioactive lysophospholipids exhibit pleomorphic effects on multiple cell lineages, including breast and ovarian cancer cells (58, 59). LPC activates the ovarian cancer G protein-coupled receptor family of G protein-coupled receptors, known to regulate cellular motility, growth, differentiation, and gene transcription, all of which are factors central to the biology of cancer (60). The effects of LPC on intracellular signaling in breast cancer cells is translated to functional changes such as decreased production of multiple growth factors from breast cancer cells, including interleukin-6 and -8, which are potent regulators of neovascularization (61). The mechanisms regulating the production and degradation of lysophospholipids are just beginning to be elucidated.

MSI analysis of MDA-MB-231-HRE-ttdTomato breast tumor tissue also revealed the presence of SM(d18:1/16:0), which was detected primarily in hypoxic tissue regions. SM(d18:1/16:0) is known to be involved in tissue inflammation and acts as a precursor of sphingolipid mediators (62, 63). Sphingolipids such as SM(d18:1/16:0) are detected by MSI in the same mass range as PC species. However, the protonated ions of PC species have even m/z values, while those of SM species have odd m/z values; therefore, although they occupy similar mass ranges, the peaks do not overlap, as evident in Fig. 4 (51). Sphingolipids contribute to the pathogenesis of cancer as well as neurological and autoimmune disorders (64). Several sphingolipid-regulated functions are implicated in tumor initiation, promotion, progression, and response to chemotherapy due to their importance in regulating growth, survival, adhesion, and migration of cancer cells (65). Sphingolipids have been proposed as cancer markers (66), and tumors often display alterations in sphingolipid composition (66). Sphingolipid intermediates may trigger proliferation and impair the ability of damaged cells to undergo apoptosis (65). Signaling-induced hydrolysis of sphingolipids produces a series of messenger lipids, such as ceramides and ceramide-1-phosphate, which, as part of signaling cascades, act by recruiting cytosolic proteins. Sphingosine, sphingosine-1-phosphate, ceramide, and ceramide-1-phosphate have pivotal roles in apoptosis, inflammation, and arachidonic acid signaling, all of which are pathways that are impaired in many cancers (67, 68).

Carnitine and acylcarnitines are essential compounds in FA metabolism (69). Carnitine assists with transport of FAs into mitochondria, where FAs are oxidized by β-oxidation as a major source of energy. We have detected and imaged two acylcarnitines directly from MDA-MB-231-HRE-ttdTomato breast tumor tissue sections by applying MSI together with on-tissue IMS-MS/MS analysis combined with a Human Metabolome Database search for their identification. IMS was successfully employed for removing background ion fragments. We have identified ions at m/z 400.3 and m/z 428.3 as protonated ions of palmitoylcarnitine and stearoylcarnitine, respectively, which both localized to hypoxic tumor regions. This accumulation of acylcarnitines in hypoxic tumor regions probably occurred because of a blockage of oxidative processes in these hypoxic tumor regions (70). The inhibition of β-oxidation during hypoxia leads to an accumulation of acylcarnitines in the cytoplasm of hypoxic cells (69). The concentration of long-chain acylcarnitines increased rapidly within minutes after the onset of ischemia in vivo or hypoxia in vitro (71), as shown in plasma and hearts of hypoxic rats (72-74), a newborn rat model of hypoxia-ischemia (75), adult canine myocytes (76), rabbit fetal lungs (70), and blood samples from neonates (77). Acylcarnitines are highly polar, zwitterionic molecules and therefore tend to elute together with phospholipids such as phosphatidylcholine in many chromatographic systems. Here, we have demonstrated that due to the presence of their positively charged quaternary ammonium group, acylcarnitines can also be easily detected and localized by MALDI-MSI.

CONCLUSIONS

We detected, identified, and visualized several important lipid species in a human breast tumor xenograft model.
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REFERENCES


Supplemental Material can be found at: www.jlr.org/content/suppl/2012/08/28/jlr.M207961.DC1.html
SUPPLEMENTAL FIGURE LEGENDS

Supplemental Figure 1. Fragmentation spectra of protonated phosphatidylcholine molecular ions detected from MDA-MB-231-HRE-tdTomato breast tumor xenograft tissue.

Supplemental Figure 2. Fragmentation spectra of sodium and potassium adducts of phosphatidylcholine molecular ions detected from MDA-MB-231-HRE-tdTomato breast tumor xenograft tissue.

Supplemental Figure 3. Fragmentation spectrum of the sphingomyelin molecular ion detected from MDA-MB-231-HRE-tdTomato breast tumor xenograft tissue.

Supplemental Figure 4. Fragmentation spectra of lysosphatidylcholine molecular ions detected from MDA-MB-231-HRE-tdTomato breast tumor xenograft tissue.

Supplemental Figure 5. Fragmentation spectra of palmitoyl- and stearoylcarnitine molecular ions detected from MDA-MB-231-HRE-tdTomato breast tumor xenograft tissue.
Supplemental Figure 1.

$m/z$ 734.5 [M+H]$^+$ PC(16:0/16:0)

$NL$ 256.2 fatty acid 16:0

$m/z$ 125.0

$NL$ 256.2 fatty acid 16:0

$NL$ 59

$NL$ 183 phosphocholine head group

$m/z$ 184.0 [phosphocholine+H]$^+$

$m/z$ 760.5 [M+H]$^+$ PC(16:0/18:1)

$m/z$ 125.0

$NL$ 256.2 fatty acid 16:0

$NL$ 59

$NL$ 183 phosphocholine head group

$m/z$ 184.0 [phosphocholine+H]$^+$
$m/z$ 816.6 [M+H]$^+$ PC(16:0/22:1)

NL 206.2 fatty acid 16:0

$m/z$ 125.0

NL 338.3 fatty acid 22:1

NL 183 phosphocholine head group

$m/z$ 184.0 [phosphocholine+H]$^+$

$2011115_20.34$ (5.985) Cm (1.43)

$154.00 \times 54$

$m/z$ 786.6 [M+H]$^+$ PC(18:1/18:1)

NL 282.2 fatty acid 18:1

$m/z$ 125.0

NL 183 phosphocholine head group

$m/z$ 164.0 [phosphocholine+H]$^+$
$m/z$ 788.6 [M+H]$^+$ PC(18:0/18:1)

NL 284.2 fatty acid 18.0

NL 282.2 fatty acid 18:1

$NL$ 183 phosphocholine head group

$m/z$ 125.0

NL 59

$m/z$ 184.0 [phosphocholine+H]$^+$
Supplemental Figure 2.
m/z 808.6 [M+Na]^+ PC(18:1/18:1)

m/z 810.6 [M+Na]^+ PC(18:0/18:1)
Supplemental Figure 3.

$m/z$ 725.5 [M+Na]$^+$ SM(d18:1/18:0)

$m/z$ 294.2 sphingoid base

$m/z$ 125.0

NL 256.2 fatty acyl 16:0

NL 183 phosphocholine head group

$m/z$ 184.0 [phosphocholine+H]$^+$
Supplemental Figure 4.

**m/z 496.3 [M+H]^+ PC(16:0/0:0)**

- CH$_2$=CH-N(CH$_3$)$_3$
- m/z 163 Da
- m/z 125 Da
- NL 265.2 fatty acid 16:0
- NL 183 phosphocholine head group
  - m/z 184.0 [phosphocholine+H]^+

**m/z 522.3 [M+H]^+ PC(18:1/0:0)**

- CH$_2$=CH-N(CH$_3$)$_3$
- m/z 163 Da
- m/z 125 Da
- NL 282.2 fatty acid 18:1
- NL 183 phosphocholine head group
  - m/z 184.0 [phosphocholine+H]^+
Supplemental Figure 5.

1. $m/z$ 400.3 [M+H]$^+$ palmitoylcarnitine

2. $m/z$ 428.3 [M+H]$^+$ stearoylcarnitine