Supported Membrane Composition Analysis by Secondary Ion Mass Spectrometry with High Lateral Resolution

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ABSTRACT The lateral organization of lipid components within membranes is usually investigated with fluorescence microscopy, which, though highly sensitive, introduces bulky fluorophores that might alter the behavior of the components they label. Secondary ion mass spectroscopy performed with a NanoSIMS 50 instrument also provides high lateral resolution and sensitivity, and many species can be observed in parallel without the use of bulky labels. A tightly focused beam (~100 nm) of Cs ions is scanned across a sample, and up to five of the resulting small negative secondary ions can be simultaneously analyzed by a high-resolution mass spectrometer. Thin layers of 15N- and 19F-labeled proteins were microcontact-printed on an oxidized silicon substrate and imaged using the NanoSIMS 50, demonstrating the sensitivity and selectivity of this approach. Supported lipid bilayers were assembled on an oxidized silicon substrate, then flash-frozen and freeze-dried to preserve their lateral organization. Lipid bilayers were analyzed with the NanoSIMS 50, where the identity of each specific lipid was determined through detection of its unique secondary ions, including 12C1H, 12C2H, 13C, 12C14N, and 12C15N. Steps toward obtaining quantitative composition analysis of lipid membranes that varied spatially in isotopic composition are presented. This approach has the potential to provide a composition-specific analysis of membrane organization that compliments other imaging modalities.

INTRODUCTION

The lateral organization and dynamic rearrangements of lipid and protein components within biological membranes are essential for their functions. The lipid bilayer is the common motif for all membranes; some fraction of the lipid molecules are free to diffuse and mix in the two-dimensional plane of the bilayer, whereas other components are believed to be organized into domains, for example, lipid rafts, that are distinguished by correlated motion and emergent function. Investigation of the associations, organization, and dynamics of membrane components on a length scale that is greater than that of the individual molecules but much less than that which is accessible with light microscopy is a major challenge. Furthermore, since fluid membrane properties, especially those of the lipid components, involve delicate interactions that are highly sensitive to their specific chemical and physical properties, elucidation of the spatial organization of all the components present without the use of large and chemically complex labels, such as those used in fluorescence microscopy, is most desirable. Herein, we demonstrate that secondary ion mass spectrometry (SIMS), implemented using a recently developed instrument from Cameca Instruments (Courbevoie, France), the NanoSIMS 50, can be combined with isotopically labeled membrane components and tools developed for supported bilayer patterning (Groves and Boxer, 2002) to obtain information on the lateral composition of a membrane without altering the chemical composition of the components of interest. Moreover, because the ultimate resolution of the NanoSIMS 50 is on the order of 50 nm, this technique surpasses the lateral resolution of fluorescence microscopy. Through a combination of sample preparation, isotopic substitution strategies, data acquisition, and analysis, this method may ultimately be a powerful technique to probe membrane composition and spatial organization in a manner that complements current approaches.

A comprehensive view of the complex lateral organization of biological membranes requires the application of many methods in parallel. To date, most studies of lipid domains utilize model membrane systems and a multitude of techniques, including atomic force microscopy (AFM) (Lawrence et al., 2003; Rinia et al., 2001; Giocondi et al., 2001; Yuan et al., 2002), differential scanning calorimetry (Gandhavadi et al., 2002; Shaikh et al., 2001), single particle tracking (Dietrich et al., 2001a), electron spin resonance and infrared spectroscopy (Veiga et al., 2001), x-ray diffraction (Gandhavadi et al., 2002), nuclear magnetic resonance (Filippov et al., 2004), and fluorescence microscopy (Dietrich et al., 2001a,b; Hwang et al., 1998; Korlach et al., 1999; Samsonov et al., 2001; Scherfeld et al., 2003; Veatch and Keller, 2003). Fluorescence microscopy is the most powerful approach for achieving the highest sensitivity. However, this requires the addition of fluorescent labels that might interfere with the delicate interactions that occur within membranes (Burns, 2003) and, of course, only the component that is conjugated to the label can be detected. There is an extensive literature on the differential partitioning of fluorescent labels into different phases in monolayers (Dietrich et al., 2001a; Ianoul et al., 2003; Keller, 2002, 2003; Keller et al., 2000; Strottrup et al., 2004), supported

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NanoSIMS 50 uses a tightly focused Cs+ ion beam that preserves the surface organization with minimal perturbations. In our experiments, the Cameca NanoSIMS 50 instrument. The incoming focused Cs-ion beam generates secondary ions; negative ions are collected and analyzed in a high-resolution mass spectrometer (not shown). The identities of different components are encoded by isotopic or atomic labeling.

MATERIALS AND METHODS

Silicon/SiO2 substrates

To dissipate charge on the surface, samples for NanoSIMS 50 measurements must be 100 nm thick. Soft protein and lipid membranes described in this report have been previously characterized with the NanoSIMS 50. These materials present many challenges, and we address a first set of these in the following by imaging protein adlayers and supported lipid membranes that contained simple variations in their lateral composition.
Lipid Bilayers Analyzed with a NanoSIMS

Protein stamping

Proteins were stamped onto the wafer surface by microcontact printing (James et al., 1998) using polydimethylsiloxane (PDMS) stamps (Sygald 184 silicon elastomer, Dow Corning, Midland, MI). The PDMS was cured on silicon masters that were patterned with photoresist, generating a 1 μm raised grid, in which the bars of the grid were either 2.5 μm or 5 μm wide and spaced 25 μm apart. A droplet of protein solution was deposited on the PDMS stamp and incubated for 2 min. The excess protein solution was absorbed with a tissue and the surface of the stamp was dried under a stream of nitrogen gas. The protein-coated PDMS stamp was pressed against the oxidized silicon substrate with a 12 g weight, which typically left a 5–10 nm thick adlayer of protein on the surface (imaged by AFM, see below). In certain cases, these printed proteins could be used to pattern the assembly of supported lipid bilayers (Kung et al., 2000).

Three proteins were utilized in this work. Fibronectin (FN, C22972H3564O7362N6412S180) (Sigma-Aldrich, St. Louis, MO) was diluted to a concentration of 100 μg/ml in 10 mM Heps;150 mM NaCl solution. For some samples, FN was labeled with Cy3.5 (Amersham Biotech, Piscataway, NJ) for visualization. Acyl carrier protein 2 that was uniformly isotopically labeled with 15N to ~90% (15N-Acp2, C520H841O152N167 S1) was generously provided by V. Y. Alekseyev in the Khosla lab (Stanford University). Under certain conditions, the purity of this protein was greater than 98%, as assessed by SDS-PAGE (Wu et al., 2002). A sample of recombinant human dihydrofolate reductase (DHFR) grown in the presence of 4-fluorophenylalanine (F-DHFR, C841H1275O245N235F) was generously provided by J. T. Schulte in the Swartz lab (Stanford University). Under previous methods (Juneja et al., 1989). Thin layer chromatography was performed according to reported procedures (Juneja et al., 1989). Thin layer chromatography analysis (65:25:4 chloroform/methanol/water) showed the purified 15N-egg PC comigrated with an egg PC standard. 1,2-Dioleoyl-sn-glycero-3-phosphatidylcholine (DOPC) (Avanti Polar Lipids) was purchased from Sigma-Aldrich, Inc. (St. Louis, MO) for use in supported lipid bilayers (Kung et al., 2000).

Supported lipid bilayers

Egg phosphatidylcholine (egg PC, Avanti Polar Lipids, Alabaster, AL), Texas Red 1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine (TR-DHPE, Molecular Probes, Eugene, OR) and 1-palmitoyl-D17:2-oleoyl-sn-glycero-3-phosphocholine (D17:2-POPC, Avanti Polar Lipids) were purchased in chloroform. Egg phosphatidylcholine-15N (15N-egg PC) was prepared from egg phosphatidylethanolamine (Avanti Polar Lipids) and choline chloride-15N (99 atom % 15N, Icon, Summit, NJ) following previously reported procedures (Juncera et al., 1989). Thin layer chromatography analysis (65:25:4 chloroform/methanol/water) showed the purified 15N-egg PC comigrated with an egg PC standard. 1,2-Dioleoyl-sn-glycero-3-phosphatidylcholine (DOPC) (15C, Sigma-Aldrich) according to previous methods (Patel et al., 1979). The purified product coprecipitated with a DOPC standard using the aforementioned thin layer chromatography solvent system. Typically, 0.5 mol % TR-DHPE was added to the bilayer samples to aid in imaging the membrane with fluorescence microscopy, which was performed to evaluate the homogeneity of the sample before and after freeze-drying, and for comparison with the composition acquired with the NanoSIMS 50. The desired lipid mixtures were dissolved in chloroform, dried under a nitrogen stream, desiccated under vacuum for at least 3 h, and reconstituted with Millipore (Bedford, MA) water (18 MΩ) to a concentration of 5 mg/ml. Small unilamellar vesicles were formed by passing the lipid suspension through a 50 nm polycarbonate membrane 19 times. The extruded vesicles were stored at 4°C and used within 1 week. Supported bilayers were formed by exposing the oxidized silicon substrate to the vesicle suspension, and excess vesicles were removed by rinsing with Millipore water.

Although methods are available to prepare precise variations in lipid composition across a patterned surface (Kam and Boxer, 2000), the small dimensions of the NanoSIMS 50 substrates led to a simplified method for these first experiments. Two 1 μl droplets, each containing an isotopically distinct vesicle population, were deposited on a glass coverslip such that they were separated by ~2–3 mm. A protein-patterned substrate was then placed over the droplets, causing the two droplets to come into contact with one another and begin to mix. Since membrane formation on the substrate was more rapid than the mixing of the two vesicle suspensions, the ratio of the two lipid components within the resulting bilayer varied from corral to corral. The composition within each corral was allowed to become fully mixed before freezing the samples.

Freeze-drying

Supported bilayers are only stable when bathed by aqueous solutions and delaminate from the surface upon contact with air (Cremer and Boxer, 1999; Holden et al., 2004). To prepare the samples for the NanoSIMS 50 experiments, which are conducted under UVH, the water must be removed without disrupting the lateral organization of the membrane (at least on the scale of the 100 nm area sampled by the Cs+ beam). The freeze-drying method was derived from those employed in freeze-fracture and cryoelectron microscopy experiments (Ryan, 1992), which involves flash-freezing and careful removal of bulk water. After assembly, the supported bilayer samples were manually plunged into liquid ethane (cooling rate ~15,000°C/s (Ryan, 1992) with a well-separated melting and boiling point, 90 K and 184 K, respectively, that minimizes ice crystallization). Samples were then transferred into a precooled vessel that was submerged in liquid nitrogen to keep the water in a frozen amorphous state during the freeze-drying process, then the ice was removed under reduced pressure (90 μbar) for at least 16 h using an oil-free scroll pump (Varian, Lexington, MA) and a liquid nitrogen trap to minimize surface contamination. During the first 4 h that the samples were subjected to reduced pressure, they were kept cold by immersing the vessel that contained the samples in liquid nitrogen. This process evolved by trial and error, in which the primary criteria for the quality of preservation was gauged by the homogeneity of the lipid membrane within each corral, as determined by imaging the dehydrated sample with fluorescence microscopy and AFM. The most common problem was slow transfer of the sample from the water into the liquid ethane, which adversely affected the bilayer structure. Artifacts of the freeze-drying process were easily identified by fluorescence microscopy (see below).

Secondary ion mass spectrometry

SIMS was performed using the Lawrence Livermore National Laboratory Cameca NanoSIMS 50 in simultaneous secondary ion collection mode with pulse counting on the electron multipliers. A microcosmic source was used to generate 16 kV Cs+ primary ions for sample interrogation. The primary beam was tuned to a nominal spot size of ~100 nm, which determines the lateral resolution of these measurements (100 nm), with ~2 pA Cs+, and stepped over the sample in a 256 × 256 or 1024 × 1024 pixel raster to generate quantitative secondary ion images. The same region was rerastered (1–7 times), creating a series of images, one for each raster plane, that were compiled together to create the final image, which is presented in this article. The dwell time was 1 ms/pixel and the magnetic field in the mass spectrometer was derived from those employed in freeze-fracture and cryoelectron microscopy experiments (Ryan, 1992), which involves flash-freezing and careful removal of bulk water. After assembly, the supported bilayer samples were manually plunged into liquid ethane (cooling rate ~15,000°C/s (Ryan, 1992) with a well-separated melting and boiling point, 90 K and 184 K, respectively, that minimizes ice crystallization). Samples were then transferred into a precooled vessel that was submerged in liquid nitrogen to keep the water in a frozen amorphous state during the freeze-drying process, then the ice was removed under reduced pressure (90 μbar) for at least 16 h using an oil-free scroll pump (Varian, Lexington, MA) and a liquid nitrogen trap to minimize surface contamination. During the first 4 h that the samples were subjected to reduced pressure, they were kept cold by immersing the vessel that contained the samples in liquid nitrogen. This process evolved by trial and error, in which the primary criteria for the quality of preservation was gauged by the homogeneity of the lipid membrane within each corral, as determined by imaging the dehydrated sample with fluorescence microscopy and AFM. The most common problem was slow transfer of the sample from the water into the liquid ethane, which adversely affected the bilayer structure. Artifacts of the freeze-drying process were easily identified by fluorescence microscopy (see below).

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electron multiplier (EM5) ( Cameca_NanoSIMS), and the four movable electron multipliers were positioned to detect $^{12}\text{C}^{-}$, $^{12}\text{C}^{14}\text{N}^{-}$, $^{12}\text{C}^{14}\text{N}^{-}$, $^{19}\text{F}^{-}$, $^{12}\text{C}^{14}\text{N}^{-}$, $^{13}\text{C}^{14}\text{N}^{-}$, or $^{31}\text{P}^{-}$. The secondary mass spectrometer was tuned for $\sim 6800$ mass resolving power to resolve isobaric interferences, which arise from mass interferences that occur whenever another ion has the same nominal mass as the analyte ion. The slit at the entrance of the multiplier was typically set at 80 μm; however, when the resolution of $^{13}\text{C}^{1}\text{H}^{-}$ (mass charge ratio) = 14.0106304 amu) (atomic mass unit), $^{12}\text{C}^{1}\text{D}^{-}$ (m/e = 14.0135524 amu), and $^{12}\text{C}^{1}\text{H}_{2}^{-}$ (m/e = 14.0151014 amu) was desired, a slit width of 25 μm was employed. Samples were also simultaneously imaged using the secondary electrons that were detected by a photomultiplier. Samples were first visually located using a charge-coupled device camera. Next, the samples were moved to the analysis position in the NanoSIMS 50 and positioned exactly using real-time secondary electron imaging. A map of their overlapping regions matched. For the analysis of gradients, the sample was obtained by assembling a series of measurements such that the sample was FN because its position differed from that of $^{15}\text{N}-\text{Acp2}$, as shown in panel B. Closer inspection of the map of the $^{12}\text{C}^{14}\text{N}^{-}$ secondary ion signal reveals the third protein grid, $^{15}\text{N}-\text{Acp2}$, which has one-fourth the signal intensity of the other two proteins. The detection of a $^{12}\text{C}^{14}\text{N}^{-}$ secondary ion signal from the $^{15}\text{N}-\text{Acp2}$ protein grid reflects the incomplete (90%) isotopic substitution of $^{15}\text{N}$ for $^{14}\text{N}$. The $^{12}\text{C}^{-}$ and $^{32}\text{S}^{-}$ signals, which are shown in Fig. 2, D and E, were generated by all three protein grids. Defects in the protein patterns are visible in panels A and B; the $^{15}\text{N}-\text{Acp2}$ and $^{19}\text{F}-\text{DHFR}$ protein grids were not continuous in the regions where the substrate was already patterned with protein, indicating protein-on-protein deposition typically did not occur. Additionally, protein over stamping also produced defects in the protein grid that was previously deposited on the silicon substrate. As shown in panels C–E, a portion of the FN grid was removed by subsequent contact with the $^{15}\text{N}-\text{Acp2}$ and $^{19}\text{F}-\text{DHFR}$ stamps. For panels A, B, C, D, and E in Fig. 2, the regions of maximal signal intensity denoted by circles in the schematic drawings are 1, 2, 11, 9, 2, 27, and 6.5 relative to the $^{19}\text{F}^{-}$ signal, respectively. For comparison of the ion yields in each sample, the signal intensities of all five secondary ions were measured at a localized region on each sample, which is designated with a star. As expected, the ratio of the $^{12}\text{C}^{15}\text{N}^{-}/^{12}\text{C}^{14}\text{N}^{-}$ signal intensities measured for $^{19}\text{F}$- DHFR (Fig. 2 A) and FN (Fig. 2 C) correspond to the natural abundance ratio of $^{15}\text{N}/^{14}\text{N}$ (0.4%). The $^{19}\text{F}^{-}$ yield was 16 times that of the $^{12}\text{C}^{14}\text{N}^{-}$ yield for $^{19}\text{F}$-DHFR, and the relative $^{32}\text{S}^{-}$ yield varied between 4 and 6 times that of $^{12}\text{C}^{14}\text{N}^{-}$ for all three different proteins, where the yield was calculated by taking the ratio of the relative SIMS signal intensity to the relative theoretical number of species present in the protein. The high ion yield and uniqueness of the $^{19}\text{F}^{-}$ ions demonstrates that fluorine is an attractive atomic label for future work, especially because only a relatively small amount of fluorine incorporation is required for detection. Comparison of yields between proteins was not performed because variations in protein thickness can drastically influence their secondary ion signal intensities, especially when the sample is rastered multiple times, since a thicker adlayer continues to produce secondary ions after a thinner adlayer is sputtered away. This complication is absent for lipid membranes since their thickness is roughly constant.

### Supported lipid bilayers

Although the thickness of a supported lipid bilayer in an aqueous environment (5 nm) is similar to that of microcontact-printed proteins, the potential amount of high-yielding secondary ions (i.e., $^{12}\text{C}^{14}\text{N}^{-}$) produced by the lipid
components is significantly lower than that of the protein. For compatibility with UHV conditions, supported lipid bilayers were flash-frozen and freeze-dried to preserve their lateral organization. Fig. 3A shows an AFM image of a freeze-dried supported lipid bilayer (egg PC doped with 0.5 mol % TR-DHPE) that was patterned with a FN grid, which prevents the lipid components that were deposited within each corral from mixing with neighboring corrals. The width of the protein grid measured by AFM and SIMS are very similar, being 3.4 ± 0.2 μm and 3.8 ± 0.2 μm, respectively. The protein barrier protruded 3 nm above the lipid membrane within the grid, which was extremely flat with a root mean-square roughness of 0.2 nm over an area of 100 μm², similar to that of the clean oxidized silicon substrate (0.18 nm over the same area). The presence of the lipid membrane within the corral was confirmed with fluorescence microscopy, which detected the TR-DHPE within the membrane. Although freeze-drying decreased the fluorescence intensity of the TR-DHPE, the fluorescence from the lipid membrane was still 3.3 times greater than that detected for the protein grid after background subtraction. Analysis of the same region using the NanoSIMS 50 demonstrates that supported bilayers can be detected. (To test the sample stability over time, this sample was re-measured after 48 h in UHV, then stored in ambient air for 6 months and reintroduced to the UHV two more times, giving a total time under vacuum of 270 h. The 12C/13C ratios measured for proteins and lipids after all four different measurements varied by a total of only 4–5%, suggesting that materials did not sublime off the surface under the UHV conditions.) The location of the lipids and proteins was evidenced by contrast in the 12C/13C (Fig. 3B), 31P/13C (Fig. 3C), and, when appropriate, 12CD/13C (vide infra, see Figs. 4A and 5B) signals. The 12C, 31P, and 12C14N signals from the supported bilayers were 61%, 1150%, and 9%, respectively, compared to that obtained from the protein grid.

Although the silicon substrates were carefully cleaned, contamination of the substrate by residual material from microcontact printing and contaminants from the air and buffers was difficult to avoid. Therefore, whether the differences in the signal intensities detected for the protein grid and the lipid membrane-covered regions could be attributed to surface contamination was investigated. The background signals collected from the lipid- and protein-free regions of the oxidized silicon substrate were 61%, 1150%, and 9%, respectively, compared to that obtained from the protein grid.

The 12C image, also present in all three proteins. The dots in the schematic guide correspond to regions with maximal intensities, which are 527, 1068, 6280, 1436, and 3402 counts/pixel s, respectively. All five of the secondary ions were measured at a localized region on each protein, which is denoted by the star. The total dwell time was 4 ms (1 ms each serial image; total of four serial images).

FIGURE 2 SIMS data for a set of differently labeled proteins deposited in a grid pattern by microcontact printing. Fibronectin (FN, natural abundance) was printed first, followed by 15N-labeled-Acp2, and then 19F-Phe-substituted DHFR such that the protein grids were rotated and offset from each other. The left panels present the SIMS data for different species, and a schematic guide is presented to the right. (A) The 19F image exclusively corresponds to 19F-DHFR. (B) The 12C15N image corresponds to 15N-Acp2. (C) The 12C14N image, which shows two separate protein grids, 19F-DHFR and FN. (D) The 32S image (32S is present in all three proteins). (E)
values as an estimate of the magnitude of the background signal in the lipid membrane samples, the signal intensities observed for the lipid membrane relative to those for FN were appreciably greater than the background, confirming that the lipid membrane did, indeed, produce the secondary ions collected in that region. Because each lipid has only a single phosphorous atom (two per ~60 Å²), the detection of \( \text{P}^- \) demonstrates both the extraordinary sensitivity of the NanoSIMS 50 and its ability to image thin films of biological materials with high contrast, where contrast refers to the difference between the light and dark regions of the image produced by the NanoSIMS 50, which is dictated by the signal intensities at these regions. Although the presence of an element that is unique to the phospholipids, phosphorous, simplifies differentiating the membrane from the protein adlayer, the NanoSIMS 50 can also distinguish isotopically substituted lipids from other components. Fig. 4A shows the \( ^{12}\text{C}^- \) secondary ion signal of a D₃₁-POPC lipid membrane patterned with a FN grid. The \( ^{12}\text{C}^- \) secondary ion signal was 6.1 times greater than that of the protein adlayer, which contains deuterium at its natural abundance level (0.02%). Likewise, the presence of an \(^{15}\text{N}\)-egg PC membrane doped with 0.5 mol % TR-DHPE was evidenced by a \( ^{12}\text{C}^{15}\text{N}^- \) signal that was 5.0 times that of FN, which contains \(^{15}\text{N}\) at natural abundance (0.4%). The lower than expected excess of \( ^{12}\text{C}^- \) and \( ^{12}\text{C}^{15}\text{N}^- \) suggests that the lipid membrane also contained a significant amount of isotopically unmodified lipid; however, as mentioned in the previous section, the differences in the thickness of the protein and lipid layers makes comparison of their ion yields unreliable.

\**Bilayer composition variations**

Although the results presented above demonstrate that biological materials can be imaged with high contrast and high lateral resolution by exploiting differences in the masses of their secondary ions, ultimately, quantitative information on the composition of a mixture is most desirable. As a first step in this direction, we investigated whether the NanoSIMS 50 could detect variations in membrane composition by...
forming a lipid bilayer that spatially varied in isotopic composition on a substrate patterned with protein grids, which serve as barriers to lateral diffusion. As described above, variations in the composition within the corralled regions were created by allowing two drops of different vesicle solutions to mix on a patterned surface during membrane deposition. Fig. 5 shows results for a freeze-dried sample with a compositional gradient between partially deuterated lipids (D31-POPC) and isotopically unmodified egg PC (doped with 0.5 mol% TR-DHPE for visualization). Fig. 5 A shows a fluorescence image of this surface, where the fluorescence intensity was digitized at a magnification of 400. The unlabeled microcontact-printed FN barriers appeared dark, but the fluorescence intensity, and, therefore, the composition of the lipid membrane, varied from left to right. Subsequent AFM measurements of this sample indicated that the regions covered by the lipid membrane were topographically homogeneous (data not shown), similar to that shown in Fig. 3 A. The normalized 12CD− ([12C]/[12C]−) signal acquired by the NanoSIMS 50 at the same region is shown in Fig. 5 B. As expected, the natural isotopic abundance FN barriers appear homogeneously dark in contrast to the lipid membrane. The normalized 12CD− ([12C]/[12C]−) signals generated by each corralled membrane patch varied in intensity in parallel with the fluorescence intensity, suggesting that the quantity of deuterium-enriched lipid within each corral differed. The rectangular region of low secondary ion signal intensity, represented by a dark silhouette located approximately six corrals from the left side of Fig. 5 B, resulted from exposing this area to the Cs+ beam for 3 min before acquiring the entire image. Further investigation with AFM revealed this region corresponded to a crater with a depth of 0.2 nm, indicating that the damage produced by extended exposure of the sample to the Cs+ ion beam was limited to the soft adlayer and did not significantly affect the silicon substrate.

To verify that the secondary ion signal intensities were quantitatively dependent on the lipid components within the sample, we examined the relationship between the intensities of component-specific secondary ions and the fluorescence signal. Both the normalized 12CD− ([12C]/[12C]−) and 12C1H− ([12C]/[12C]−) signals are plotted as a function of the fluorescence intensity in Fig. 5 C, where each data point represents the average fluorescence and secondary ion signal collected from one of the eight corrals depicted in the lower row of Fig. 5, A and B (highlighted with arrows). Because the natural abundance egg PC contained the fluorescent lipid and the deuterium-enriched lipid was not doped with a fluorescent lipid, a rise in the normalized 12C1H− signal should accompany an increase in fluorescence, whereas the 12CD− signal should increase when the fluorescence falls if the secondary ion signal is quantitative. Indeed, both plots exhibited these trends, and the normalized 12CD− and 12C1H− signals were also linearly correlated to the fluorescence intensity (R2 = 0.97 and 0.95, respectively), suggesting the possibility for quantitative analysis of lipid mixtures using the NanoSIMS 50. Exploration of the quantitative aspects of this technique is under way and will be presented in a detailed report.

Choice of isotopic labels

The commercial availability of many deuterium-enriched compounds, primarily due to their widespread application for NMR experiments, renders deuterium an obvious choice as an isotopic label for a component of interest. However, to investigate the spatial organization of multiple lipids within a mixture, each unique component of interest must be individually labeled with a specific stable isotope that is exclusive to that component. Besides deuterium, there are three more potential isotopic labels for lipids: 13C, 18O, and 15N. 15N can be selectively substituted into the headgroup of a phospholipid, but this generates only a single unique nuclide per lipid molecule. Although there is not a simple method to replace the six oxygen atoms within a phospholipid with 18O, an 18O label can be easily incorporated into cholesterol (Hudgins et al., 1988). Preliminary results indicate that the background signal from the natural abundance of 18O (0.2%) in the oxidized silicon substrate overwhelms the 18O− secondary ion signal collected from

![Image](https://example.com/image.png)
the sample, obscuring visualization of the $^{18}$O-cholesterol (substrates prepared with $^{18}$O-depleted oxygen may curtail this background signal). Therefore, $^{15}$N- or $^{13}$C-enriched lipids were employed in the following experiments. To determine the efficacy of $^{15}$N- and $^{13}$C-labeled lipid detection by the NanoSIMS 50, a single substrate that supported an array of compositionally distinct lipid bilayers was employed, in which the composition of the membrane assembled within each corral exhibited a smooth transition from $^{15}$N-egg PC doped with 0.5 mol % TR-DHPE to $^{13}$C-enriched lipids (DOPC-$^{13}$C$_3$, N,N,N-trimethyl-$^{13}$C$_3$ choline, 99 atom % $^{13}$C) at the other. After freeze-drying, the compositional gradient was clearly evidenced by a variation in the fluorescence signal intensity of the membrane, which increased from left to right in Fig. 6A, where the fluorescence intensity was digitized at a magnification of 100. In addition, small regions of bright fluorescence were distributed across the sample; these regions were disregarded in the experimental analysis that follows but can be exploited as discussed in the next section. The normalized $^{12}$C$^{15}$N$^{-}$/$^{12}$C$^{15}$N$^{-}$/$^{12}$C$^{-}$ signal collected from the region shown in Fig. 6A using the NanoSIMS 50 corroborates the presence of a compositional variation in the sample. Similar to the fluorescence, the $^{12}$C$^{15}$N$^{-}$/$^{12}$C$^{15}$N$^{-}$/$^{12}$C$^{-}$ signal intensity increased from left to right in Fig. 6B, which was expected since the $^{15}$N-egg PC solution also contained the fluorescent lipid. In contrast, a change in the composition of the membrane within this region was not visible from the image of the normalized $^{13}$C$^{-}$ signal ($^{12}$C$^{-}$/$^{13}$C$^{-}$) produced by DOPC-$^{13}$C$_3$ (not shown), indicating the $\sim7\%$ $^{13}$C-enrichment (relative to 1.1\% $^{13}$C natural abundance) was insufficient for unambiguous visualization of the compositional gradient with the NanoSIMS 50. (We have not yet explored the use of other combinations of isotopically labeled CN$^{-}$, such as $^{13}$C$^{15}$N$^{-}$ or $^{13}$C$^{14}$N$^{-}$ as potential component-specific secondary ions. However, $^{13}$C$^{14}$N$^{-}$ and $^{12}$C$^{13}$N$^{-}$ cannot be simultaneously detected due to constraints on the physical placement of the detectors.) Fig. 6C graphically illustrates the relationship between the normalized secondary ion and fluorescence signal intensities acquired from the 15 corrals displayed in the indicated row in Fig. 6. A and B, where each point depicts the average signal intensities collected from a small region (100–180 $\mu$m$^2$) within a single corral that was devoid of abnormally bright fluorescent spots. Again, increases in the fluorescence were paralleled by a rise in the $^{12}$C$^{15}$N$^{-}$ signal with a linear correlation ($R^2 = 0.99$) between the two signals. Although the gradual change in membrane composition was not evident from the SIMS image of the normalized $^{13}$C$^{-}$ signal, its presence was suggested in the quantitative analysis and plot of the $^{13}$C$^{-}$ versus fluorescence signal intensities. A decrease in the normalized $^{13}$C$^{-}$ signal was linearly correlated ($R^2 = 0.45$) with an elevation in fluorescence intensity, as predicted a priori since the $^{13}$C-enriched lipid was not doped with a fluorescent label. The correlation factor $R^2$ increases to 0.9 if only the last nine corrals are considered, where a significant variation in the fluorescence intensity (from 216 to 536) also indicates that the membrane composition changes appreciably. Nevertheless, the maximum normalized $^{13}$C$^{-}$ signal was only 1.3 times greater than its minimum value, so a more highly enriched $^{13}$C-labeled lipid would be necessary for future experiments.

**Detection and analysis of microdomains**

As mentioned above, fluorescence imaging revealed localized regions of elevated fluorescence scattered across some of the corrals in the $^{15}$N- and $^{13}$C-enriched lipid gradient sample. The origin of this effect is unknown, but we find that these bright spots are more frequently observed when TR-DHPE is employed for visualization with fluorescence microscopy, and protein barriers are used to pattern the membrane, compared to the use of 12-(N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)amino)-labeled lipids and chrome grids for these respective roles. Although these bright spots were unintended defects within the membrane, we can exploit their presence to test whether the NanoSIMS 50 can image structures within a membrane (i.e., microdomains). The elevated fluorescence signal that typified these defects suggested they contained a TR-DHPE concentration that

![FIGURE 6 Isotopically labeled bilayer compositional array was analyzed using fluorescence and the secondary ion signals collected by the NanoSIMS 50. A gradient between $^{15}$N-egg PC (containing 0.5 mol % TR-DHPE) and DOPC-$^{13}$C$_3$ (terminal methyl in choline headgroup labeled with carbon-13) was formed by combining FN-patterned surfaces with two droplets of vesicles, each consisting of one of the two isotopically labeled lipids ($^{15}$N-egg PC or DOPC-$^{13}$C$_3$). (A) Fluorescence image showing the variation in composition from little fluorescence to maximum fluorescence. (B) $^{12}$C$^{15}$N$^{-}$/$^{12}$C$^{15}$N$^{-}$/$^{12}$C$^{-}$ SIMS signal from the same region indicates the substitution of a single nitrogen-15 atom into each lipid is adequate for detection with the NanoSIMS 50 (the total dwell time was 2 ms). (C) Normalized $^{13}$C$^{-}$/$^{12}$C$^{-}$ and $^{13}$C$^{14}$N$^{-}$/$^{12}$C$^{-}$ signals as a function of the fluorescence taken along the lower row highlighted with arrows in panels A and B. The normalized $^{13}$C$^{14}$N$^{-}$ signal also increased with the fluorescence. Note that the data range of SIMS signals differs from Fig. 5C due to lower intensities of $^{15}$C$^{14}$N$^{-}$ and $^{13}$C$^{-}$. The best fit to a linear relationship is plotted through the points.
was higher than the rest of the membrane, but lower than that which leads to self-quenching. Because TR-DHPE is the only component that contains sulfur, the $^{32}\text{S}^-$ signal was used to image TR-DHPE, as shown in Fig. 7. The fluorescence signal (green) is shown in Fig. 7 A and the $^{32}\text{S}^-$ SIMS signal is presented in Fig. 7 B (red). Both images are overlaid in Fig. 7 C, where an increased $^{32}\text{S}^-$ signal (Fig. 7, B and C) and a depressed $^{12}\text{C}^{15}\text{N}^-$ signal (not shown) was detected at a majority of the highly fluorescent regions, confirming they contained an elevated abundance of TR-DHPE. Though the microdomains examined in this experiment differ in both complexity and chemical composition from those of biological relevance, these results demonstrate that the NanoSIMS 50 has the sensitivity and resolution to detect composition variations and identify their constituents.

**CONCLUSION**

By combining isotopic labeling strategies with SIMS, we have demonstrated that thin biological films can be detected and imaged by the NanoSIMS 50 without the necessity of bulky chemical labels that may change the properties of the labeled component. Isotopically labeled proteins can be imaged and detected with high sensitivity and selectivity. The high relative yield of F$^-$ secondary ions and relative ease of introducing F-Phe into proteins makes this an attractive atomic label for proteins. A lipid membrane can be distinguished from a protein adlayer through detection of the P$^-$ secondary ions, which exclusively originate from the single phosphorous atom within each lipid during sample interrogation with the NanoSIMS 50. Likewise, isotopically substituted lipid membranes can be differentiated from protein adlayers and even each other with the proper selection of unique isotopic labels. Our experiments indicate that quantitative information about the composition of a lipid mixture might be extracted from the intensity of the signal of interest, and small heterogeneous regions within a single membrane can be detected, and components that reside there identified, with the NanoSIMS 50. We expect that rapid progress in multifaceted isotopic labeling strategies, membrane patterning techniques, and quantitative methods will follow and prove to be a useful addition to the tools available for characterizing complex biological membranes.

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