Mass spectrometry imaging with high resolution in mass and space

Andreas Römppp · Bernhard Spengler

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Abstract   Mass spectrometry (MS) imaging links molecular information and the spatial distribution of analytes within a sample. In contrast to most histochemical techniques, mass spectrometry imaging can differentiate molecular modifications and does not require labeling of targeted compounds. We have recently introduced the first mass spectrometry imaging method that provides highly specific molecular information (high resolution and accuracy in mass) at cellular dimensions (high resolution in space). This method is based on a matrix-assisted laser desorption/ionization (MALDI) imaging source working at atmospheric pressure which is coupled to an orbital trapping mass spectrometer. Here, we present a number of application examples and demonstrate the benefit of 'mass spectrometry imaging with high resolution in mass and space.' Phospholipids, peptides and drug compounds were imaged in a number of tissue samples at a spatial resolution of 5–10 μm. Proteins were analyzed after on-tissue tryptic digestion at 50-μm resolution. Additional applications include the analysis of single cells and of human lung carcinoma tissue as well as the first MALDI imaging measurement of tissue at 3 μm pixel size. MS image analysis for all these experiments showed excellent correlation with histological staining evaluation. The high mass resolution \( R = 30,000 \) and mass accuracy (typically 1 ppm) proved to be essential for specific image generation and reliable identification of analytes in tissue samples. The ability to combine the required high-quality mass analysis with spatial resolution in the range of single cells is a unique feature of our method. With that, it has the potential to supplement classical histochemical protocols and to provide new insights about molecular processes on the cellular level.

Keywords    Mass spectrometry imaging · Tissue analysis · High-resolution mass spectrometry · Accurate mass measurements · Mass spectrometry-based histology · MALDI mass spectrometry

Abbreviations

DHB    Dihydroxybenzoic acid
ESI    Electrospray ionization
H&E    Hematoxylin and eosin staining
LC    Liquid chromatography
MALDI    Matrix-assisted laser desorption/ionization
MS    Mass spectrometry
MS/MS    Tandem mass spectrometry
PC    Phosphatidylcholine
Ppm    Parts per million
RMS    Root mean square
SIMS    Secondary ion mass spectrometry
SM    Sphingomyelin
TOF    Time-of-flight (mass spectrometer)

Introduction: Mass spectrometry imaging

Mass spectrometry (MS) is a universal and prominent tool in analytical chemistry. This method can provide qualitative as well as quantitative information for a wide variety of compound classes. The molecular mass of a compound, calculated from the \( m/z \) value of the corresponding ion, can
be used for (tentative) identification. Ion fragmentation experiments (often called ‘tandem MS’ or ‘MS/MS’ experiments) provide information about the molecular structure of an analyte. Peak intensities in mass spectra can be calibrated for analyte quantitation. Mass spectrometry imaging combines these capabilities with information on the location of analytes within a sample. The spatial distribution can provide valuable additional information about the function and properties of an analyte, which would be lost in typical MS measurements of bulk (homogenized) samples. The first spatially resolved measurements in mass spectrometry were obtained with laser desorption ionization (LDI) in the 1970s (Hillenkamp et al. 1975a, b), but were limited to the analysis of smaller ions, predominantly of inorganic compounds. The investigation of larger (bio)molecules by mass spectrometry was made possible by the introduction of ‘soft ionization methods,’ namely electrospray (ESI) (Fenn et al. 1989) and matrix-assisted laser desorption/ionization (MALDI) (Karas et al. 1985; Karas and Hillenkamp 1988) in the 1980s. The first imaging application of MALDI was described in 1994 (Spengler et al. 1994). These experiments demonstrated that larger organic molecules such as peptides, which are thermally labile and which require matrix assistance, can be detected in a spatially resolved manner using a highly focused pulsed ultraviolet (UV) laser beam. This concept was taken up and extended to the analysis of tissue in the following years (Caprioli et al. 1997; Stoeckli et al. 2001).

Today mass spectrometry imaging is one of the most active fields in mass spectrometry. While MALDI and secondary ion mass spectrometry (SIMS) (Boxer et al. 2009) are the most widely used ionization techniques for mass spectrometry imaging, a number of alternative techniques have been developed in recent years. In particular, atmospheric pressure ionization techniques such as desorption electrospray (DESI) (Eberlin et al. 2010), low-temperature plasma (LTP) (Liu et al. 2010) and laserspray ionization (Richards et al. 2011) are increasingly operated in imaging mode. MALDI can also be operated at atmospheric pressure (Laiko et al. 2000), but it has mostly been employed in high vacuum ion sources.

A schematic representation of a MALDI mass spectrometry imaging experiment is shown in Fig. 1. The sample (tissue section) is covered with matrix in order to allow desorption and ionization of bioorganic analytes. A laser beam illuminates a defined area of the sample and desorbs and ionizes material from the surface (Fig. 1a). The resulting ions are transferred to the mass spectrometer, and a mass spectrum is acquired (Fig. 1b). Subsequently, the sample is moved by a defined distance, and the next position on the sample is analyzed the same way. Several thousand spectra are typically acquired sequentially in this manner. MS images of a selected analyte peak are generated after the measurement by extracting the signal intensity within a certain $m/z$ window. The intensities are plotted as gray scale values for each pixel in a grid representing the corresponding positions on the sample (Fig. 1c). These ‘MS images’ or ‘selected ion images’ represent the spatial intensity distribution of a certain $m/z$ signal, which can be assigned to a certain compound. Separate MS images can be generated for each signal in the mass spectrum. Therefore, mass spectrometry imaging is an ‘untargeted’ and multiplexed method. All desorbed and ionized compounds in the sample are detected, regardless of being known or unknown, expected or unexpected. This is a major advantage compared to other imaging techniques such as histochemical staining which requires the availability of a suitable antibody.

A number of compound classes have been analyzed by MALDI imaging including peptides (Chen et al. 2010; Stoeckli et al. 2002), proteins (Chaurand et al. 1999; Reyzer and Caprioli 2007), lipids (Jackson et al. 2005; Trim et al. 2008) and drug compounds (Stoeckli et al. 2007a). More details about MS imaging of these compound classes are covered in the respective sections below.

Today mass spectrometry imaging is used in a broad range of applications. The vast majority of studies are focusing on tissue originating from animal models or on human clinical tissue originating from surgical interventions. The targeted diseases include Parkinson’s (Stauber et al. 2008) and Alzheimer’s disease (Rohner et al. 2005) as well as Duchenne muscular dystrophy (Benabdellah et al. 2009b). Numerous studies of clinical tissue focused on cancer, for example prostate cancer (Cazares et al. 2009), breast cancer (Balluff et al. 2010), gastric cancer (Balluff et al. 2011a), adenocarcinoma (Djidja et al. 2009) and ovarian cancer (El Ayed et al. 2010). Additional topics in MS imaging studies included the analysis of plants (Lee et al. 2012; Kaspar et al. 2011; Peukert et al. 2011), insects (Vanickova et al. 2012) and microorganisms (Esquenazi et al. 2008; Yang et al. 2009).

A number of review articles about mass spectrometry imaging have been published in recent years. An extensive overview that covers instrumental topics as well as application areas is found in McDonnell and Heeren (2007). The main ionization techniques, namely MALDI, SIMS and DESI, are compared in Pól et al. (2010). The perspectives of MS imaging for clinical research focusing on protein analysis are covered in Rauser et al. (2010a) and Balluff et al. (2011b). A review by Schwamborn is specifically targeted at biomarker discovery approaches by MALDI

\textsuperscript{1} The term ‘MS image’ will be used in this work to describe the outcome of an MS imaging experiment in a general meaning. ‘Selected ion image’ will be used when referring to the spatial intensity distribution of a specific ion.
imaging (Schwamborn 2012). Multi-modal approaches and details on matrix application methods are included in Chughtai and Heeren (2010). Watrous et al. (2011) cover current applications and future perspectives of MS imaging in biological research.

In this review, we discuss MALDI-based mass spectrometry imaging with a focus on spatial resolution and molecular information content. We describe the methodology and application of a recently developed mass spectrometry imaging method with high resolution in mass ($R = 30,000$) and space (pixel size 3–10 μm).

**Critical parameters in mass spectrometry imaging**

Mass spectrometry imaging is a highly complex and interdisciplinary analytical technique. This section covers selected fundamental and practical aspects that have a significant effect on the outcome of an MS imaging experiment.

**Sample preparation**

Sample preparation is probably the most critical step in mass spectrometry imaging. All procedures should be optimized as to retain the native spatial distribution and molecular identity of analyte compounds in the sample as far as possible. Mass spectrometry imaging experiments are typically performed on (thin) tissue sections. However, classical embedding material for sectioning such as ‘optimal cutting temperature’ compound (OCT) should be avoided if possible. These polymers often result in substantial ion suppression and background signals which can severely interfere with mass spectrometric analysis. Therefore, cryosections of tissue without embedding are used if possible. More delicate samples can be prepared with embedding materials that have been reported to be compatible with mass spectrometry imaging, this includes carboxymethyl cellulose (CMC) (Kawamoto 2003), gelatine (Altealaar et al. 2005), a polymer compound (Strohalm et al. 2011) and tragacanth gum (Brignole-Baudouin et al. 2012).

The second crucial step in sample preparation is the application of matrix which is necessary to desorb and ionize sample components. The matrix needs to interact with the sample (surface) in order to extract analytes from the tissue and incorporate them into matrix crystals. At the same time, spatial integrity of the sample has to be preserved and analyte migration should be limited as far as possible. A number of sample preparation methods have been developed for MALDI imaging. A simple way of matrix application is spraying of a concentrated matrix solution by means of a thin-layer chromatography (TLC) sprayer (Stoeckli et al. 2007b). This method provides...
effective analyte incorporation, but can also lead to substantial spatial spreading of analytes if spraying conditions are not carefully controlled. An increasing number of commercial matrix application devices have been developed recently. They include the ‘ImagePrep’ (Bruker Daltonik, Bremen, Germany), which utilizes ‘vibrational vaporization’ to generate a spray of matrix droplets (Schuenenberg et al. 2007), and the ‘SunCollect’ sprayer (SunChrom, Friedrichsdorf, Germany), which employs a pneumatic sprayer. Both systems can achieve matrix crystal sizes below 50 µm in diameter. Spotting devices for MALDI matrix include the ‘CHIP-1000’ (Shimadzu Corp., Kyoto, Japan), which is based on piezoelectric technology, and a spotter from Labcyte (Sunnyvale, CA, USA), which is based on acoustic droplet deposition (Aerni et al. 2006). These systems provide matrix deposition spots of about 200 µm in diameter. High spatial resolution in MALDI imaging requires matrix crystals which are in the range of the laser spot size, that is the intended pixel size. Therefore, a two-step approach including matrix sublimation and subsequent recrystallization was developed by Bouschen et al. (2010). This method provides homogeneous matrix coverage and allows MALDI imaging at 1–2 µm spatial resolution. Similar approaches were developed by Thomas et al. (2012) and Yang and Caprioli (2011), which were applied to lipids and proteins, respectively. A pneumatic spraying device capable of obtaining matrix crystals below 10 µm was developed by Bouschen et al. (2010).

Irrespective of the matrix application method, a certain level of experience and practice is needed in order to obtain suitable results. Great care should be taken during each step of sample preparation, and experimental parameters should be adapted for each application (sample type and targeted compounds). This includes the choice of matrix which has a significant effect on crystal size and the set of compound classes that can be detected. The first MALDI imaging experiments were performed with 2,5-dihydroxybenzoic acid (DHB), and this matrix is still used in many studies, mainly for lipids and small molecules (Debois et al. 2010; Groseclose et al. 2007; Spengler and Hubert 2002; Spengler et al. 1994). α-Cyano-4-hydroxycinnamic acid (CHCA or HCCA) is also mainly used for small molecules, especially for peptides (Schwartz et al. 2003) and drug compounds (Prideaux et al. 2011; Stoeckli et al. 2007a). Proteins are typically analyzed with sinapinic acid (SA, 3,5-dimethoxy-4-hydroxycinnamic acid) (Grey et al. 2009; Leinweber et al. 2009; Yang and Caprioli 2011). While most MALDI imaging experiments were performed in positive ion mode, an increasing number of studies in negative ion mode has been reported, often using 9-aminonocacidine (9AA) as matrix (Shroff et al. 2008; Benabdellah et al. 2009a; Burrell et al. 2007). Matrix compounds that can be used in positive as well as in negative mode include 4-nitroaniline and 1,5-diaminonaphthalene (Thomas et al. 2012).

Mass spectral data

An important factor in generating MS images is the ‘bin width,’⁴ which defines the integration range on the \(m/z\) scale for a certain imaged ion signal. Intensity values for pixels in the MS image are calculated by integrating a small \(m/z\) window within the corresponding mass spectrum. Intensity values of two peaks are added up if they fall in the same bin. Thus, the bin width has significant influence on the generated MS image and has to be chosen according to the complexity of the sample and the resolution and accuracy of the mass spectral measurement (examples are shown in Figs. 6, 8). Consequently, the choice of the mass analyzer has significant impact on the quality of MS images.

The most widely used detectors for MALDI imaging are time-of-flight (TOF) mass spectrometers. These instruments are sensitive and are available in many (bio)analytical laboratories. They have been employed for a wide range of applications, such as MS imaging of lipids (Benabdellah et al. 2009b; Thomas et al. 2012; Chen et al. 2009), drugs/metabolites (Benabdellah et al. 2009a; Stoeckli et al. 2007a) and peptides/proteins (Stoeckli et al. 2002; Mange et al. 2009; Lemaire et al. 2006, 2007b; Francese et al. 2009; El Ayed et al. 2010; Rohner et al. 2005). Time-of-flight mass spectrometers in reflection mode reach a mass resolving power of up to \(R = 30,000\) and mass accuracies better than 5 ppm. However, this performance is only obtained for measurements of homogenized thin-layer samples. In MS imaging experiments, the three-dimensional structure of the sample affects ion flight times and results in significantly lower mass resolution and mass accuracy. Height differences, which cannot be entirely eliminated, lead to inhomogeneity in the acceleration field and thus to a shift in detected flight time (and the corresponding \(m/z\) value). Mass accuracy of axial TOF mass spectrometers under tissue imaging conditions is in the range of several tens of ppm if the measurement is controlled carefully. However, mass deviations up to \(m/z\) 0.5 are also not uncommon in MS imaging experiments. One way to reduce these detrimental effects are orthogonal TOF systems where the ions are accelerated perpendicularly to the initial extraction direction. These instruments are typically equipped with a quadrupole filter in front of the time-of-flight analyzer (QTOF) to allow MS/MS experiments (Loboda et al. 2000). Mass accuracies reported for MS imaging are in the range of 20–30 ppm.

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⁴ This parameter is also described as ‘bin size,’ ‘integration range’ or ‘mass window.’ The term ‘bin width’ will be used in this work.
et al. 2010). Applications of these systems include drug compounds (Stoeckli et al. 2007a), peptides (Djidja et al. 2009; Djidja et al. 2010; Trim et al. 2010) and lipids (Prideaux and Stoeckli 2012; Trim et al. 2008). MALDI–QTOF systems have also been combined with ion mobility spectrometry separation (Stauber et al. 2010; Djidja et al. 2009). This method adds an additional gas-phase separation step (based on the collisional cross section), which can help to resolve the complexity of biological samples.

The highest mass resolving power and mass accuracy are obtained with Fourier transform ion cyclotron resonance (FTICR) mass spectrometers (Marshall and Hendrickson 2002; Amster 1996; Marshall 2000; He et al. 2001; Römpp et al. 2005). These analyzers have the advantage that the cyclotron frequency (and thus the measured m/z value) is independent of the kinetic energy of the ions. This results in mass accuracies of better than 1 ppm which can also be obtained in MS imaging experiments. FTICR mass spectrometers have been used for MS imaging of peptides (Taban et al. 2007) and drug compounds (Cornett et al. 2008). Spatial resolution in these cases was between 200 and 300 μm. Sensitivity was lower than for regular MALDI–TOF instruments. The combination of an FTICR mass spectrometer with a MALDI imaging ion source at a spatial resolution of 7 μm was demonstrated for artificial targets by Koestler et al. (2008).

Development of the orbital trapping mass spectrometer (Orbitrap, Thermo Fisher Scientific GmbH, Bremen, Germany) greatly improved limits of detection (LOD) for accurate mass measurements (Makarov 2000; Makarov et al. 2006). This system is based on the detection of ions trapped in an electrostatic field and does not require a superconducting magnet. Apart from easier handling and lower maintenance costs, this also allows for more efficient ion transmission. Ions are accumulated in a so-called C-trap which is located only centimeters away from the analyzer cell. Orbital trapping mass spectrometers are widely used in bioanalytical applications, typically coupled to liquid chromatography (LC) systems by an electrospray ionization source (ESI). They were also used to image lipids (Landgraf et al. 2009) and peptides (Chen et al. 2010). Orbital trapping mass spectrometers were used for all our measurements which are presented in this review.

Spatial resolution

Spatial resolution in MALDI imaging experiments primarily depends on the size of the laser focus. While the majority of MALDI imaging studies were performed at a spatial resolution between 50 and 200 μm per pixel, there are a number of notable exceptions. As mentioned above, the first MALDI imaging results were obtained at 1 μm pixel size (Spengler et al. 1994; Spengler and Hubert 2002). The obtained MS images showed in detail the spatial distribution of peptides and matrix compounds (dihydroxybenzoic acid, DHB) in the crystals of a standard MALDI preparation. The same system was used to image cancer cells at 1 μm step size revealing the distribution of several compounds up to m/z 5,000 (Bouschen et al. 2010).

A similar optical setup built by Bernhard Spengler was used by Chaurand et al. (2007) to image proteins at 10 μm pixel size. Recently MALDI imaging at 10 μm pixel size using (slightly modified) commercial systems was demonstrated for lipids (Thomas et al. 2012) and proteins (Yang and Caprioli 2011).

The pixel size of the MS image is determined by the distance that the sample is moved between two measurements. This step size is usually chosen in the same range as the laser focus diameter in order to avoid overlapping of analyzed areas. An alternative is to move the sample by less than the focus size resulting in ‘oversampling.’ If the laser ablates all material of the sample, this method can generate reasonable MS images with a pixel size that is several times smaller than the laser focus size. This was demonstrated for lipids at 15 μm pixel size (Snel and Fuller 2010) and for peptides prepared from cells (Jurchen et al. 2005). It has to be noted that oversampling improves pixel resolution on the cost of sample loss. This is because a large-focus beam has a much lower beam intensity slope at the focus edges than a small-focus beam. The shallow intensity gradient significantly leads to premature sample evaporation outside the ionization zone and thus to a lower overall ion yield.

Another method of obtaining MS image pixels smaller than the laser spot size is the so-called microscope mode (Altelaar et al. 2007; Froesch et al. 2010; Luxembourg et al. 2004). In this case, the laser irradiates an area of 200 μm in diameter and the ions are extracted and detected in a spatially resolved manner. This method can deliver a pixel size of about 4 μm, but practical issues such as detection speed make it difficult to acquire MS imaging data over a larger mass range with this technique (Stauber et al. 2008).

Mass spectrometry imaging with high resolution in mass and space

MS images contain mass spectral and spatial information. Consequently, data need to be acquired in high quality for both dimensions. As described above, high spatial resolution and high mass resolution were achieved from MALDI imaging experiments earlier, but only in separate experiments. We have recently developed a method that for the first time combines a spatial resolution in the low micrometer range and high mass accuracy for the analysis of biological samples. Phospholipids (Römpp et al. 2010a), neuropeptides (Guenther et al. 2011) and drug compounds (Römpp et al. 2011b) were imaged with accurate mass at a...
pixel size between 5 and 10 μm. Spatial resolution for trypic peptides after on-tissue digestion is lower, but 50 μm pixel size was demonstrated for this application (Schober et al. 2012a). In all cases, this method provided a significant improvement in spatial resolution and/or reliability of identification compared to published methods as discussed below.

**Methodology**

An important aspect in order to obtain high-quality and reproducible results from tissue samples was to systematically evaluate and optimize every step of the analytical workflow for the specific application. This included sample selection, sample preparation, instrumental measurement parameters, but also acquisition, processing and interpretation of data. The following describes general aspects of the workflow which are common to most of our studies. More specific details and additional information are provided below in the individual sections covering the different applications.

**Sample preparation**

Tissue samples were snap-frozen immediately after the animal was sacrificed and stored at −80 °C. No embedding or fixation was used. Thin sections of tissue with a thickness of 10–20 μm were prepared with a microcryotome (HM 525, Thermo Scientific, Dreieich, Germany) and thaw-mounted on regular microscope glass slides. Conductive coating of glass slides, indium tin oxide (ITO), is not required, but was used in some applications. Tissue sections were covered with matrix by a pneumatic sprayer (Bouschen et al. 2010). 2,5-Dihydroxybenzoic acid (DHB) at a concentration of 30 mg/mL in acetone/water (0.1 % TFA) 1:1 v/v was used as matrix solution. Matrix crystals in the low micrometer range were obtained with this system. Tissue sections were stained after the MS imaging measurement in order to allow for a detailed analysis of histological features. This included staining with hematoxylin and eosin (H&E), toluidine and Luxol fast blue (Riedelsheimer et al. 2010).

**Mass spectrometry imaging**

Experiments were performed using a home-built atmospheric pressure scanning microprobe matrix-assisted laser desorption/ionization (AP-SMALDI) imaging source attached to a linear ion trap/Fourier transform orbital trapping mass spectrometer (LTQ Orbitrap Discovery, Thermo Scientific GmbH, Bremen, Germany). The setup of the AP-SMALDI imaging source is described in detail elsewhere (Koestler et al. 2008). The source works at atmospheric pressure and thus allows analyzing biological samples in their native form without the need for dehydration. A nitrogen laser (wavelength 337 nm) with a repetition rate of 60 Hz was used. The laser beam was focused onto the sample by a centrally bored objective lens. In this system, the optical diameter of the focused laser beam is 8.4 μm (as defined by a decrease in beam intensities at the edges to 1/e² (13.5 %) of its peak intensity). The diameter of the ablation area (which defines the effective spatial resolution) depends on the laser fluence on the sample and is typically adjusted to 5 μm. A detailed characterization of the optical system is described in Guenther et al. (2010). The sample is moved in front of the stationary capillary/lens setup by an x, y, z-stage with a precision better than 1 μm. MS imaging data was acquired with a mass resolving power of 30,000 at m/z 400 in profile mode (i.e. mass peaks were not centroided for data reduction). Ions from 30 laser pulses were accumulated in the linear ion trap for each mass spectrum. Automatic gain control (AGC) was disabled during the measurement, and ion injection time was set to a fixed value (typically 650 ms). Mass accuracy was improved by applying internal calibration using the ‘lock mass’ feature of the Orbitrap mass spectrometer. The dimer [2DHB − H_2O + H]^+, trimer [3DHB − 2H_2O + NH_4]^+ and pentamer [5DHB − 4H_2O + NH_4]^+ of DHB were typically used as reference masses for this procedure. This resulted in a mass accuracy of better than 2 ppm (RMS) achieved for MS imaging experiments of tissue. Mass spectrometer, laser and sample stage are controlled and synchronized by a microcontroller and home-built software. More details on the instrumental setup and experimental parameters can be found in Römpp et al. (2010a).

**Software for image generation**

MS images were generated by the software package MI-RION (Paschke et al. submitted). The imaging software is able to create ion images from any of the detected m/z values with any selected mass window (bin width). MS images can be generated semiautomatically based on a number of selection parameters such as mass range and pixel coverage. Alternatively selected ion images of a defined m/z value can be generated manually. Up to three different ion images are overlayed in red–green–blue (RGB) images by the software to display different ion species in parallel.

**Reporting of MS images and mass spectral data**

‘Images’ in mass spectrometry imaging are constructed in the computer from individual mass spectra. Consequently,
the information content of the resulting image can only be as good as that of the underlying mass spectral data. Therefore, we take extra care to ensure the quality and validity of our reported MS imaging data. If not explicitly stated differently, all data presented in this work adhere to the following rules. Intensity values in selected ion images were normalized to the highest intensity within the image, measured for each ion species separately. MS images do not include any spatial interpolation, smoothing or pixel-wise normalization, in order to allow for an unbiased evaluation. No overlay with optical images is used for MS image presentation. Black-and-white images of selected raw data are usually reported in the supporting information of the original research publications. Step size of the stage movement is identical in x and y direction, that is square pixels are measured and displayed.

The terms ‘mass deviation,’ ‘mass accuracy’ and ‘maximum mass deviation’ are used according to the definition by Zubarev and Mann (2007). All measurements are based on high-resolution mass spectrometry data (mass resolving power \( R > 20,000 \) @ \( m/z \) 700), and images are generated with a bin width of \( \Delta m/z = 0.01 \). In almost all cases, this proved to be necessary in order to resolve the complexity of biological samples. Mass spectra acquired from single pixels are presented in order to allow an evaluation of the quality of the underlying measurements. Mass accuracy is not (solely) discussed on the basis of selected mass spectra, but is reported as values of the root mean square of \( m/z \) deviations across the whole imaging measurement. Accurate mass measurements (RMS \( \leq 3 \) ppm) are used for assignment of imaged compounds. Identification is confirmed by additional MS/MS experiments whenever possible. All these factors are relevant in order to report and evaluate the data in a reliable and comprehensive way.

**Phospholipids**

Phospholipids are major structural components of all cell membranes and play a crucial role in the cell metabolism of all organisms. Phospholipid derivatives play an important role in signal transduction and are thus of high relevance for numerous pathological processes (Abrass 2004; Lee et al. 2003). Changes in lipid metabolism are major factors in diabetes and cancer (Paradisi et al. 2010; Santos and Schulze 2012). Lipids have been analyzed by mass spectrometry imaging in a number of studies (Sparvero et al. 2010; Jackson et al. 2005; Benabdellah et al. 2009b; Thomas et al. 2012).

The first application of our high-resolution mass spectrometry imaging method in the field of tissue analysis was the analysis of phospholipids in mouse urinary bladder (Römpp et al. 2010a). A measurement with 10 \( \mu m \) pixel size is shown in Fig. 2a. The MS image shows excellent agreement with classical histological information (after toluidine staining, Fig. 2b). Certain phospholipid signals could be clearly assigned to different tissue regions (all lipids discussed here were detected as potassium adducts). The outer region of the urinary bladder was found to be characterized by the sphingomyelin compound SM (34:1)\(^3\) (\( m/z \) 741.5307, Fig. 2a, blue). This histological region could be further subdivided into the adventitial layer, the outermost thin layer as indicated by increased signal intensities of SM (34:1) and the detrusor muscle mainly consisting of smooth muscle tissue.

Following the muscle layer is the loose connective tissue of the lamina propria, characterized by intense signals of \( m/z \) 743.5482 (Fig. 2a, red). Colocalization with SM (34:1) displayed in blue results in the magenta color of this region in the RGB overlay. Toward the bladder lumen, a thin layer of subepithelial myofibroblasts was found to be characterized by again high signal intensities of SM (34:1) (Fig. 2a, blue). The high cell densities of both layers were expressed in the stained image (Fig. 2b). The innermost layer of the mucosa is composed of the bladder epithelium (urothelium), indicated by the phosphatidylycholine species PC (34:1) at \( m/z \) 798.5410 (Fig. 2a, green). This discussion shows that our MS imaging method can reproduce very detailed histological features, close to cellular resolution. In addition, it also provides highly reliable molecular information.

A mass spectrum acquired from a single 10-\( \mu m \) pixel from the urinary bladder imaging experiments is shown in Fig. 2c. Several peaks are labeled based on accurate mass measurements. Mass deviation of assigned peaks in this spectrum was below or equal to 1 ppm. Mass accuracy as determined over the whole measurement (10965 spectra) was 0.6 ppm (root mean square of all mass deviations) for PC (34:1) at \( m/z \) 798.5410. This demonstrates that accurate mass detection is stable over the whole measurement under imaging conditions. This is a prerequisite for reliable identification of compounds and for specific image generation. Another important factor that influences MS image quality is mass resolution. The effect of different bin widths used for image generation is discussed in detail in Römpp et al. (2010a), (Figure S5). MS images with \( \Delta m/z = 0.01 \) result in complementary spatial intensity distributions of two individual compounds, whereas a bin width of \( \Delta m/z = 0.1 \) (which is often used in MS imaging

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\(^3\) Lipids are denoted by the abbreviation for the respective compound class followed by the total number of C atoms and the number of double-bonds in the fatty acid chains. The chain lengths of the individual fatty acid residues and the location of the double-bonds cannot be resolved on the basis of accurate mass. This can be achieved through additional MS/MS measurements.
A second measurement of the same urinary bladder section (adjacent area) was measured with a pixel size of 5 \( \mu \)m. The selected ion images of two phospholipids in a region of two adjacent tissue regions are shown in Fig. 2d. This measurement also shows excellent agreement with histological staining by toluidine (Fig. 2e). The MS image provides sharp separation between the muscular layer \([m/z \ 741.5307, \text{SM (34:1), green}]\) and the urothel \([m/z \ 798.5410, \text{PC (34:1), red}]\). These data demonstrate that the effective analytical resolution is indeed in the range of 5 \( \mu \)m.

These results represented the first combination of high-resolution, high-accuracy mass spectrometry with a pixel size in the low micrometer range for mass spectrometry imaging of biological samples. Both features are necessary in order to obtain reliable molecular information from tissue sections on a cellular level. These results also show that phospholipids are highly specific for different cell types and can thus provide detailed histological information. This information can be used to differentiate and assign features in tissue section which might not be apparent when using classical histological techniques.

### Drug compounds

Detection of drug compounds is a major application of mass spectrometry imaging (Prideaux and Stoeckli 2012). The ability for label-free detection of and differentiation between drug compounds and metabolites are its major advantages compared to autoradiography, which is the routine method to investigate the spatial distribution of drug compounds in animal models (Solon et al. 2010). The spatial resolution in MS imaging measurements of drug compounds typically has been in the range of 200–500 \( \mu \)m, which is sufficient to analyze whole body sections (Khatib-Shahidi et al. 2006; Stoeckli et al. 2007a) or individual rat organs (Cornett et al. 2008), but insufficient to resolve detailed features in mouse organs. The highest spatial resolution reported before for drug compounds was 100 \( \mu \)m (Acquadro et al. 2009; Stoeckli et al. 2007a). MS imaging experiments with accurate mass detection for drug compounds were performed at 350 \( \mu \)m spatial resolution (Cornett et al. 2008). Drug compounds are usually detected...
in MS/MS mode in order to increase the specificity of the measurement and to prevent interference of matrix cluster ions which are typically abundant in the low mass range (Stoeckli et al. 2007a). The number of compounds that can be detected in one experiment is limited in this approach.

Following the promising results on phospholipids, we investigated the distribution of drug compounds by ‘mass spectrometry imaging with high resolution in mass and space (HR² MSI)’ (Römpp et al. 2011b). The anticancer drug imatinib (a tyrosine kinase inhibitor) was analyzed in mouse kidney at 35 μm pixel size. Selected ion images of phospholipids are shown in Fig. 3a. The distribution of these phospholipids corresponds very well to histological features of the kidney as determined by H&E staining (Fig. 3c). The distribution of the drug compound imatinib in the mouse kidney is shown in Fig. 3b. Imatinib ([M + H]⁺, m/z 494.2662) is displayed in green, while two phospholipids (red and blue) are included to indicate the histological features of the kidney. Identification and structural characterization of tissue regions, which usually requires examination of the H&E-stained section by a trained histologist, is possible based on the MS imaging data of lipids. This information can be used to determine the exact location of the targeted analyte in the mouse kidney. The colocalization of imatinib (Fig. 3b, green) and PC (40:6) (Fig. 3a, blue) indicates that imatinib is accumulated in the outer stripe of the outer medulla.

A mass spectrum acquired from a single 35-μm pixel is shown in Fig. 3d. The molecular ion peak ([M + H]⁺) of imatinib was detected with a mass deviation of −0.4 ppm and a mass resolution of R = 30,000 in this spectrum. The overall mass accuracy for imatinib in the imaging measurement was ±0.4 ppm. The overall mass accuracy for imatinib in the imaging measurement was ±0.4 ppm.
measurement calculated from 5862 individual spectra was 2 ppm (root mean square of mass deviations in all spectra). The identity of imatinib was verified by additional MS/MS experiments, which showed a characteristic fragmentation pattern.

A smaller area of an adjacent mouse kidney section was imaged with 10 μm pixel size in order to investigate the distribution of imatinib in more detail. The resulting MS images again showed excellent agreement with histological features of the mouse kidney and confirmed that imatinib is specifically accumulated in the outer stripe of the outer medulla [see Römpf et al. (2011b) for details]. This can be tentatively associated with a selective accumulation of drug molecules in the proximal tubules. Imatinib was detected with a mass accuracy of 1.8 ppm RMS (calculated from 5816 individual spectra) in this 10 μm measurement.

Mass spectrometry imaging with high resolution in mass and space (HR² MSI) provides detailed information on spatial distribution of compounds. The described measurements represented the first mass spectrometric imaging analysis of drug compounds at 10 μm pixel size. This constituted a 10-fold increase in spatial resolution compared to previous studies. At the same time, the measurements provided detailed information on histological features based on the spatial distribution of phospholipids and other endogenous compounds. Correlation of different images allowed for fast and easy interpretation of the drug compound distribution, and areas of accumulation could be directly linked to certain tissue regions.

**Neuropeptides**

Neuropeptides are important messenger molecules that control a wide range of regulatory functions and processes within an organism. They are formed in vivo by cleavage of precursor proteins and often undergo posttranslational modifications (PTM). These modifications directly affect the biological function of neuropeptides. Investigation of neuropeptides therefore requires specific detection and reliable identification. This can be difficult with histochemical methods as antibodies are not always able to differentiate PTMs. Previous MS imaging approaches for neuropeptide analysis included high spatial resolution (Altelaar et al. 2006, 2007), high mass resolving power and mass accuracy (Taban et al. 2007; Chen et al. 2010) or MS/MS capabilities (Chen et al. 2009; Verhaert et al. 2007; DeKeyser et al. 2007), but these features were never combined in one experiment.

Therefore the aim of our study (Guenther et al. 2011) was to demonstrate that MS imaging of neuropeptides in mammalian tissue is possible with high mass resolution and with MS/MS analysis at cellular resolution. The pituitary gland is one of the best known neuropeptide-secreting organs in mammals and was therefore chosen for our experiments. It consists of three distinct histological regions, called anterior, posterior and intermediate lobe. Each of these regions has different functions and produces different neuropeptides. In a first experiment, a section of a complete mouse pituitary gland (approximately 1.5 mm × 2.5 mm) was imaged at a pixel size of 10 μm using a mass range of m/z 200–4,000. Figure 4 shows the optical image (a) and examples for MS images of neuropeptides (b–d). The distribution of phosphatidylcholine PC (38:4) (shown in green) was used to indicate the overall shape of the pituitary gland. Neuropeptides originating from the same precursor were found to be colocalized. For example, vasopressin and two variants of copeptin, which are all formed from the prohormone AVP-NPII, were detected in the posterior lobe. The neuropeptides POMC 76–85 (γ-MSH), POMC 103–120 (joining peptide), POMC 124–136 (α-MSH) and POMC 141–162 (CLIP) were detected in the intermediate lobe, where the corresponding prohormone POMC is known to be synthesized. In addition, two modifications of α-MSH, the acetylated and diacetylated forms, were detected in this region. This demonstrates that posttranslational modification of neuropeptides can be differentiated with our method. The determined spatial distributions of all detected neuropeptides were in agreement with their expected biological function. In total, 10 neuropeptides were detected in this measurement.

A single-pixel (10 μm) mass spectrum acquired from the intermediate lobe is shown in Fig. 4e. Peptides were identified by comparing their accurate mass with theoretical values for known neuropeptides. A maximum mass tolerance of 2 ppm was used for this procedure. The identity of the neuropeptides was confirmed by additional MS/MS measurements which were obtained in profiling mode from selected areas of the pituitary gland. The identification of all 10 neuropeptides was confirmed by detailed evaluation of product ion peaks [see Guenther et al. (2011) for details]. Furthermore, the spatial distribution of two peptides was investigated by MS/MS in imaging mode at 10 μm spatial resolution. The resulting product ion images were in excellent agreement with the distributions of their peptide precursor ions, confirming their identity and excluding possible analyte interferences.

In an attempt to resolve the fine structure of the pituitary gland, a smaller area covering all three lobes (indicated by a yellow square in Fig. 4b) was imaged at 5 μm pixel size. An overlay of selected ion images of the neuropeptides is shown in Fig. 4f. Diacetyl-α-MSH (POMC 124–136) was found to be located in the intermediate lobe (green). Vasopressin (AVP-NPII) was detected in the center of the posterior lobe (blue). Oxytocin (OT-NPI 20–28) was also
present in the posterior lobe, but was primarily detected in the outer region of this lobe. The different distributions of these two peptides are in agreement with the known origin of the corresponding prehormones and can be directly linked to physiological processes on a cellular level [see Guenther et al. (2011) for details].

This study demonstrated that MS imaging of neuropeptides in mammalian tissue is possible at cellular resolution. For the first time, such experiments were performed by combining accurate mass analysis, high mass resolving power and high spatial resolution in one experiment. Peptide sequences were verified by MS/MS experiments, including posttranslational modifications which cannot be differentiated by antibody staining methods.

**Proteins (on-tissue tryptic digestion)**

Proteins are one of the most studied compound classes in biomedical applications. The analysis of their spatial distributions can provide additional information about their role in physiological and pathological processes. Numerous studies have been published on MS imaging of proteins (Chaurand et al. 2007; Leinweber et al. 2009; Rauser et al. 2010a, b; Shimma et al. 2008; Walch et al. 2008). Spatial resolution for proteins is typically in the range of 100–250 μm. A recent study showed measurements at 10 μm pixel size (Yang and Caprioli 2011). However, protein analysis by MS imaging still faces a number of problems (McDonnell et al. 2010). Direct
High mass accuracy MS imaging of tryptic peptides on tissue

The goal of our first study involving tryptic peptides was to improve the reliability of identification by acquiring mass spectra with accurate mass (Schober et al. 2011). In addition, a combination of on-tissue tryptic digestion and complementary LC–MS/MS analysis of the corresponding proteome was applied to increase the number of identified peptides. The highest mass accuracy reported earlier for imaging of tryptic peptides was 30 ppm on a time-of-flight system (Djidja et al. 2010), while most studies reported values in the range of 100 ppm.

In our study, trypsin solution was deposited manually on mouse brain tissue resulting in trypsin spots of about 1 mm diameter (Schober et al. 2011). MS images at a pixel size of 100 μm were acquired from the mouse brain section. In contrast to the other MS imaging experiments discussed in this work, these sections were analyzed with the commercial MALDI source manufactured by Thermo Fisher Scientific (Bremen, Germany). Selected ion images of two peptides are shown in Fig. 5a, b. In the next step, a mouse brain tissue section, adjacent to the one used for MALDI imaging, was analyzed by LC–ESI–MS/MS in order to identify proteins which are present in this tissue. Prior to analysis, this sample was homogenized and fractionated by ultracentrifugation resulting in four subcellular fractions: cytosol, mitochondria/nuclei, endoplasmatic reticulum/golgi and plasma membranes. More than 1,100 peptides corresponding to 500 proteins were identified in this sample. The data from MALDI imaging (tissue section) and from the LC–ESI–MS/MS experiment (homogenate sample) were combined as shown in Fig. 5c, d. In analogy to our previous work (Römpp et al. 2007), a maximum mass deviation of 3 ppm between m/z values of MALDI imaging peaks and masses of peptides identified by LC–ESI–MS/MS was accepted in this study. The two peaks detected by MALDI imaging (Fig. 5c) match with masses of peptides that were identified in the mouse brain tissue by LC–ESI–MS/MS (Fig. 5d). Therefore, the corresponding images were assigned to these two peptides. An unequivocal match of MALDI imaging and LC–ESI–MS/MS data is only possible with a mass accuracy in the low ppm range for both measurement modes, while less accurate measurements would lead to ambiguous assignments. For example, at 30 or 100 ppm mass tolerance, more than one possibility of linking the two measurements existed and would have obstructed an unequivocal interpretation of imaging results [see Schober et al. (2011) for details].

The combination of MALDI imaging and LC–ESI–MS/MS led to the identification of more than 147 peptides corresponding to 101 proteins. More than 75% of these proteins had a molecular weight of more than 30 kDa and
would therefore not be accessible for a top-down approach imaging intact proteins directly on tissue. A total of 24 proteins were detected with two or more peptides and could be considered as reasonable matches. These numbers are in line with comparable studies (Djidja et al. 2009; Gustafsson et al. 2010), but the reliability of identification is significantly higher in our measurements due to accurate mass data.

This study showed that the identification of tryptic peptides without accurate mass and without complementary approaches such as bulk identification methods can easily produce misleading results. In light of these results, the identification results of previous studies have to be considered with care if no independent histochemical confirmation was performed. In particular, in the case of digested tissue sections of high complexity, reliable identification of proteins significantly benefits from high mass accuracy of measurements. This proof-of-concept study was based on low-spatial-resolution imaging. Results and implications with respect to accurate mass measurements are, however, equally relevant for experiments with smaller trypsin spots as demonstrated in the following section.

**Improved spatial resolution for MS imaging of tryptic peptides**

The second study on MS imaging of tryptic peptides (Schober et al. 2012a) was focused on increasing the accessible spatial resolution while maintaining the high reliability of the identification step. The critical step in increasing spatial resolution for tryptic peptides is the application of the enzyme solution (usually trypsin). In most of the reported studies, trypsin was deposited in discrete positions by a spotting device (Groseclose et al. 2007; Stauber et al. 2008). Only recently, homogeneous application of trypsin by a spraying device has been reported (Djidja et al. 2010). The spatial resolution of studies reported so far was typically in the range of 200–250 μm. The smallest pixel size for MS imaging of tryptic peptides reported earlier was 150 μm (Stauber et al. 2010).

In our study, trypsin was deposited by a spraying device similar to that used for matrix application (Bouschen et al. 2010). All parameters of sample preparation were carefully optimized (Schober et al. 2012a). Trypsin was deposited in several aliquots (of 3 μL each) in intervals of 15 min, preventing excessive accumulation of liquid on the tissue. The sample was placed in an incubator (37 °C, 100 % relative humidity) between spraying cycles and was incubated overnight after the last trypsin deposition. Another important factor is the employed buffer solution, NH₄HCO₃, which proved to be compatible with subsequent matrix application. We also used a trypsin concentration that was twice as high as in most published studies. The resulting tryptic peptides in a mouse brain section were analyzed by mass spectrometry imaging at 50 μm pixel size. Peptides were identified by comparison with LC–ESI–MS/MS as described above [see Schober et al. (2011)]. One of the first measurements using this new approach is shown in Fig. 6. The mass spectrum acquired from a single 50-μm pixel shows two peaks at nominal mass m/z 726 (Fig. 6a).

The myelin peptide HGFLPR (theoretical m/z 726.4046) was detected with a mass deviation of −0.28 ppm and a mass resolution of R = 24,000 in this spectrum. The overall mass accuracy for this peptide in the imaging measurement calculated from more than 500 individual spectra was 1.0 ppm (root mean square of all mass deviations). Another signal at the same nominal mass was detected at m/z = 726.5114 and was discriminated from the myelin peptide signal. This second peak was identified as an isotopologue peak of the ammonium adduct of the lipid phosphatidylserine PS (30:0) (theoretical

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**Fig. 5** a, b Selected ion images of two identified peptides with an imaging bin width of Δm/z = 0.01. c Section of the averaged mass spectrum of high mass accuracy MS imaging of mouse brain tissue, containing two identified tryptic peptides. d m/z values of all tryptic peptides identified in homogenate LC/ESI–MS/MS measurements in the displayed mass range (m/z 1,256.50–1,256.75). Modified from original figure in Schober et al. (2011). Copyright © 2011 John Wiley & Sons, Ltd.
Fig. 6 Mouse brain (coronal section) after on-tissue tryptic digestion. a Mass spectrum from a single 50-μm pixel. b Optical image of adjacent section after staining for myelin (Luxol fast blue). c, d, e MS images, 50 μm pixel size, 92 × 128 pixels: c selected ion image of \( m/z = 726.40–726.41 \) corresponding to myelin peptide. d Selected ion image of \( m/z = 726.40–726.60 \). e Selected ion image of \( m/z = 726.51–726.52 \) corresponding to lipid isotopologue peak. Details on method can be found in Schober et al. (2012a).

\( m/z \) 726.5128). Fig. 6d shows a selected ion image generated with a bin width of \( \Delta m/z = 0.2 \). In this image, no differentiation between the peptide and the lipid isotopologue was possible; the myelin peptide signal was masked by the lipid isotopologue. Figure 6c, e show selected ion images generated with a bin width of \( \Delta m/z = 0.02 \) for the myelin peptide and the lipid isotopologue, respectively. In this representation, both analyte distributions were clearly resolved.

The myelin peptide shows a well-defined distribution with high intensities in the corpus callosum area of the mouse brain which is in accordance with its physiological function (Sidman 2011). Figure 6b shows the optical image of an adjacent mouse brain section after staining of the myelin sheath (Luxol fast blue). The myelin distributions in the MS image and the histological staining show excellent correlation. This confirms that the spatial distribution of myelin was preserved and no substantial migration of tryptic peptides occurred during the digestion process.

A similar measurement with a higher number of identified peptides is discussed in Schober et al. (2012a). In total, 9 peptides corresponding to myelin and 19 peptides corresponding to other proteins were identified and imaged. All myelin peptides showed well-defined distributions which were highly correlated with each other. The number of identified peptides in this work is in the same range as that of studies reported in the literature (Groseclouse et al. 2007; Franck et al. 2009; Shimma et al. 2008), but was obtained at significantly smaller pixel size. Peptides were also identified with a very low mass tolerance of 3 ppm and thus with higher confidence.

These measurements represented the first MS imaging experiment of tryptic peptides with a spatial resolution of 50 μm. The small pixel size shown here cannot be achieved with spotting devices since the size of individual droplets and thus the resulting trypsin spots are too large. Diffusion processes within these trypsin spots on tissue lead to unwanted spreading of peptides. The homogeneous application of trypsin by a spraying device can overcome this problem if spraying conditions such as flow rate, distance and buffer solution are carefully controlled. As for all MS imaging experiments, sample preparation and instrumental parameters have to balance analytical sensitivity and spatial resolution. The experiments in this study were optimized for ultimate spatial resolution. Exact conditions need to be optimized for each tissue type and targeted application.

The obtained results represented a significant improvement over previously reported methods in terms of image quality for tryptic peptides. The improved spatial resolution made smaller histological features accessible. High mass resolving power improved the specificity of the spatial distribution information by reducing interferences with neighboring peaks. Accurate mass allowed for reliable identification of tryptic peptides. Each of these factors was
found to be critical in order to obtain reliable information on biologically relevant compounds. The capabilities of the method were demonstrated in a proof-of-concept study on mouse brain tissue.

Current experiments demonstrate that comparable results in terms of reliable identification can also be obtained from clinical tissue samples. Newest developments also indicate that the number of identified peptides can be significantly increased and that spatial resolution can most likely be further improved.

**Single-cell MALDI imaging**

Cell cultures are a common model system in biological and biomedical research. The influence of pathogenic and/or environmental factors can be studied under controlled conditions. Metabolic studies in cell cultures are usually based on the analysis of bulk samples. The analysis of individual cells would instead provide much more specific and informative data. Previous attempts to use MS imaging for this application faced a number of problems. Time-of-flight secondary ion mass spectrometry (TOF-SIMS) offers sub-micrometer spatial resolution and is able to image the spatial distribution of metabolites inside single cells (Col-liver et al. 1997; Ostrowski et al. 2004). However, TOF-SIMS in single-cell analysis is typically limited in mass range to \( m/z < 500 \), and detected analytes can often not be identified (Szakal et al. 2011). This is due to severe fragmenta
tion of analyte ions and limited MS/MS capabilities in imaging mode. MALDI imaging, in contrast to that, offers higher mass range and higher mass resolution. However, direct analysis of single cells with this method resulted in only one pixel per cell due to the limited spatial resolution of 40–50 \( \mu \)m in reported studies (Miura et al. 2010; Zimmerman et al. 2011).

Based on the instrumentation and methodology described above, we developed a dedicated workflow for imaging metabolite distribution in single cells (Schober et al. 2012b). Sample preparation is a critical step in obtaining useful results in mass spectrometry imaging, especially at high spatial resolution. Fixation methods with ethanol or formaldehyde resulted in the loss of the cells during sample preparation. The best results for analysis with MALDI MS imaging were obtained by fixation of cells with glutaraldehyde. Experimental details such as temperature of solutions, buffer concentration and incubation times proved to be critical and were carefully optimized. After fixation, cells were stained with 3,3′ dihexyloxacarbocyanine iodide [DIOC\(_6\)(3)], which is a fluorescence dye that stains cell membranes (Invitrogen 2011). After matrix application, samples were measured with our AP-SMALDI ion source on an ‘Exactive Orbitrap’ mass spectrometer (Thermo Fisher Scientific GmbH, Bremen, Germany). This system provided improved sensitivity and mass resolution \( (R = 100,000 \times m/z 200) \) compared to the previously used ‘LTQ Orbitrap.’

The images of two individual HeLa cells are shown in Fig. 7. The signal of DIOC\(_6\)(3) in optical fluorescence mode (wavelength \( \lambda = 501 \) nm, Fig. 7a) and MS imaging mode (selected ion image of \( m/z 445.285 \), Fig. 7b) show very good correlation. The shape of the cells is clearly visible in the MS image, and their extensions of only a few micrometers in diameter are detectable. This implies that the spatial structure of the cells was not affected significantly by matrix application and the measurement procedure. Figure 7c, d show selected ion images of two phospholipids. The (sodium adducts of) phosphatidylcholines PC (16:0/16:1) and PC (16:0/18:1) were detected primarily in the center of the cell where higher concentrations of membranes are expected. The fatty acid chain length of these two phospholipids was determined by additional MS/MS experiments [see Schober et al. (2012b) for details].

A mass spectrum acquired from a single 7-\( \mu \)m pixel is shown Fig. 7e. Numerous phospholipids such as phosphatidylcholine (PC), sphingomyelin (SM), phosphatidylserine (PS), as well as di- (DG) and triglycerides (TG) were found in the mass range \( m/z \) 620–810. As in all our studies, the imaged compounds were detected with accurate mass (better than 3 ppm RMS for the complete image) and therefore allow reliable substance identification. In addition to phospholipids, smaller metabolites such as nucleic acids, amino acids as well as cholesterol were identified in the lower mass range [see Schober et al. (2012b)]. A broad metabolic profile could thus be acquired from a defined area of the cell with high confidence. Alternatively, several spectra acquired from one cell can be combined in order to obtain better statistics for quantitative analysis.

Current experiments focus on the mass range of phospholipids. Invasion mechanisms of pathogens are known to lead to changes in the lipid pattern through enzymatic degradation (Vazquez-Boland et al. 2001). These effects can be investigated with high molecular specificity in single cells at different stages of the infection process with the methodology described here. Another application that is currently evaluated is monitoring the differentiation of stem cells on a molecular level. In general, our method can be used for a detailed investigation of metabolic changes in single cells and can provide more detailed and/or more reliable information than previous MS imaging approaches.

**Analysis of lung carcinoma tissue**

We typically test our methodology with mouse brain tissue as this is a widely used model system for MS imaging. The
molecular composition of this tissue is well known, and extensive histological information is available in brain atlas resources (Sidman 2011; Hochheiser and Yanowitz 2007). Once a certain method is optimized, it is applied to more relevant samples including clinical tissue sections. An example that was first shown in a conference presentation (Römpf et al. 2011a) is the analysis of a human non-small-cell lung carcinoma that was induced into a severe combined immunodeficiency (SCID) mouse model (Fig. 8). This tissue section was imaged at 10 \( \mu \text{m} \) pixel size. MS image analysis (Fig. 8a) shows excellent correlation with the different regions within the cancerous tissue that were identified by the pathologist in the H&E staining (Fig. 8b). Cerebrosides Cer (42:2) (Fig. 8a, green) was found to be almost exclusively present in the connective tissue of this section. The tumor region can be further differentiated in the invading tumor edge with high signal intensities of phosphatidylcholine PC (32:1) (Fig. 8a, blue) and the ischemic tumor region that was characterized by high signal intensities of phosphatidylcholine PC (40:6) (Fig. 8a, red). Purple color results from colocalization of PC (32:1) (blue) and PC (40:6) (red). Multiple other lipid species were detected and identified for each tissue region in addition to the examples shown here. Figure 8c shows the distribution of lyso-phosphatidylcholine LPC (16:0) (green), primarily localized in the area of the tumor that was identified as necrotic zone. Lyso-lipids are degradation products of lipids and are therefore expected in areas where cell degradation occurs. This is another example where our method provides more insights about the molecular information that is not directly accessible by classical histological methods. This approach can be applied to investigate the molecular processes in tumorigenesis and to differentiate tumor states.

Lipids in this measurement were identified by accurate mass data. As demonstrated above for tryptic peptides (Fig. 6), the bin width used for image generation significantly affects the result of an MS imaging experiment. Generating MS images with a bin width of \( m/z = 0.1 \), as common for many MALDI–TOF measurements, would result in the distribution shown in Fig. 8d. However, the healthy and tumor tissue regions can only be differentiated by using a bin width of \( \Delta m/z = 0.01 \) (Fig. 8e, f). These black-and-white images represent the raw data acquired from the mass spectrometer without applying any processing steps. Overlay of these two signals shows very sharp separation between (healthy) connective tissue and the tumor region (Fig. 8g). Similar effects were observed for nearly all signals in the lipid mass region. Consequently, the reliable characterization of complex samples such as mammalian tissue sections by mass spectrometry imaging greatly benefits from the use of high-resolution mass spectrometry.

MALDI imaging at 3 \( \mu \text{m} \) pixel size

In an attempt to further improve the accessible spatial resolution of MALDI imaging, the lateral ventricle region of a coronal mouse brain section was imaged with 3 \( \mu \text{m} \) pixel size. The MS image of this experiment, which was included in a conference presentation (Römpf et al. 2010b), is shown in Fig. 9a. The selected ion image of phosphatidylcholine PC (38:4) shows the detailed structure...
of the choroid plexus (green), while the surrounding tissue is characterized by high intensities of phosphatidylcholine PC (34:1). The optical image of the tissue sample prior to MS analysis is shown in Fig. 9b. The poor quality of the optical image is due to the fact that the sample is completely consumed under these highly focused laser conditions, and subsequent staining of tissue was therefore not possible. The quality of the mass spectral data is, however, not significantly affected by the reduction in pixel size. A mass spectrum acquired from a single pixel (corresponding to $3 \times 3 \mu m^2$ tissue area) is shown in Fig. 9c. Phospholipids in this mass spectrum were detected with a mass deviation of $<2$ ppm and a mass resolution of about $R = 20,000$. Lower signal intensities make recalibration based on matrix cluster signals slightly more challenging in this experiment. These measurements demonstrate that under slight oversampling conditions (at an ablation spot diameter of $5 \mu m$ in diameter) still high-quality mass spectral data can be obtained in high-resolution AP-SMALDI imaging and that there is still potential for a further increase in spatial resolution.

Spatial resolution and measurement speed

It is often argued that imaging at high spatial resolution takes too much time. This might be true if two experiments of the same measurement area are compared. But high spatial resolution imaging is typically targeted at smaller structural features which would not be resolved otherwise. Therefore, the spatial resolution (pixel size) of an experiment should be adjusted according to the structure of interest. The speed of our method of about 1 s per pixel is comparable to that of other mass spectrometry imaging systems. Recent experiments indicate that significant increase in speed can be obtained (with almost constant mass accuracy and moderately lower mass resolution). At present, our method already provides reliable MS imaging measurements at different spatial resolution settings as demonstrated in Fig. 10. All three MS images show detailed histological features (at different spatial scales). Figure 10a shows the phospholipid spatial distribution in mouse brain (coronal section) at $5 \mu m$ pixel size. The structure of the putamen and the corpus callosum (upper right) are clearly visible. Figure 10b shows a horizontal
section of mouse brain at 50 μm pixel size. The two imaged phospholipids show detailed features in the corpus callosum, striatum and cerebellum region. Figure 10c shows a measurement of the intestinal tract of a whole body rat section at 200 μm pixel size. A phospholipid (red) that provides histological information can be directly used to indicate the localization of a drug compound (green) [see Römpp et al. (2011b) for details]. A useful approach for balancing measurement time and spatial information is to scan a larger area with lower spatial resolution in order to identify areas of interest. These areas can then be scanned with higher spatial resolution in order to investigate the detailed distribution of analytes.

Measurement speed is obviously an important factor when comparing quantitative differences between two biological conditions. In this case, biologically relevant results can only be obtained if the experiment is repeated with several samples (biological replicates). Consequently, higher speed allows for a higher number of measurements and thus provides more reliable results. Another approach is the detailed molecular analysis of a (tissue) sample without prior knowledge. In such an untargeted approach, data analysis and interpretation can be much more time-consuming than the actual measurement itself. In this case, specificity and sensitivity of the measurement is of highest priority and measurement speed is of less importance.

**Data analysis: imzML—a common data format for MS imaging**

Mass spectrometry imaging produces large and complex data sets, and thus efficient strategies for processing and
analyzing are essential. A common data format for the flexible exchange and processing of mass spectrometry imaging data was developed within the EU joint project COMPUTIS (www.computis.org). The data format is in part based on the HUPO-PSI format mzML (Martens et al. 2011) and was thus called imzML for ‘imaging mzML.’ The concept of imzML and application examples are summarized in Schramm et al. (2012). Additional information and all technical details necessary to implement imzML are provided in a book chapter (Römpp et al. 2010c) and on the Web site www.imzml.org.

MS imaging data in imzML format are divided into two separate files. Metadata (e.g. instrumental parameters and sample details) are stored in an XML file. Information in this file is organized in a controlled vocabulary in order to avoid ambiguities (Smith et al. 2007). Mass spectra are stored in a separate binary file in order to ensure efficient data storage. The XML structure in combination with the controlled vocabulary ensures that the data format can be easily adapted to new instrumentation and experimental strategies.

Data sets from more than 10 different MS imaging platforms have been converted into imzML so far. Examples shown in Fig. 11 include data from matrix-assisted laser desorption/ionization (MALDI) as well as desorption electrospray ionization (DESI) and secondary ion mass spectrometry (SIMS). A growing number of processing options are available for the imzML format. The lower part of Fig. 11 shows software tools which were developed by COMPUTIS partners. imzML can also be employed for processing data with different tools sequentially as demonstrated in Schramm et al. (2012). Additional examples of imzML conversion and processing options are displayed in the ‘imzML gallery’ on www.imzml.org. imzML-based software was also used to process data from our high-resolution mass spectrometry imaging method. While the small pixel size is not a problem, some software tools need to be adapted to handle high mass resolution data (i.e. to generate MS images with a narrow bin width of $\Delta m/z = 0.01$).

ImzML can also be used to compare data from different instrument platforms. Options for data processing (e.g. binning, normalization) and visualization (e.g. interpolation options, color schemes) vary strongly between different MS imaging software tools and can make a direct comparison of data difficult. This problem can be avoided if all data sets are converted to imzML and the data are displayed with identical parameters of one software tool [see Schramm et al. (2012) for details].

The concept of imzML is also actively supported by major vendors of MS imaging instrumentation including Thermo Fisher Scientific (Bremen, Germany), Waters

![Fig. 11 Combination of data from different MS imaging platforms and software tools on the basis of the common data format imzML. See Schramm et al. (2012) for details](image-url)
Corporation (Manchester, UK) and Bruker Daltonik (Bremen, Germany). An updated list of available tools and workflows can be found on the imzML Web site. This also includes the first commercial software tools, which are compatible with imzML, from Imabiotech (2012) and PREMIERBiosoft (2012).

imzML provides more flexibility in data processing as the user is no more limited to proprietary software but can choose the processing option that best fits the purpose. Data can also easily be shared between collaborators as several of the software tools are freely available. Measurement results can be provided to partners in biological or clinical laboratories without the loss of information (which is the case when sharing only the images). Finally, a common data format provides more transparency in reporting MS imaging results. Consequently, imzML can facilitate everyday work with MS imaging data.

Conclusion and perspective

The results presented above demonstrate that mass spectrometry imaging with high molecular specificity is possible at or close to cellular dimensions. Lipids, (neuro)peptides and drug compounds were imaged in a wide range of biological samples at a spatial resolution of 5–10 μm. A measurement of mouse urinary bladder at 5 μm step size showed pixel-sharp separation between tissue regions. A drug compound (imatinib) was imaged in mouse kidney at 10 μm pixel size. Correlation with spatial distributions of (simultaneously detected) phospholipids allowed to directly link areas of imatinib accumulation to histological structures. More than 10 neuropeptides were imaged at 5 and 10 μm pixel size in mouse pituitary gland. Peptide identification by accurate mass was confirmed by additional MS/MS imaging experiments.

On-tissue digestion was employed for the analysis of proteins. Tryptic peptides were imaged for the first time with accurate mass and identified by complementary LC–ESI–MS/MS measurements of an adjacent tissue section. This approach significantly increased both the reliability and the number of peptide identifications. In subsequent experiments, the spatial resolution for tryptic peptides was improved to 50 μm while maintaining the high molecular specificity. This was achieved by applying trypsin with a spraying device.

A dedicated sample preparation protocol was established for the analysis of cell cultures. Phospholipids and smaller metabolites such as amino acids and nucleic acids were imaged in single cells. A full metabolic profile was obtained from a single pixel of 7 × 7 μm².

The feasibility of obtaining high-quality mass spectrometry imaging data from clinically relevant tissue was demonstrated by the analysis of lung carcinoma tissue. The highest spatial resolution of a MALDI imaging experiment of tissue so far was obtained by analyzing a mouse brain section at 3 μm pixel size.

MS image analysis for all these experiments showed excellent correlation with histological evaluation. In addition, it provided highly specific molecular information. In many cases, components of very similar mass (Δm/z < 0.1) showed distinctly different spatial distributions. This demonstrated the need for accurate mass determination in order to obtain reliable information from MS imaging experiments of biological samples. The ability to combine high mass resolution and accuracy with a high spatial resolution in the low micrometer range is a unique feature of the presented method.

Each of these studies represented a significant improvement in spatial resolution and/or reliability of identification compared to studies reported earlier. These methodological improvements were made possible by the development of an integrated workflow that is optimized for ‘mass spectrometry imaging with high resolution in mass and space.’ The experiments presented above were not merely the application of one method to different samples, but involved new experimental approaches for each study. For example, sample preparation was carefully adjusted for each application. This was especially important for the analysis of single cells and the detection of tryptic peptides, but also for the analysis of highly water-soluble analytes such as neuropeptides and drug compounds.

While small pixel size is not necessarily a prerequisite for high-quality MS imaging, higher mass accuracy and mass resolution will always result in more reliable data. The high complexity of biological samples inevitably leads to multiple peaks for each nominal mass. High mass accuracy significantly improves the reliability of identification as demonstrated in the examples above. This is even more important when different MS methods are combined as shown for the example of tryptic peptides. Therefore, MS imaging experiments should always be designed as to deliver the highest mass accuracy and mass resolution possible (for a given instrumental setup). More frequent mass calibration steps, internal standards and/or postrun recalibration can be used to achieve this goal.

Up to now, we have applied our mass spectrometry imaging method to a wide variety of applications. Targeted compound classes included lipids, amino acids, peptides, proteins, nucleic acids, carboxylic acids and carbohydrates. In addition to the examples discussed above, we have analyzed several sample types including muscle, testis, spinal disk as well as adrenal gland tissue originating from mouse, rat, pig or cow. In these experiments ‘mass spectrometry imaging with high resolution in mass and space’ proved to be a versatile and robust method for the highly detailed analysis of biological samples.
A major focus of our current and future studies is the analysis of clinical tissue samples. Several types of human tumor biopsies are currently being analyzed. Sets of carefully selected samples will be analyzed in close collaboration with clinical partners. On the methodological side, the current activities are aimed at further improving measurement speed, sensitivity and spatial resolution. In addition to optimizing existing protocols, new application areas are being explored. This includes the analysis of material from plants, insects and fungi.

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