

The Need for Speed in Matrix-Assisted Laser Desorption/Ionization Imaging Mass Spectrometry

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Abstract

Imaging mass spectrometry (IMS) has emerged as a powerful analytical tool enabling the direct molecular mapping of many types of tissue. Specifically, matrix-assisted laser desorption/ionization (MALDI) represents one of the most broadly applicable IMS technologies. In recent years, advances in solid state laser technology, mass spectrometry instrumentation, computer technology, and experimental methodology have produced IMS systems capable of unprecedented data acquisition speeds (>50 pixels/second). In applications of this technology, throughput is an important consideration when designing an IMS experiment. As IMS becomes more widely adopted, continual improvements in experimental setups will be important to address biologically and clinically relevant time scales.

Keywords: Imaging mass spectrometry, MALDI, high-speed imaging, TOF

Introduction

Mass spectrometry (MS) has long been recognized for its high sensitivity, high throughput, and molecular specificity. Advances in MS instrumentation have enabled the use of MS technology as an imaging modality.¹⁻² Specifically, matrix-assisted laser desorption/ionization imaging mass spectrometry (MALDI IMS), offers an untargeted approach to the regiospecific measurement of molecules in tissue specimens. The untargeted nature of this technology provides for the ability to simultaneously measure the complex array of molecular species present in biological tissue, producing molecular maps that can be correlated with anatomical tissue features without any required prior knowledge of the analytes present and without using any specialized reagents (*e.g.*, antibodies). MALDI IMS analyses have been successfully applied to the study of eye physiology,³⁻⁷ neurology,⁸⁻⁹ skin cancer,¹⁰⁻¹¹ pancreatic cancer,¹² and breast cancer.¹³⁻¹⁴

In a typical MALDI IMS experiment, a thinly sectioned tissue specimen is mounted onto a flat

target, such as a microscope slide, and then coated with a MALDI matrix (Figure 1). The MALDI matrix is typically a small organic molecule with strong absorbance at the wavelength of the MALDI laser and is applied to the sample in a manner that preserves the spatial integrities of the analytes of interest (*e.g.*, metabolites, lipids, peptides, proteins). A raster of the tissue surface is performed, generating a mass spectrum at each x, y coordinate (*i.e.*, a single pixel in 'microprobe' imaging mode).^{1, 15} Ion intensity maps can be constructed as a function of x, y position across the tissue surface for any ion of interest. Ions can then be identified through one or a combination of several techniques, including exact mass measurements,¹⁶⁻¹⁷ on-tissue tandem mass spectrometry (MS/MS),¹⁸⁻²³ or off-tissue liquid chromatography-tandem mass spectrometry (LC-MS/MS).²⁴

Early MALDI IMS experiments required 1-2 minutes/pixel for data acquisition, often requiring total analysis times of many hours or

even days.¹ However, recent advances in laser technology,²⁵⁻²⁷ data acquisition, and scanning methodology^{18, 23, 28-29} have provided for mass spectrometry systems which are capable of scanning at speeds as high as 50 pixels/second. The increasing use of IMS technology in many biological and clinical applications necessitates that continuing improvements in experimental times be achieved in order to perform IMS investigations on reasonable time scales.

Experimental

Transverse and coronal rat brain (Pel-Freez Biologicals, Rogers, AR) specimens were cryosectioned at 10 μm and thaw-mounted onto indium-tin oxide (ITO) slides. A custom built sublimation apparatus was used to apply a 1,5-diaminonaphthalene (DAN, Sigma Aldrich, St. Louis, MO) matrix layer (110° C, 8 min, ~50

mtorr).³⁰⁻³¹ IMS experiments were performed on either a SimulTOF 300 Tandem MALDI tandem time of flight (TOF/TOF) MS (SimulTOF Systems, Sudbury, MA) in positive ion reflectron MS mode or a RapifleX MALDI Tissue typer TOF MS (Bruker Daltonics, Billerica, MA) in negative ion reflectron mode. The SimulTOF instrument is equipped with a 349 nm, diode-pumped, frequency-tripled Nd:YLF laser capable of laser repetition rates up to 5 kHz. This system employs continuous laser raster sampling and was scanned using a 1 mm/second stage speed, 1 kHz laser frequency, and 50 hardware averaged shots per spectrum.²³ The RapifleX instrument is equipped with a Smartbeam 3D 10 kHz 355 nm Nd:YAG laser and was set to 200 shots per pixel using a single shot laser configuration and 85% focus setting.²⁴

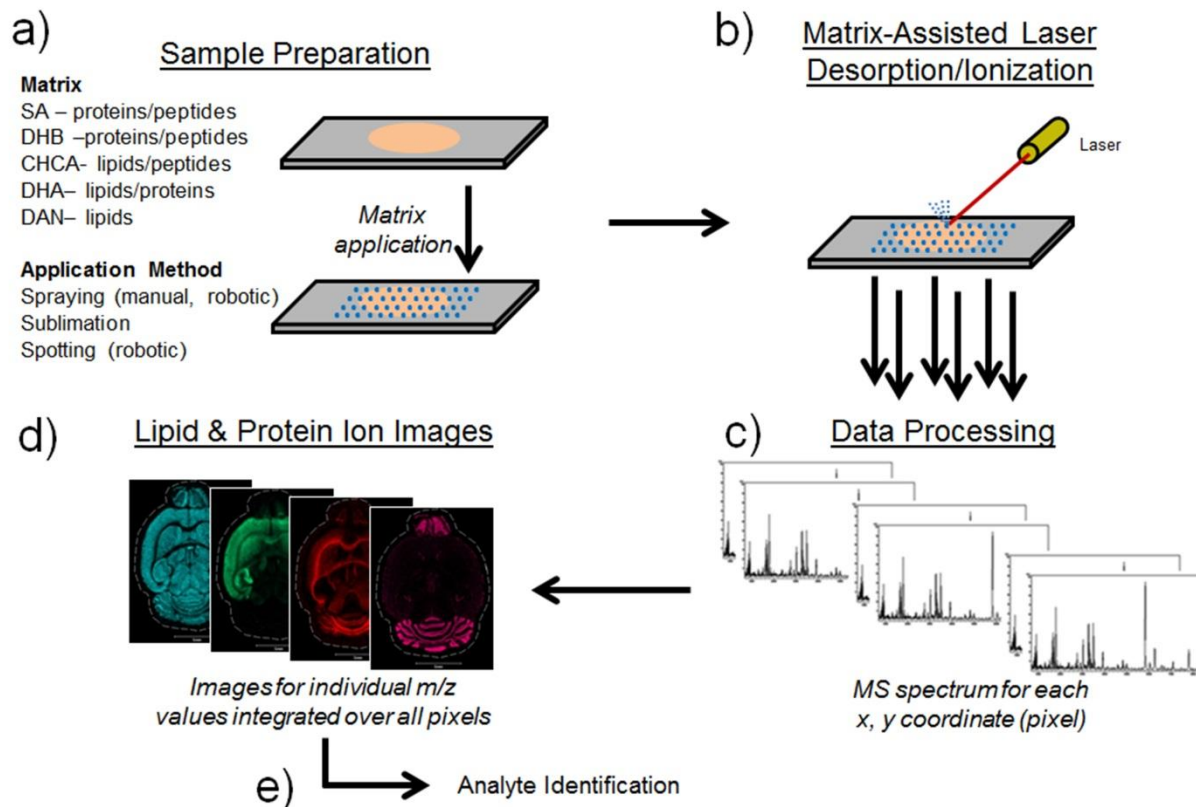


Figure 1. IMS workflow. a) Specimens are prepared for analysis by mounting thinly cut tissue sections onto slides. Matrix application is then performed via any number of methods prior to b) MALDI analysis. c) Mass spectra generated at each x, y position are then used to d) construct intensity map images for any single ion of interest. e) Analyte identification can be performed by one or a combination of several techniques.

Results

All analytical technologies experience trade-offs in various aspects of performance; as one particular characteristic is optimized, there are consequences that can affect the performance of other aspects of the technology. One such trade-off in IMS is the interplay between the amount of area sampled and the time required for data acquisition. The total number of pixels in an IMS experiment is a function of both the spatial resolution of the image (*i.e.*, defined by the pitch or spacing between each pixel) and the total imaged area. As area is a square function, the number of pixels required to sample larger and larger amounts of tissue at a defined spatial resolution increases rapidly. For example, a 1 cm by 1 cm area of tissue sampled at 100 μm spatial resolution requires 10,000 pixels to analyze the entire region, whereas a 5 cm by 5 cm area of tissue analyzed at the same spatial resolution requires 250,000 pixels. Similarly, as the spatial resolution of an experiment is adjusted to a finer setting, the experimental time also increases. An example of the consequences of throughput is shown in Figure 2 and Table 1. Assuming a 1.5 cm diameter circular tissue section and a pixel acquisition speed of 1 pixel/second, a fairly common acquisition speed, the region of tissue that can be sampled in a reasonable 12 hour acquisition time decreases dramatically as the desired spatial resolution is altered (Figure 2). Similarly, the amount of time required to sample the entire tissue section using a 1 pixel/second acquisition speed at 5 μm spatial resolution can take over 81 days (Table 1)! Certainly, this acquisition time is not practical for most biological and clinical applications. However, at

increased acquisition speeds, imaging the same area of tissue at finer spatial resolutions can again become practical. It is important to note that the acquisition times in Table 1 are reported for only a single tissue section; the effect on throughput is multiplicative when one considers the number of tissue sections in an experiment (*e.g.*, normal versus disease state tissue comparisons), the number of technical replicates, and the number of biological replicates.

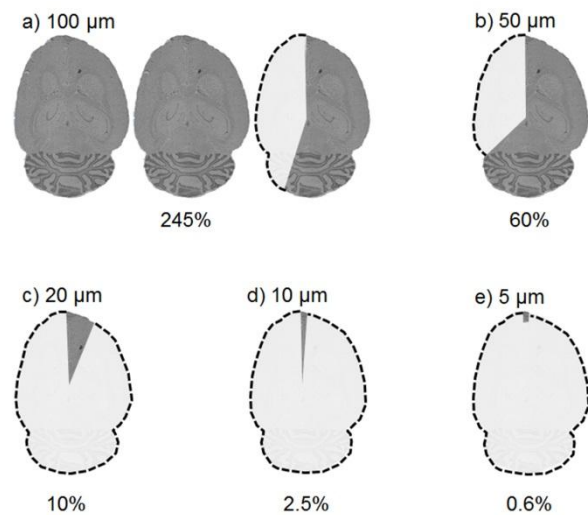


Figure 2. A depiction of an IMS experiment showing that, given a defined circular tissue area ($d=1.5$ cm, represented by one cartoon brain image, images not shown to scale) and a 1 pixel/second acquisition speed, the amount of tissue that can be sampled in 12 hours is dependent upon the spatial resolution of the experiment. Percentages indicate the approximate proportion of one brain section that can be measured at the indicated spatial resolution.

Table 1. The number of pixels required to sample a defined circular tissue area is determined by the spatial resolution of the experiment. Depending on the type of IMS instrumentation utilized, acquisition speeds can vary and will have a dramatic influence on the total analysis time.

Spatial Resolution (μm)	Number of Pixels Required to Sample Entire Tissue ($d=1.5$ cm)	Analysis Time (1 pixel/sec acquisition)	Analysis Time (40 pixels/sec acquisition)
100	~17,670	4.9 hours	7.4 minutes
50	~70,690	19.6 hours	29.5 minutes
20	~441,790	122.7 hours	3.1 hours
10	~1,767,150	20.5 days	12.3 hours
5	~7,068,580	81.8 days	49.1 hours

New advances in MS instrumentation have provided for systems capable of imaging at well above single pixel/second acquisition speeds. One such system utilizes continuous laser raster sampling to acquire data while continually firing the laser and scanning the stage (SimulTOF 300 Tandem).²³ This is in contrast to more traditional scanning methods that move the stage in discrete steps under a stationary laser, pausing the stage to fire the laser before turning the laser off and moving to the next raster step. By using continuous raster sampling, a 39,073 pixel image of a coronal rat brain section sampled at 50 μm spatial resolution was acquired in approximately 45 minutes (Figure 3). This corresponds to an acquisition rate of roughly 15 pixels/second. In this image, many brain substructures are clearly visible, including the corpus callosum, fornix, anterior commissure, and ventricles.³² It is also

evident that various ions in the mass spectrum display quite different spatial localizations. For example, the ion at m/z 734 is localized primarily to the ventricles (Figure 3c) while the ion at m/z 778 is primarily localized to the corpus callosum, fornix, and anterior commissure (Figure 3d). Given that this image was acquired in positive ion mode under sample conditions conducive to lipid ionization, these ions are likely forms of phosphatidylcholine ions, a class of glycerophospholipids that constitutes the major phospholipid present in most cellular membranes.³³ While the identity of these two ions can be hypothesized based on the experimental polarity and the nominal mass of the peak, further rigor should be employed when assigning identities to ions from these types of experiments (*vide infra*).

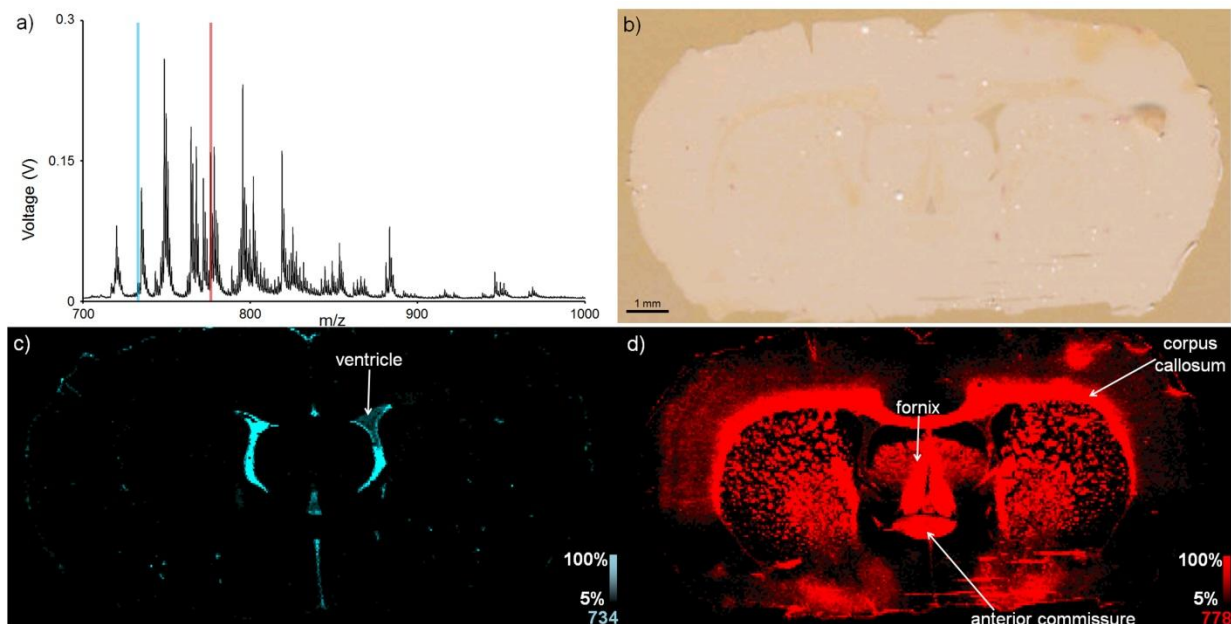


Figure 3. A 39,073 pixel MALDI image of a coronal rat brain section was sampled at 50 μm spatial resolution and was acquired in approximately 45 minutes. a) An average mass spectrum of 100 pixels shows the various lipid ions that were detected. b) A scanned optical image shows the tissue section following MALDI matrix application. Ion images of nominal masses c) m/z 734 and d) m/z 778 are plotted as m/z 734.6 ± 0.6 and m/z 778.6 ± 0.6 , respectively, and show differential localization in brain substructures.

Another high-speed system employs a 10 kHz laser and moves the laser and the stage independently from one another, allowing the stage to be moved continuously while still acquiring discrete pixel spots (Rapiflex Tissuetype).²⁴ Using this setup, a 644,134 pixel image of a transverse rat brain section sampled at 20 μm spatial resolution was acquired in approximately 345 minutes (Figure 4a). This corresponds to an acquisition rate of roughly 30 pixels/second. At this fine spatial resolution, the gray and white matter regions of the cerebellum

are clearly well resolved (Figure 4b). However, when conducting an IMS experiment, the user should not simply ask: “how high of a spatial resolution can I achieve?” Rather, the user should consider the experimental question for the biological or clinical task at hand and ask: “what spatial resolution is sufficient for this experiment?” By considering the level of image detail required and the area required to be imaged, the user can define a reasonable spatial resolution and analysis time.

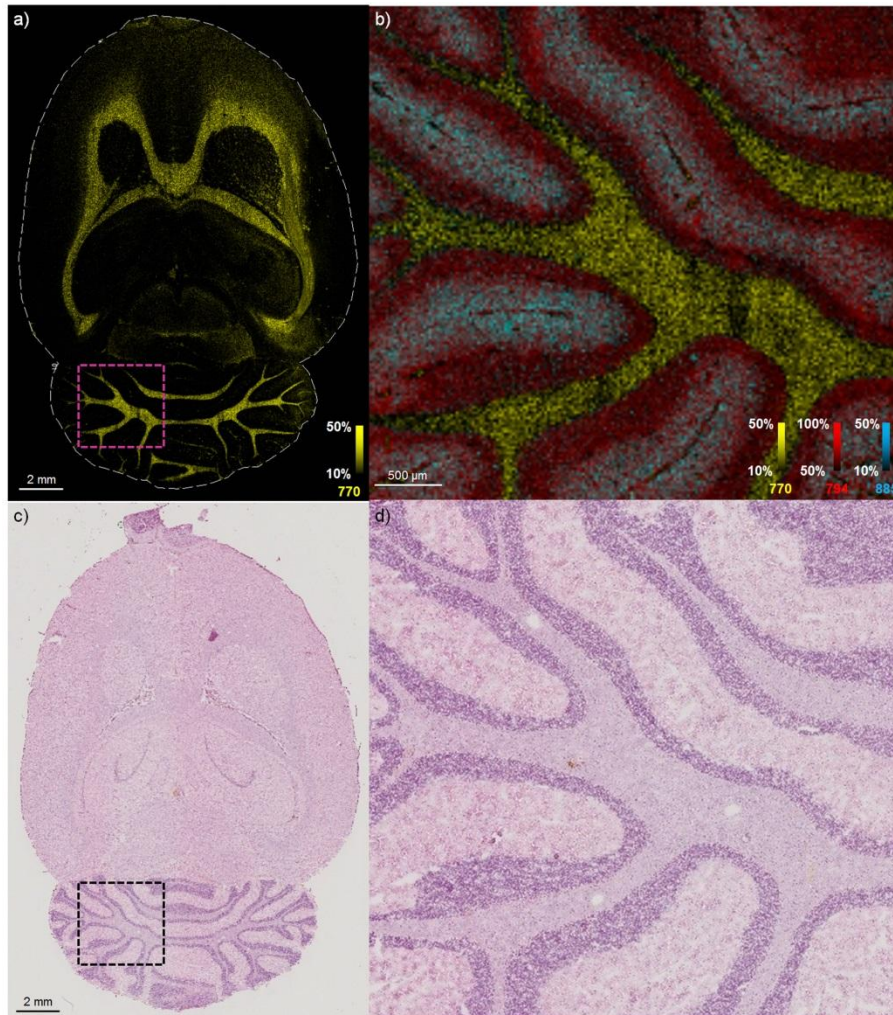


Figure 4. A 644,134 pixel MALDI image of a transverse rat brain section was sampled at 20 μm spatial resolution and was acquired in approximately 345 minutes. a) An ion image of nominal mass m/z 770 depicts the entire brain. b) Ion images of nominal masses m/z 770, m/z 794, and m/z 885 are shown in an enlarged region of the cerebellum highlighted in (a) using the pink box and are plotted as m/z 770.6 \pm 0.5, 794.7 \pm 0.5 and m/z 885.6 \pm 0.5, respectively, using root mean square (RMS) normalization. c) A hematoxylin and eosin stain of the tissue section is performed following MALDI IMS analysis. d) The enlarged region of the cerebellum is highlighted in (c) using the black box.

Such considerations not only have implications on throughput, but also on other aspects of IMS performance. For example, as the number of pixels in an image increases, the data file size also increases.³⁴⁻³⁶ In some cases, file sizes can be well over a terabyte in size! Additionally, at smaller laser diameters required for high-spatial resolution analyses, sensitivity can become a concern.³⁷⁻³⁹ While the images shown here were acquired on high-speed TOF systems, other systems with slower acquisition speeds can be extremely important for high-quality IMS studies. As mentioned above, the ability to identify an ion of interest should not rely solely on the measured nominal mass of the ion. Exact mass measurements using high-mass accuracy, high-resolving power mass spectrometers such as Fourier transform ion cyclotron resonance (FTICR) instruments, Orbitrap instruments, and quadrupole-TOF (Q-TOF) instruments can be used to identify the empirical formula of an ion of interest, facilitating analyte identification.^{16-17, 40} For example, the ion of nominal m/z 885 in Figure 4b was determined to have an exact m/z of 885.550 by tissue profiling on a 15T FTICR MS (Bruker Daltonics, Billerica, MA) (data not shown). Searching this experimental exact mass through an online lipid database (LIPID MAPS, Lipidomics Gateway, www.lipidmaps.org) allows for the identification of this lipid with an accuracy of <1 ppm as a phosphatidylinositol, PI(38:4), a glycerophospholipid commonly found in brain tissue.⁴¹ While extremely powerful, high-resolving power instrument platforms are often the most expensive and have larger data file sizes. MS/MS enabled ion trap, TOF, and hybrid instruments^{18, 20-22, 40, 42-43} as well as ion mobility-mass spectrometry (IM-MS) instruments^{40, 44} can also be used to provide structural information to aid in identification. While not always the case, many of these instruments have IMS acquisition speeds closer to 1 pixel/second, making them much slower than the high-speed TOF systems detailed above. However, diminished throughput is countered by an increase in molecular specificity with these instruments, an extremely important aspect of any MS experiment.

Conclusions

As IMS workflows are extended to more and more applications, it will be imperative that experimental setups provide biologically or clinically relevant time scales. In addition to continuing advances in instrumentation, advances in data acquisition also provide promising avenues for increasing experimental throughput. Histology-directed MS profiling experiments, where MS data is only acquired from several small predefined regions of the tissue, allow for dramatic increases in throughput.⁴⁵ Additionally, predictive imaging modalities that employ multivariate regression⁴⁶ or pan-sharpening algorithms⁴⁷ can allow for the mathematical combination of multiple imaging datasets, allowing for poorer spatial resolution IMS experiments to be sharpened to finer resolution IMS experiments. This 'image fusion' approach can also be used to provide out-of-sample ion distribution predictions for regions of unanalyzed tissue.⁴⁶ Combined with high-speed MS instrumentation, these methods and bioinformatics tools will be vital to the widespread adoption of IMS technology.

Acknowledgements

This work was sponsored by the National Institutes of Health/National Institute of General Medical Sciences under Award 5P41 GM103391-05. B.M.P. was supported by the National Institutes of Health/National Institute of Diabetes and Digestive and Kidney Diseases under Award F32 FDK105841A.

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