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Analyzing cell-type-specific dynamics of metabolism in kidney repair

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A common drawback of metabolic analyses of complex biological samples is the inability to consider cell-to-cell heterogeneity in the context of an organ or tissue. To overcome this limitation, we present an advanced high-spatial-resolution metabolomics approach using matrix-assisted laser desorption/ionization mass spectrometry imaging (MALDI-MSI) combined with isotope tracing. This method allows mapping of cell-type-specific dynamic changes in central carbon metabolism in the context of a complex heterogeneous tissue architecture, such as the kidney. Combined with multiplexed immunofluorescence staining, this method can detect metabolic changes and nutrient partitioning in targeted cell types, as demonstrated in a bilateral renal ischemia-reperfusion injury (bIRI) experimental model. Our approach enables us to identify region-specific metabolic perturbations associated with the lesion and throughout recovery, including unexpected metabolic anomalies in cells with an apparently normal phenotype in the recovery phase. These findings may be relevant to an understanding of the homeostatic capacity of the kidney microenvironment. In sum, this method allows us to achieve resolution at the single-cell level in situ and hence to interpret cell-type-specific metabolic dynamics in the context of structure and metabolism of neighboring cells.

Spatial omics methods are becoming an increasingly important means to provide insights into in situ tissue cellular heterogeneity and how cells interact and behave within their microenvironments. The aim of analyzing the entirety of cellular metabolites at single-cell resolution is challenging owing to enormous structural diversity, rapid turnover, and low analyte abundances in a limited sample volume. Previously, MALDI-MSI-based metabolomics methods have been proposed as a way to measure spatial distribution of metabolites and lipids at single-cell resolution^{1,2}. To date, MALDI-MSI-based single-cell metabolomics studies have, however, focused on analyzing single-timepoint metabolic ‘snapshots,’ which provide valuable information on in situ metabolic heterogeneity but crucially lack insight into the dynamic component of cellular metabolism^{3–5}; hence, there is a need for approaches that provide a truly comprehensive understanding of the interplay between biochemical

alterations and cell-type-specific functions, metabolic fluxes, and dynamic interpretations. To this end, isotope tracing has been used to model dynamic metabolic processes using bulk metabolomics⁶, in which complete tissues are homogenized and reduced to a single metabolic profile, disregarding tissue and metabolic heterogeneity. The combination of isotope tracing and subsequent analysis by MALDI-MSI could be considered as an approach to assess metabolic dynamics in situ^{7–9}. Here, we describe a platform based on high-spatial-resolution MALDI-MSI (5 × 5 μm² pixel size) that uses isotope tracing to allow for in situ cell-type-specific dynamic metabolism measurements to uncover cell metabolism within the architecture of the tissue that the cells reside in. The average diameter of most cells in kidney tissue is around 10 μm, and the proposed method provides a single measurement at subcellular-level resolution because each cell has a size of approximately 4 pixels in MALDI-MSI images. To gain the most information from the measured area, all pixels, not just those segmented into individual cells, will be used for the cell-type-specific metabolomics analysis that we describe below.

Our method uses ¹³C-labeled nutrients that allow tracing of the spatiotemporal incorporation of the ¹³C isotopes into the main intermediates of glycolysis and the tricarboxylic acid (TCA) cycle (Fig. 1a). Using different labeled nutrients allows not only visualization of dynamic changes over time, but also resolution of their contributions to specific metabolic pathways. To accomplish this, we used a tissue culture system in which 350-μm-thick vibratome slices of fresh mouse kidney¹⁰ (Fig. 1b) were incubated for up to 2 hours. During incubation, ¹³C-isotope-labeled nutrients were introduced in parallel to the tissue culture medium at selected timepoints, which led to efficient and biochemically meaningful labeling of metabolically active cells¹¹. We could then measure metabolic changes in a non-steady state. Subsequently, tissue slices were quenched with liquid nitrogen and sectioned into 10-μm-thick sections, thaw-mounted on conductive glass slides, and coated with a chemical matrix for MALDI-MSI measurements. Negative-ion-mode MALDI-MSI at subcellular resolution (that is, pixel size of 5 × 5 μm²) was used to detect metabolites and (phospho)lipids¹² (Extended Data Table 1) from the collected tissues.

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Following MALDI-MSI measurements, the sections were stained and subsequently imaged using multiplexed immunofluorescence (IF) microscopy for cell-type identification (Fig. 1c). By combining IF and spatial segmentation of the MALDI-MSI data (on the basis of uniform manifold approximation and projection (UMAP) of the lipid signals, in particular), cell-type-specific signatures were established using the following assumptions: (1) MALDI-MSI lipid profiles are cell-type specific and (2) major phospholipid species, important for cell typing, are mainly cell-membrane components¹³ and thus are stable during the 2 hours of isotopic labeling (Extended Data Fig. 1). Providing functional evidence for this approach, MALDI-MSI and molecular histology of tissues incubated with [$U\text{-}^{13}\text{C}$]linoleate, [$U\text{-}^{13}\text{C}$]glutamine, or [$U\text{-}^{13}\text{C}$]glucose revealed similar distributions of lipid signatures for cell typing (Extended Data Fig. 1f). MALDI-MSI measurements were performed on sections at different timepoints and with different ^{13}C -enriched nutrients to obtain the labeling datasets for each condition. Thus, in order to show all the dynamic metabolic changes within 1 pixel without batch effects, a two-step process was followed, as introduced by Stuart et al. in the Seurat package¹⁴: (1) single-pixel lipid profiles were used to identify anchors between two datasets; (2) ^{13}C -labeled-metabolite production was imputed into the control dataset by transferring the abundance of all measured ^{13}C -enriched metabolites from the labeling datasets using k -nearest neighbors (KNN) analysis (Fig. 1d). An imputed dataset containing the complete ^{13}C -labeling information, representing the predicted values of mass isotopomers from the pixels with a similar lipid profile for each tracing experiment, from different timepoints and nutrients could thus be established. Dynamic metabolic calculations, including metabolic changes and pathway convergence, and assessment of ^{13}C enrichment of isotopologues were performed on single-pixel data. Finally, the in situ heterogeneity of metabolic dynamics was visualized in pseudoimages generated from the calculated values (Fig. 1e).

This approach allowed us to detect both endogenous and ^{13}C -labeled metabolites from glycolysis and the TCA cycle, as well as branching metabolites, including hexose, glycerol-3-phosphate (G3P), 3-phosphoglycerate (3PG), ribose-5-phosphate/xylulose-5-phosphate (R5P/X5P), lactate, glutamate, glutamine, malate, aspartate, and linoleate (Extended Data Fig. 2). To validate metabolite imputation performance, a 'leave-one-factor-out' cross-validation¹⁵ was applied to a bisected MALDI-MSI dataset from a single kidney section after 2 hours of incubation with [$U\text{-}^{13}\text{C}_6$]glucose (Extended Data Fig. 3). The 'new' datasets (samples 1 and 2) created from the single MALDI-MSI measurement underwent identical sample handling and thus had comparable technical variation and metabolic activity (Extended Data Fig. 3a). The correlation between the imputed and measured metabolites at the single-pixel level was moderate (mean $r=0.49$, $P<0.001$, Extended Data Fig. 3b). To test imputation performance at the cell-type level, we divided all pixels into 29 clusters on the basis of their lipid profiles

(Extended Data Fig. 3c). Following clustering, cell-type validation was performed using Spearman's correlation analysis on the average value of each cluster. A very strong correlation (mean $r=0.94$, $P<0.05$) was observed between the imputed and detected values (Extended Data Fig. 3d). In addition, when comparing imputed and detected metabolite abundance, the average ^{13}C enrichment of isotopologues in relation to endogenous metabolites in each cluster was highly accurate (ratio= 0.99 ± 0.04) (Extended Data Fig. 3e–g). We continued to test the MALDI-MSI data from two kidneys (samples 3 and 4) and found a strong correlation between imputed and detected metabolite intensity at the cluster level (mean $r=0.82$, $P<0.001$, Extended Data Fig. 3h–j). Following validation of the dynamic metabolism analysis, which revealed that the imputed cell-type-specific ^{13}C -enrichment calculations were very reliable, pseudoimages were generated to show the spatial changes in dynamic metabolism within one tissue (Extended Data Fig. 4).

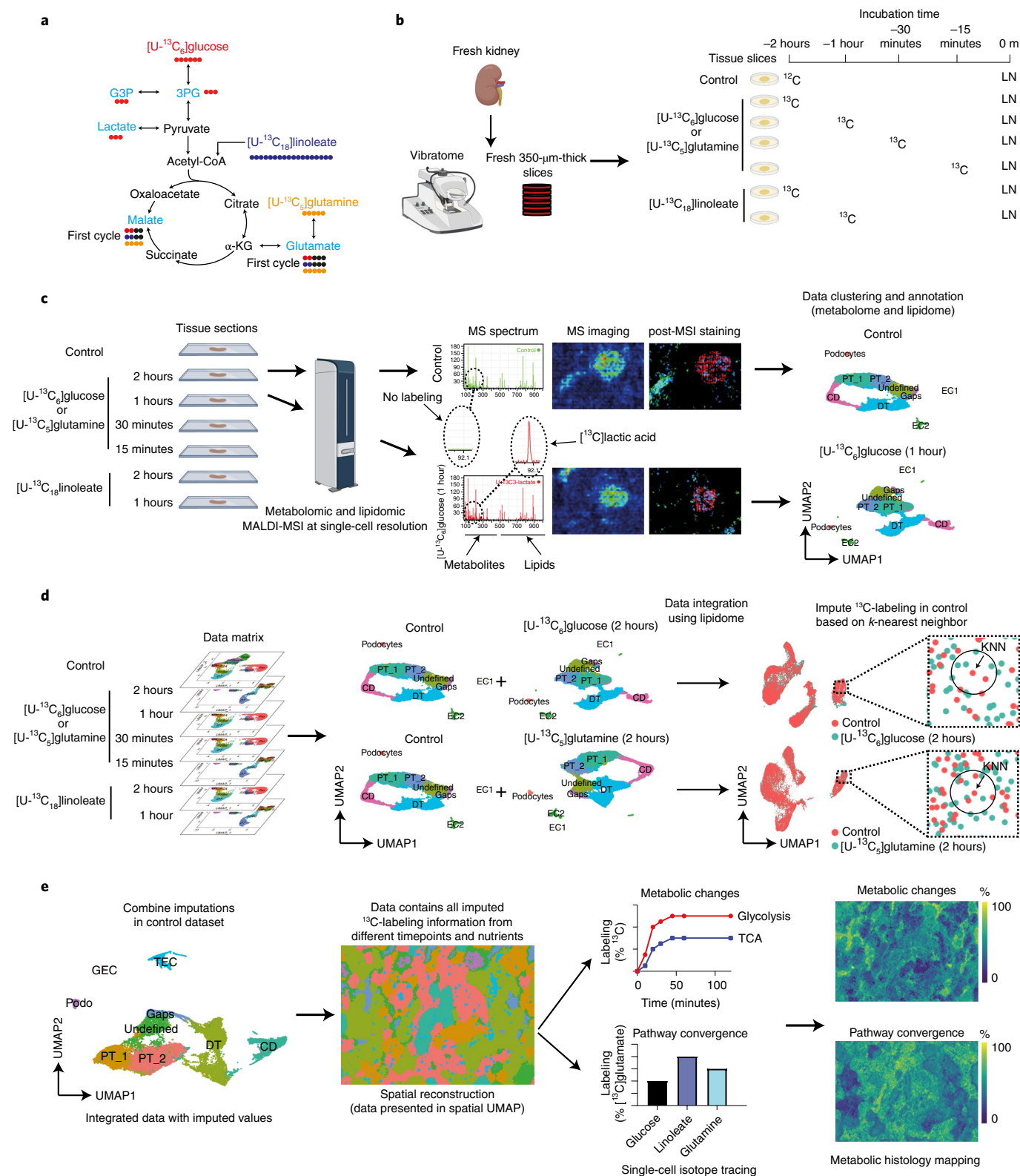
To demonstrate how the method can be used, we induced moderate ischemic injury to mouse kidneys, similar to that observed in a kidney-transplantation setting. We then studied the tissue metabolism 2 weeks after the injury. At this timepoint, part of the initial proximal tubular epithelial injury has regenerated (Extended Data Fig. 5a) and has taken up a normal phenotype again (characterized by *Lotus tetragonolobus* lectin (LTL) positivity and kidney injury molecule-1 (KIM1) negativity), while other epithelial cells may still have a maladaptive inflammatory phenotype, as exemplified by vascular cell adhesion molecular 1 (VCAM1) expression in the presence of LTL, or may have lost the tubular epithelial phenotype all together (LTL negative)¹⁶. To visualize overall metabolic heterogeneity in whole kidney tissue sections, we used MALDI-MSI at a $20\times 20\mu\text{m}^2$ pixel resolution, which spatially and molecularly resolved all major renal structures on the basis of their specific lipid signatures (Fig. 2a and Extended Data Fig. 1). Two weeks after kidney bIRI, we found additional clusters (Injured_1, Injured_2, Injured_3, and Injured_4) with unique lipid signatures that corresponded with loss of LTL and E-cadherin staining (Fig. 2a–c and Extended Data Fig. 5b–d). A loss of the healthy proximal tubule (PT) lipid signature and an increased presence of oxidized lipid species (m/z 865.5; Extended Data Table 1) were observed in the areas with persisting epithelial injury (Fig. 2c). We then generated spatial segmentation images using an integrated 3D UMAP analysis, which displayed the kidney molecular histology. These images showed the presence of ongoing tubular injury in both the cortex and outer medulla of bIRI kidneys (Fig. 2d). There was substantial loss of PT structures from the outer stripe of outer medulla (PT-S3) compared with the renal cortex (PT-S1/PT-S2) (Fig. 2e). These observations were also confirmed using a histopathological staining (Fig. 2f).

Considering that a tissue's microenvironment is an important cue for cell-specific metabolism within that tissue, we next used high-spatial-resolution MALDI-MSI measurements with a pixel

Fig. 1 | Workflow of cell-type-specific dynamic metabolic measurements and analysis. **a**, Overview of the traced isotopes in the primary carbon metabolism. The contributions of $U\text{-}^{13}\text{C}$ -labeled nutrients ([$U\text{-}^{13}\text{C}_6$]glucose, [$U\text{-}^{13}\text{C}_{18}$]linoleate, and [$U\text{-}^{13}\text{C}_5$]glutamine) to glycolytic and TCA intermediates (light blue) were traced. **b**, Fresh mouse kidney tissue was cut into $350\text{-}\mu\text{m}$ -thick slices using a vibratome. For ^{13}C isotope tracing in tissue culture, different ^{13}C -labeled nutrients were added to a well-defined medium described in the Methods section at different timepoints. Samples were quenched using liquid nitrogen (LN). **c**, The metabolome and lipidome were measured in all samples using MALDI-MSI at high spatial resolution ($5\times 5\mu\text{m}^2$ pixel size). MALDI-MSI data were preprocessed and transferred into a data matrix. Cell types were identified on the basis of lipid profiles and IF staining after MALDI-MSI. Images in **b** and **c** were created using Biorender. **d**, Cell-type-specific (phospho)lipid data were used to characterize various cell types. The lipid data were used for anchor-based data integration of the 'control' data matrix with data matrices of sections from ^{13}C -isotope tracing measurements. We used KNN analysis to impute the molecular information contained in the ^{13}C -labeling timecourse data matrices into the control data matrix. **e**, Establishment of the imputed dataset, in which each pixel contains all added ^{13}C -labeling information from each timepoint and labeled nutrient. Dynamic metabolic calculations were performed on single pixels, including metabolic rates and pathway convergence. To visualize the heterogeneity in tissue metabolic dynamics, a series of pseudoimages, which were generated from calculated values, were created by tracing pixel coordinates back to the original spatial information from the MALDI-MSI analysis. $\alpha\text{-KG}$, alpha-ketoglutaric acid; DT, distal tubules; CD, collecting ducts; PT, proximal tubules; EC, endothelial cells; GEC, glomerular endothelial cells; TEC, peritubular endothelial cells.

size at the subcellular level ($5 \times 5 \mu\text{m}^2$) to investigate the metabolic heterogeneity in the PT of the kidney cortex and outer stripe of outer medulla and their response to injury. Following MALDI-MSI, IF staining was conducted to identify cell types (Fig. 3a). We could distinguish two PT-S1 and PT-S2 cortex segments and one medullary PT-S3 segment on the basis of their lipid profiles (Fig. 3a–c). PT-S1 and PT-S2 were combined for subsequent analyses because

we could not annotate them on the basis of LTL staining or their location. Analysis of the isotopologues showed that PT-S3 had lower hexose M+6 ($[\text{U-}^{13}\text{C}_6]$ hexose) and higher lactate M+3 ($[\text{U-}^{13}\text{C}_3]$ lactate) enrichment than did PT-S1/PT-S2 (Fig. 3d). In addition, we found lower enrichment of R5P/X5P M+3 (glycolysis branch) as well as lower enrichment of glutamate ($[\text{U-}^{13}\text{C}_2]$ glutamate; Glu M+2) from glucose carbons entering the TCA cycle in PT-S3 (Fig. 3d). To



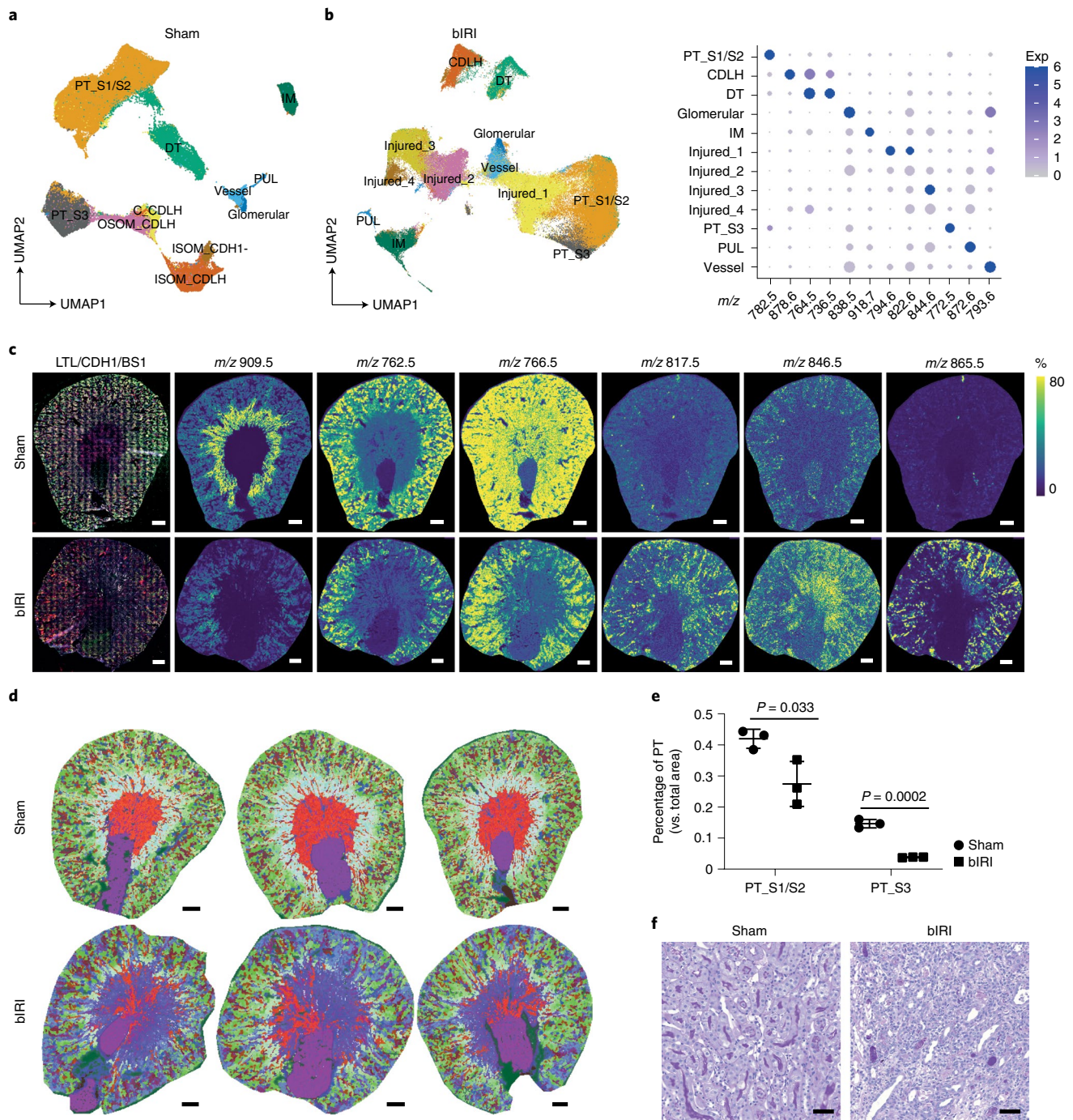
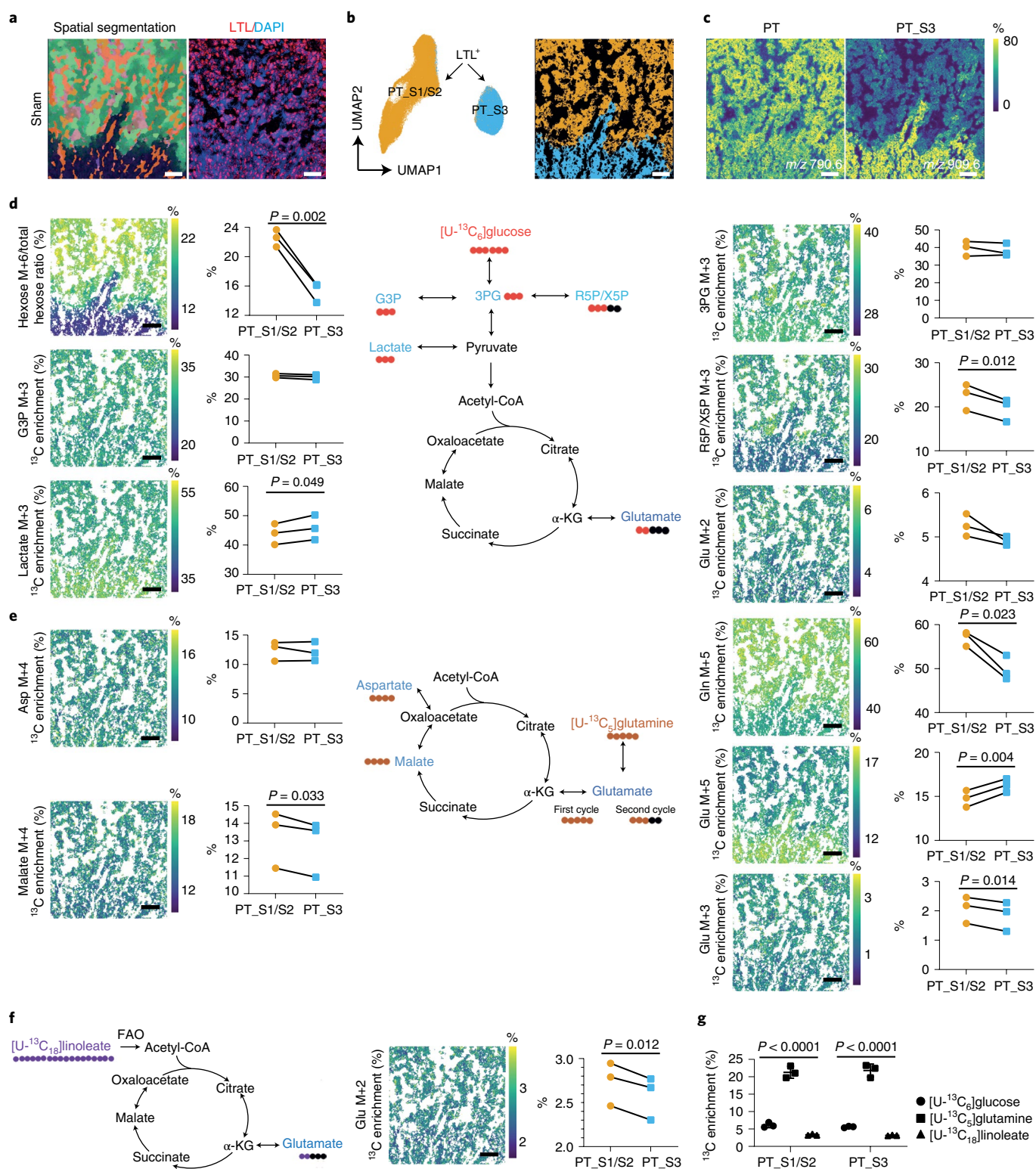


Fig. 2 | Lipid heterogeneity in mouse IRI kidneys. **a**, Lipid heterogeneity in mouse sham kidneys ($n=3$) from which mice were opened up and closed similarly as bIRI mice but no ligations were performed, visualized in a two-dimensional (2D) UMAP plot of MALDI-MSI data ($20 \times 20 \mu\text{m}^2$ pixel size). **b**, Lipid heterogeneity in mouse bIRI kidneys ($n=3$), visualized in a 2D UMAP plot of MALDI-MSI data at $20 \times 20 \mu\text{m}^2$ pixel size. The dot plot displays lipid expression of cluster-enriched signatures. Exp, expression. **c**, IF staining (LTL, green), E-cadherin (CDH1, red), and BSI-lectin (gray) of tissue that had been analyzed with MALDI-MSI. Representative images showing lipid species distributions in sham ($n=3$) and bIRI ($n=3$) kidneys, as recorded by MALDI-MSI ($20 \times 20 \mu\text{m}^2$ pixel size). Scale bars, $500 \mu\text{m}$. **d**, Molecular histology of sham ($n=3$) and bIRI ($n=3$) kidneys generated from integrated three-dimensional (3D) UMAP analysis on the basis of lipid profiles. The color code represents the position of pixels in the 3D UMAP (UMAP1: red, UMAP2: green, UMAP3: blue). Scale bar, $500 \mu\text{m}$. **e**, Comparison of PT areas between sham ($n=3$) and bIRI ($n=3$) kidneys. Two-tailed t -test was performed. **f**, PAS staining showing the tubular structures in the outer stripe outer medulla area of sham ($n=3$) and bIRI ($n=3$) kidneys. Scale bars, $50 \mu\text{m}$. PT-S1/S2, cortical proximal tubular segments 1 and 2; PT-S3, outer stripe of outer medulla proximal tubular segment; DT, distal tubule; CDLH, collecting duct and loop of Henle; IM, inner medulla; OSOM, outer stripe of outer medulla; ISOM, inner stripe of outer medulla; PUL, pelvic urothelial lining.

further evaluate the use of ^{13}C -labeled nutrients in the TCA cycle, we measured the percentage enrichment of $[\text{U-}^{13}\text{C}_5]$ glutamine-derived isotopologues, $[\text{U-}^{13}\text{C}_5]$ glutamate (Glu M+5), $^{13}\text{C}_4$ malate

(malate M+4), and $^{13}\text{C}_3$ glutamate (Glu M+3), all of which are intermediates of one oxidative TCA cycle. There was lower $[\text{U-}^{13}\text{C}_5]$ glutamine and higher $[\text{U-}^{13}\text{C}_5]$ glutamate enrichment in PT-S3



cells than in PT-S1/PT-S2 cells (Fig. 3e). Notably, the enrichment of $[^{13}\text{C}_4]$ malate and $[^{13}\text{C}_3]$ glutamate were relatively lower in PT-S3 than in PT-S1/PT-S2 from the same sample, pointing to a lower contribution of both $[\text{U}-^{13}\text{C}_5]$ glutamine and $[^{13}\text{C}_5]$ glutamate to the oxidative TCA cycle in PT-S3 (Fig. 3e). The contribution of $[\text{U}-^{13}\text{C}_{18}]$ linoleate to the $[^{13}\text{C}_5]$ glutamate isotopologue through fatty acid oxidation was lower in PT-S3 as well (Fig. 3f). By combining these data, we found that $[\text{U}-^{13}\text{C}_5]$ glutamine, rather than $[\text{U}-^{13}\text{C}_6]$ glucose or $[\text{U}-^{13}\text{C}_{18}]$ linoleate, is glutamate's main carbon source in

all PT areas (Fig. 3g). However, usage of glutamate for the oxidative TCA cycle differed between PT segments, as shown by glutamine carbon tracing to characteristic mass isotopomers (Fig. 3e). Together, these data show that, in the normal kidney, there is a marked heterogeneity among PT segments with respect to TCA metabolite consumption and glycolysis: the PT-S3 segment appears to be adapted to the low-oxygen-tension environment of the outer medulla. These observations correspond to previous studies using isolated or micro-dissected PTs^{17,18}.

Fig. 3 | Dynamic metabolic measurements on sham kidney PT cells. **a**, Left, representative molecular histology of cortical and outer stripe of outer medulla areas of sham kidney ($n=3$), generated from three-dimensional UMAP analysis on the basis of lipid profiles recorded by MALDI-MSI ($5 \times 5 \mu\text{m}^2$ pixel size). Right, LTL immunofluorescent staining on post-MALDI-MSI tissue (red). **b**, 2D UMAP plot and representative spatial segmentation showing lipid heterogeneity between LTL-positive proximal tubular cells from the cortical and outer stripe of outer medulla areas of sham kidneys ($n=3$). **c**, Representative images showing lipid species distribution in the cortical and outer stripe of outer medulla areas of sham kidney ($n=3$), as recorded by MALDI-MSI ($5 \times 5 \mu\text{m}^2$ pixel size). **d**, Dynamic metabolic measurements using $[\text{U-}^{13}\text{C}_6]\text{glucose}$ on PT cells of sham kidneys. **e**, Dynamic metabolic measurements using $[\text{U-}^{13}\text{C}_5]\text{glutamine}$ on PT cells of sham kidneys. **f**, Dynamic metabolic measurements using $[\text{U-}^{13}\text{C}_{18}]\text{linoleate}$ on PT cells of sham kidneys. Images showing the average ^{13}C enrichment of isotopologues on tissue. Graphs showing the comparison of the average ^{13}C enrichment of isotopologues between PT S1/S2 and PT S3. The average ^{13}C enrichment (area under curve (AUC) normalized to total time) of isotopologues were derived from $[\text{U-}^{13}\text{C}_6]\text{glucose}$ or $[\text{U-}^{13}\text{C}_5]\text{glutamine}$ measured at different timepoints (0, 15, 30, 60, and 120 minutes) or $[\text{U-}^{13}\text{C}_{18}]\text{linoleate}$ measured at different timepoints (0, 60, and 120 minutes). Charts showing the traced isotopes and their derived isotopologues. Two-tailed paired *t*-test was performed. All scale bars, $200 \mu\text{m}$. **g**, Direct carbon contribution of different nutrients to glutamate at the 2-hour timepoint, as measured from the glutamate isotopologues M+2, M+3, and M+5. One-way analysis of variance (ANOVA) was performed ($n=3$). Glc, glucose; 3PG, 3-phosphoglycerate; G3P, glycerol-3-phosphate; R5P/X5P, ribulose-5-phosphate/xylulose-5-phosphate; $\alpha\text{-KG}$, alpha-ketoglutaric acid; Glu, glutamate; Gln, Glutamine; Asp, aspartate.

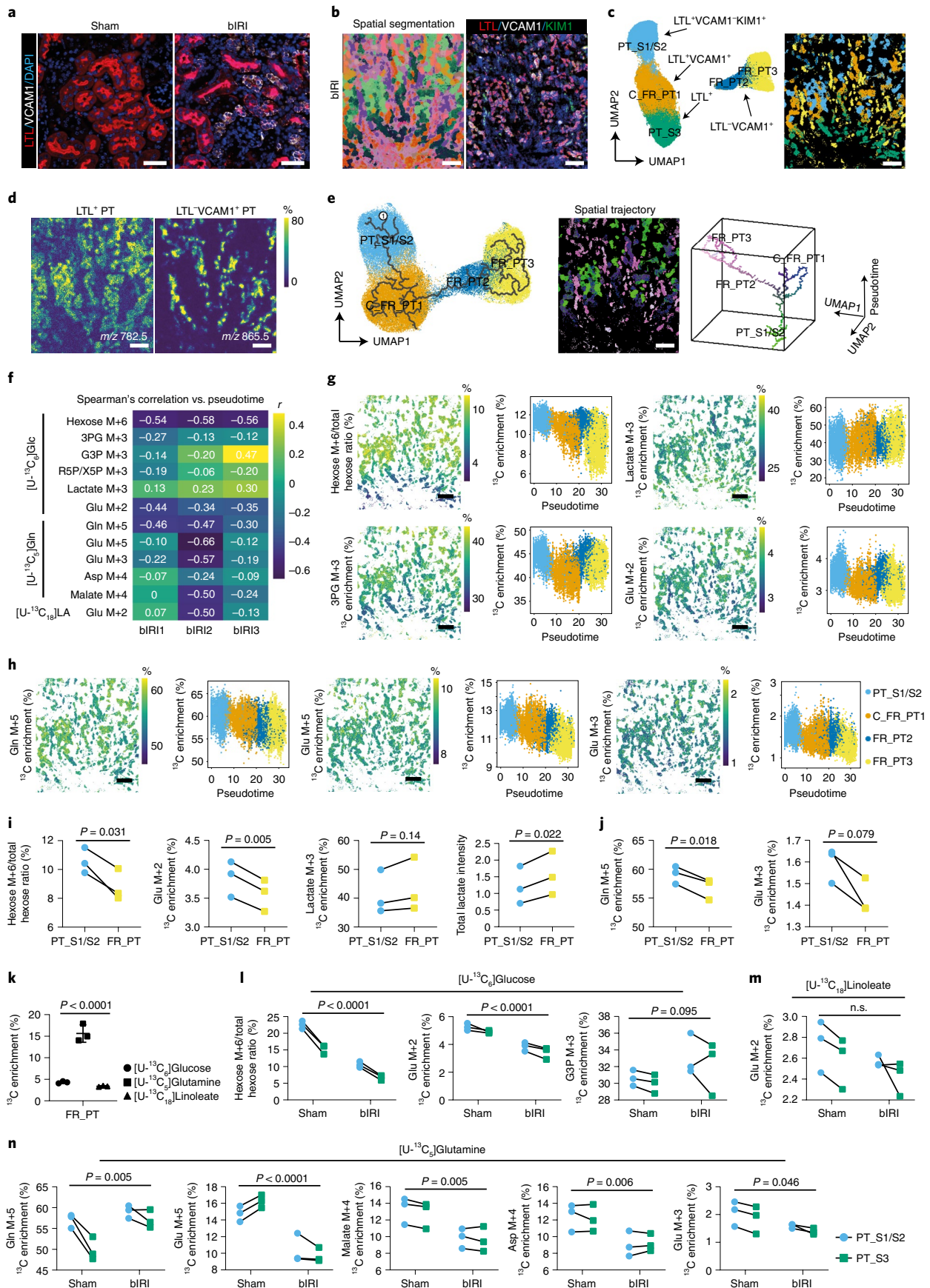
We then sought to identify metabolic (mal)adaptation after bIRI by comparing healthy PT (LTL⁺VCAM1⁻) with failed maladaptive repaired PT (VCAM1⁺; FR_PT) (Fig. 4a). Following MALDI-MSI, we selected all pixels with a PT, determined on the basis LTL, VCAM1, and KIM1 IF staining (Fig. 4b and Extended Data Fig. 6). Four types of PT could subsequently be identified on the basis of lipid profiling: LTL⁺VCAM1⁻KIM1⁻ (PT-S1/S2) and LTL⁺VCAM1⁺ (FR_PT1) in the cortex, and LTL⁻VCAM1⁺ (FR_PT2 and FR_PT3) and LTL⁺ in the outer stripe of the outer medulla (PT-S3) (Fig. 4c,d). The clusters present in the cortex areas showed a trajectory towards maladaptive repair on the basis of their lipid profiles (Fig. 4e). The spatial-trajectory map reflected progressive maladaptive repair in different areas (Fig. 4e), consistent with loss of LTL staining and increased VCAM1 expression. Notably, FR_PT and normal PT were localized in the same region (Fig. 4e), indicating that maladaptive failed repair could not be explained by a differential impact of only the ischemic insult. We then calculated the correlation of pseudotime scores with ^{13}C enrichment of several isotopologues (Fig. 4f). Following the trajectory from healthy PT areas toward progressively impaired PT areas, the pseudotime score was negatively correlated with $[\text{U-}^{13}\text{C}_6]\text{hexose}$ enrichment (mean $r = -0.56$, $P < 0.001$) and its downstream product $[\text{U-}^{13}\text{C}_2]\text{glutamate}$ (mean $r = -0.38$, $P < 0.001$) but was weakly positively correlated with $[\text{U-}^{13}\text{C}_3]\text{lactate}$ enrichment (mean $r = 0.22$, $P < 0.001$), in combination with more total lactate present in FR_PT (Fig. 4j). One possible explanation for this is a higher glycolytic flux to the production of lactate and a lower contribution of $[\text{U-}^{13}\text{C}_6]\text{glucose}$ to the TCA cycle in FR_PT (Fig. 4f,g,i), although it cannot be ruled out that other sources contribute to total lactate. A reduction of both $[\text{U-}^{13}\text{C}_5]\text{glutamine}$ enrichment and labeled-glutamine-derived isotopologues through the oxidative TCA cycle was also observed (Fig. 4f,h,j), but

glutamine, compared with glucose and linoleate, remained the main carbon source for glutamate in FR_PT (Fig. 4k). From these data, we conclude that the maladaptive repair in the PT is characterized by differences in production of lactate, which could be possibly the result of higher glycolytic activity and, as injury progresses, a concomitant reduction of TCA cycle metabolite consumption.

We then focused on normal PTs in the bIRI kidneys. Strikingly, when compared with the sham kidneys, these epithelial cells still displayed abnormal nutrient partitioning to TCA metabolism, both in the cortex as well as in the outer medulla (Fig. 4l,m). Normal PTs, 2 weeks after bIRI, had lower $[\text{U-}^{13}\text{C}_6]\text{hexose}$ and $[\text{U-}^{13}\text{C}_2]\text{glutamate}$ enrichment than sham kidneys (Fig. 4l). In addition, although normal PTs exhibited higher $[\text{U-}^{13}\text{C}_5]\text{glutamine}$ enrichment, we found a lower ^{13}C enrichment in specific isotopologues that are produced through the oxidative TCA cycle, such as $[\text{U-}^{13}\text{C}_5]\text{glutamate}$, $[\text{U-}^{13}\text{C}_4]\text{malate}$, $[\text{U-}^{13}\text{C}_4]\text{aspartate}$, and $[\text{U-}^{13}\text{C}_3]\text{glutamate}$ (Fig. 4n). This is remarkable, as the TCA is a central hub in cell function and a cell's ability to adapt to its environment. Its metabolites not only serve in the biosynthesis of nucleotides and macromolecules such as lipids and proteins, but also have recently been recognized to control gene regulation through post-translational histone modifications and DNA methylation¹⁹; hence, it will be of relevance to study how the disbalance of TCA metabolite homeostasis, even in normal PT cells after bIRI, relates to chronic progressive kidney injury or loss of repair after a new injury.

The dynamic nature of metabolism, and the metabolic heterogeneity of different cell types and phenotypes within the architecture of a single tissue, call for single-cell-resolution measurements of metabolic dynamics. Currently available methods can provide singular spatial metabolic snapshots, bulk dynamic metabolism of homogenized tissue material, or isotope tracing on tissue with

Fig. 4 | Dynamic metabolic measurements on bIRI kidney PT cells. **a**, Representative images of LTL and VCAM1 IF staining on sham and bIRI kidneys ($n=3$). **b**, Left, representative molecular histology of cortical and outer stripe outer medulla (left) areas of bIRI kidney ($n=3$), generated from UMAP of recorded lipid profiles. Right, IF staining following MALDI-MSI. **c**, UMAP plot (left) and representative spatial segmentation (right) showing lipid heterogeneity between PT cells from cortical and OSOM of bIRI kidneys ($n=3$). Spatial segmentation image showing the distribution of the UMAP clusters on tissue with same color. **d**, Representative lipid species distribution in cortical and outer stripe outer medulla areas of bIRI kidney ($n=3$). **e**, Left, Embedding of PT pixels from bIRI kidneys ($n=3$), showing the trajectory of PT injury using pseudotime lipidomics analysis (starting point 1). Middle, spatial-trajectory map showing the pseudotime score and UMAP embedding of each pixel on tissue. Right, 3D scatter plot of cell-type-specific trajectories, with the same colors as pixels in the spatial-trajectory map. **f**, Spearman's correlation between pseudotime and average ^{13}C isotopologue enrichment on all PT pixels. **g,h**, Dynamic metabolic measurements using $[\text{U-}^{13}\text{C}_6]\text{glucose}$ (**g**) or $[\text{U-}^{13}\text{C}_5]\text{glutamine}$ (**h**), and their trajectory changes on PT cells of bIRI kidneys ($n=3$). **i,j**, Dynamic metabolic comparison of $[\text{U-}^{13}\text{C}_6]\text{glucose}$ and total lactate levels (**i**) or $[\text{U-}^{13}\text{C}_5]\text{glutamine}$ measurements (**j**) between LTL⁺VCAM1⁻KIM1⁻ PT cells (PT-S1/S2) and LTL⁻VCAM1⁺ PT cells (FR_PT). A two-tailed paired *t*-test was performed. **k**, Direct carbon contribution of different nutrients to glutamate at 2 hours in FR_PT, from the glutamate isotopologues M+2, M+3, and M+5. One-way ANOVA was performed ($n=3$). Comparison of dynamic metabolic measurements using $[\text{U-}^{13}\text{C}_6]\text{glucose}$ (**l**), $[\text{U-}^{13}\text{C}_{18}]\text{linoleate}$ (**m**), or $[\text{U-}^{13}\text{C}_5]\text{glutamine}$ (**n**) on PTs from sham kidneys and healthy PTs from IRI kidneys. Two-way ANOVA was performed. Images and graphs show the average ^{13}C enrichment (area under curve (AUC) normalized to total time) of isotopologues derived from $[\text{U-}^{13}\text{C}_6]\text{glucose}$ or $[\text{U-}^{13}\text{C}_5]\text{glutamine}$, measured at different timepoints (0, 15, 30, 60, and 120 minutes), or $[\text{U-}^{13}\text{C}_{18}]\text{linoleate}$, measured at different timepoints (0, 60, and 120 minutes). Scale bars: (**a**) $50 \mu\text{m}$, (**b-e,g,h**) $200 \mu\text{m}$.



low spatial resolution^{1,2,6–9}. Other strategies, based on single-cell transcriptomics²⁰, cytometry by time of flight (CyTOF)²¹, or metabolic probes²², have yielded information on single-cell metabolism. However, these methods are based on the detection of surrogates for metabolic activity, such as metabolic enzymes and their transcripts. Here, we provide an in situ cell-type-specific measurement of metabolic dynamics through the direct detection of metabolic intermediates at single-cell resolution. Furthermore, to detect the dynamic metabolism of less abundant cell types in tissue, such as endothelial or immune cells, we introduced IF staining values to correlate with lipid profiles as a threshold for cell selection. For example, the combination of IF staining and lipid profiles allowed us to identify two populations of renal cortex endothelial cells: glomerular renal endothelial cell (gRECs) and cortical renal endothelial cells (cRECs) (Extended Data Fig. 7a–d). In line with known continuous VEGF signaling from podocytes²³, gRECs showed higher [¹³C]₃lactate enrichment than cRECs (Extended Data Fig. 7e). The combination of multiplexed IF staining on post-MSI-analyzed tissues with MALDI-MSI data shows great potential for multi-omics analysis by combining dynamic metabolomics with antibody-based spatial proteomics^{24,25}. This will link dynamic metabolism to the cell genotype and phenotype, giving more insight into metabolic regulated cell behavior. The accuracy of image co-registration of the spatial distribution of clustered pixels with IF staining, as in the bIRI study in this paper, is less critical than co-registration at the single-pixel level. To combine IF staining values with an MSI lipid dataset, a more advanced method for co-registration could be used²⁶. A single-cell segmentation step based on IF can also be used to study dispersed single-cell populations in tissues, such as immune cells²⁷. Although in this study we used pixels for UMAP and trajectory analysis, rather than individual cells after single-cell segmentation, both cell typing and trajectory were in line with IF staining, which indicates that single-cell-resolution pixels can be used for cell-type-specific analysis, as has been previously shown using spatial transcriptomics technologies^{28,29}.

To make this method potentially usable in human tissue biopsies, we used vibratome sectioning for stable-isotope tracing on tissue ex vivo, which provides the possibility of tracing different isotopes using a single tissue sample. However, the current 2-hour incubation period limits the dynamic measurement to central carbon metabolic pathways (glycolysis and TCA cycle) in metabolic non-steady state, as well as the detection of ¹³C-labeled TCA intermediates from one oxidative cycle. Nonetheless, this MALDI-MSI-based method for cell-type-specific measurement of metabolic dynamics can be applied to tissue samples or cultured cells for which longer incubation of stable isotopes is feasible, thus enabling the detection of ¹³C-labeled metabolites in a pseudo-steady-state. For instance, in endothelial cells cultured in vitro for 24 hours in the presence of [U-¹³C]₆glucose, we could detect not only the full range of isotopologues of TCA intermediates, such as citrate (M+1, M+2, M+3, M+4, and M+5) and succinate (M+1, M+2, M+3, and M+4), but also ¹³C-labeled metabolites from other metabolic pathways, such as ribose-5-phosphate/xylulose-5-phosphate (R5P/X5P), adenosine monophosphate (AMP), adenosine diphosphate (ADP), uridine diphosphate glucose (UDP-Glc), uridine diphosphate *N*-acetylglucosamine (UDP-GlcNAc), and glutathione (Extended Data Fig. 8). Owing to the limitations of MALDI-MS, we could not distinguish isomeric molecular species, such as glucose, galactose, mannose, and inositol⁹, as well as the confounders interfering with the quantification of labeling peaks, such as [¹³C]₂aspartate and [¹³C]₂malate. To solve this, more advanced instrumentation using ion-mobility separation is needed³⁰. Delocalization of metabolites is a limitation of spatial metabolomics with MALDI-MSI in general, which could cause false negatives by decreasing the signal differences between cell types, but is less likely to cause false positives³¹.

In summary, we have developed a platform to detect cell-type-specific dynamic metabolic changes at single-cell resolution. We demonstrated that this platform, applied to tissue samples, can discern metabolic heterogeneity within individual cell types in relation to the microenvironment. We used the kidney as an example of a complex and metabolically heterogeneous organ, but this method can be used to investigate tissue homeostasis in general. Our method could be applied to detect changes in cell-specific metabolism, which occur in processes such as inflammation, fibrosis, and cancer biology.

Methods

Reagents. All reagents were obtained from Sigma-Aldrich, unless stated otherwise.

Mouse studies. For studies of overall metabolic changes, post mortem material of 12-week-old male C57BL/6J mice ($n = 3$) culled as breeding surplus were used. Mice were kept and cared for in accordance with the Experiments on Animals Act (Wod, revision 2014, the Netherlands) and EU directive no. 2010/63/EU. Mice were housed at 20–22 °C in individually ventilated cages, humidity controlled (55%) with free access to food and water and a light/dark cycle of daytime (06:30–18:00) and nighttime (18:00–06:30).

For bIRI experiments, we used 12-week-old male constitutional renin reporter (B6.Ren1cCre/TdTomato/J) mice³². Six mice were divided into two groups randomly ($n = 3$ /group). In short, mice were placed under isoflurane anesthesia and controlled body temperature was kept at 36.7 °C, and renal arteries and veins were exposed through laparotomy with median incision on both sides. Subsequently, both arteries and veins were ligated using clamps for approximately 18 minutes, after which the clamps were removed to resume blood flow (reperfusion) and the abdomen was closed. For the sham controls, mice were opened up and closed similarly, but no ligations were performed. At day 14 after surgery, mice were perfused with cold PBS–heparin (5 IU/mL) via the left ventricle at a controlled pressure of 150 mmHg for 6 minutes to exsanguinate the kidneys before removal. Animal experiments were approved by the Ethical Committee on Animal Care and Experimentation of the Leiden University Medical Center (permit no. AVD1160020171145).

Vibratome sectioning and tissue slice incubation. Directly after euthanization, kidneys were collected and kept in ice-cold sterile Hanks' Balanced Salt Solution (HBSS) with 5 mM glucose and penicillin–streptomycin before vibratome sectioning. Each kidney was embedded in 4% low-melting-temperature agarose gel, and 350- μ m-thick tissue slices were obtained from fresh tissue under ice-cold HBSS with 5 mM glucose and penicillin–streptomycin using a Vibratome VT1200 (Leica Microsystems). Slicing speed was 0.1 mm/second, and vibration amplitude was 2 mm.

Tissue slices were placed into culture plates and incubated in a well-defined medium (nutrients-free Seahorse XF DMEM assay medium (Agilent, 103681), supplemented with 2% FCS, 3 mM linoleic acid (dissolved with addition of 1% BSA), 5 mM glucose, 500 μ M glutamine, 100 μ M sodium acetate, 50 μ M sodium citrate, and penicillin–streptomycin (pH adjusted to 7.4)) for up to 2 hours at 37 °C and 5% CO₂. During incubation, medium was changed to media containing various ¹³C-labeled nutrients at different time points. For the ¹³C-labeling incubation, the same amounts of either [U-¹³C]₆glucose (99%, Sigma, 389374), [U-¹³C]₆glutamine (99%, Cambridge Isotope Laboratories, Inc. CLM-1822-H), or [U-¹³C]₁₈linoleic acid (98%, Cambridge Isotope Laboratories, CLM-6855-0) were used to replace similar unlabeled nutrients in each medium. In the end, tissue slices were quenched with liquid N₂ and stored at –80 °C.

Tissue preparation and matrix deposition. Tissue slices were embedded in 10% gelatin and cryosectioned into 10- μ m-thick sections using a Cryostat NX70 cryostat (Thermo Fisher Scientific) at –20 °C. The sections were thaw-mounted onto indium-tin-oxide (ITO)-coated glass slides (VisionTek Systems). Mounted sections were placed in a vacuum freeze-dryer for 15 minutes prior to matrix application. After drying, *N*-(1-naphthyl) ethylenediamine dihydrochloride (NEDC) (Sigma-Aldrich, UK) MALDI-matrix solution of 7 mg/mL in methanol/ acetonitrile/deionized water (70, 25, 5% vol/vol/vol) was applied using a SunCollect sprayer (SunChrom). A total of 21 matrix layers were applied with the following flow rates: layer 1–3 at 5 μ L/min, layer 4–6 at 10 μ L/min, layer 7–9 at 15 μ L/min and 10–21 at 20 μ L/min (speed x, medium 1; speed y, medium 1; z position, 35).

MALDI-MSI measurement. MALDI-TOF/TOF-MSI was performed using a RapifleX MALDI-TOF/TOF system (Bruker Daltonics). Negative-ion-mode mass spectra were acquired at a pixel size of 5 \times 5 μ m² or 20 \times 20 μ m² over a mass range of m/z 80–1000. Prior to analysis, the instrument was externally calibrated using red phosphorus. Spectra were acquired with 15 laser shots per pixel (for 5 μ m measurement) or 200 laser shots per pixel (for 20 μ m measurement) at a laser repetition rate of 10 kHz. Data acquisition was performed using flexControl (Version 4.0, Bruker Daltonics) and visualizations were obtained from flexImaging

5.0 (Bruker Daltonics). All the samples from same slides were measured randomly. MALDI-FTICR-MSI was performed on a 12 T solarix FTICR mass spectrometer (Bruker Daltonics) in negative-ion mode, using 30 laser shots and 50- μm pixel size. Prior to analysis, the instrument was calibrated using red phosphorus. The spectra were recorded over a m/z range of 100–1,000 with a 2M data point transient and transient length of 0.5592 seconds. Data acquisition was performed using *ftmsControl* (Version 2.1.0, Bruker Daltonics), and visualizations were obtained from *flexImaging 5.0* (Bruker Daltonics). The images of lipid and metabolite distributions on tissue were exported from *flexImaging 5.0*. Following the MALDI-MSI data acquisition, excess matrix was removed by washing in 100% ethanol (2 \times 5 min), 75% ethanol (1 \times 5 min), and 50% ethanol (1 \times 5 min), after which this MSI-analyzed tissue section was used for IF staining, as described below.

Immunofluorescence staining. After MALDI-MSI tissues on the slide were fixed using 4% paraformaldehyde for 30 minutes, antigen retrieval was performed using antigen retrieval buffer (Dako, Agilent Technologies) in an autoclave. Slides were blocked with 3% normal donkey serum, 2% BSA, and 0.01% Triton X-100 in PBS for 1 hour at room temperature. Primary anti-mouse-KIM1 antibody (5 $\mu\text{g}/\text{mL}$, R&D Systems, MAB1817), anti-VCAM1 antibody (1:250, Abcam, ab134047), anti-CDH1 antibody (1:300, BD Biosciences, 610181), anti-NPHS1 (2 $\mu\text{g}/\text{mL}$, R&D Systems, AF4269), anti-mouse-pan-endothelial-cell-antigen (MECA32, 2 $\mu\text{g}/\text{mL}$, BD Biosciences, 553849), *Lotus tetragonolobus* Lectin (LTL, 1:300, Vector Laboratories, B-1325), or lectin from *Bandeiraea simplicifolia* isolectin B4 (BSI-TRITC, 1:200, Sigma, L5264) were incubated overnight at 4°C, followed by corresponding fluorescent-labeled secondary antibodies (donkey anti-rat-IgG AF488 (1:300, Invitrogen, A21208), donkey anti-rabbit-IgG AF647 (1:300, Invitrogen, A31573), donkey anti-mouse-IgG AF488 (1:300, Invitrogen, A21202), donkey anti-sheep-IgG AF568 (1:300, Invitrogen, A21099), streptavidin–Alexa Fluor 568 (1:300, Invitrogen, S11226), streptavidin–Alexa Fluor 647 (1:300, Invitrogen, S32357)) for 1 hour at 4°C, when necessary. Slides were embedded in Prolong gold antifade mountant with DAPI (Thermo Fisher Scientific, P36931). Fluorescent images of the slides were recorded using a 3D Histech Panoramic MIDI Scanner (Sysmex). Digital scanned images were co-registered to the MALDI-MSI data.

MSI data processing and analysis. MSI data were exported and processed in *SCIls Lab 2016b* (*SCIls*, Bruker Daltonics) with baseline correction using convolution algorithm. All MALDI-TOF-MSI data were normalized to the total ion count (TIC). Peak picking was performed (signal-to-noise ratio > 3) on the average spectrum, and matrix peaks were excluded from the m/z feature list. The m/z features which were present in both MALDI-FTICR-MSI and MALDI-TOF-MSI datasets, and which had similar tissue distributions, were further used for identity assignment of lipid species (Extended Data Fig. 9). The m/z values from MALDI-FTICR-MSI were imported into the Human Metabolome Database³³ (<https://hmdb.ca/>) after re-calibration in *mMass* and annotated for lipids species with an error < ± 5 ppm. For the small molecules detected only in MALDI-TOF, the m/z values from MALDI-TOF were imported into the Human Metabolome Database (<https://hmdb.ca/>) after re-calibration in *mMass* and annotated for metabolites with an error < ± 20 ppm. The ¹³C-labeled peaks were selected by comparing the spectrum of control and ¹³C-labeling experiments, and annotated on the basis of the presence of unlabeled metabolites and their theoretical m/z values. For measurement at 20 \times 20 μm^2 pixel size, a total of 230 high-molecular-weight features ($m/z > 400$, predominantly phospholipids¹², Extended Data Table 1) that not co-localized with matrix peaks were selected (signal-to-noise ratio > 3). For measurement at 5 \times 5 μm^2 pixel size, a total of 16 metabolites (9 unlabeled and 11 ¹³C-labeled), 227 high-molecular-weight features that were not co-localized with matrix peaks were selected (signal-to-noise ratio > 3). Peak intensities of the selected features were exported for all the measured pixels from *SCIls Lab 2016b*, which were used for the following analysis. Natural isotope abundance correction was performed for metabolites using *R* package *IsoCorrector*³⁴.

For targeted endothelial cell analysis at 5 \times 5 μm^2 pixel size, the high-resolution staining image of MECA32 and NPHS1 on the MALDI-MSI measured area was exported from *CaseViewer* (3DHISTECH). Staining in the areas not measured by MALDI-MSI was removed. The resolution of the exported image was changed to 5 \times 5 μm^2 pixel size using *Matlab R2019a* to comply with the MALDI-MSI resolution. By merging it with the previous high-resolution image, the pixels fully covering MECA32 staining were selected as endothelial cells, and non-fully-covered pixels were discarded from the image. Next, the staining values of all the pixels were exported using *Matlab R2019a* with a cutoff of 5% (values < 13 were replaced with 0). In the end, the staining values corresponding to the same pixels as MALDI-MSI measurements were integrated into MALDI-MSI data and used for data analysis as described below.

For UMAP analysis, the datasets were transformed into a count matrix by multiplying the TIC-normalized intensities by 100 and taking the integer. This count data matrix was normalized and scaled using *SCTransform* to generate a 2D UMAP map using *Seurat 3.0* in *R* (version 4.0). The distribution of the pixels from different clusters on tissues were co-registered to the IF staining, and cell

types were identified on the basis of both staining and their morphology on the kidney. Using measurements of endothelial cells, two endothelial cell clusters were identified on the basis of MECA32 staining and their position, and a podocyte cluster was identified on the basis of NPHS1 staining. For data imputation, the metabolite m/z features from the ‘control’, or unlabeled, dataset were removed, so only lipid m/z features were left, which were used as the query. Then, the MALDI-MSI data from ¹³C-labeling experiments were used as a reference to transfer metabolite production into the query using *FindTransferAnchors* and *TransferData* function from the *Seurat 3.0* package in *R*. Both the query and reference were normalized and scaled using *SCTransform*. Ultimately, all the imputed metabolite productions were combined into one dataset, which contained the ¹³C-labeling information over time.

After combining all the imputed data into one dataset, the fraction enrichment of isotopologues was calculated on the basis of the ratio of each ¹³C-labeled metabolite (isotopologue) to the sum of this metabolite abundance in each pixel. The calculated fraction enrichment of isotopologues was used to generate pseudoimages together with pixel coordinate information exported from *SCIls Lab 2016b*. The average fraction enrichment values (AUC normalized to total time) of identified clusters were used for generating graphs and statistical analysis. Hotspot removal (high quantile 99%) were applied to all the pseudoimages generated from calculated values.

The integrated data matrix from 3 IRI kidneys from *Seurat 3.0* was further used for the trajectory analysis using *Monocle3* (ref.³⁵) in *R*. A spatial trajectory map was generated using *Matlab R2019a* based on embedding information of UMAP and pseudotime values calculated by *Monocle3* and pixel coordinate information exported from *SCIls Lab 2016b* as described below.

For spatial UMAPs, the datasets were used to generate a 3D UMAP map using packages *Seurat 3.0* and *plotly*. The embedding information of the 3D UMAP was translated to RGB color coding by varying red, green and blue intensities on the three independent axes. Together with pixel coordinate information exported from *SCIls Lab 2016b*, a $\text{MxN} \times 3$ data matrix was generated and used to generate UMAP images in *Matlab R2019a*.

Data collection and analysis were not performed blind to the conditions of the experiments. A step-by-step protocol associated with this study is available in Protocol Exchange (<https://doi.org/10.21203/rs.3.pev-1912/v1>)³⁶.

Validation test of imputation performance. Validation of the imputation performance using ‘leave-one factor-out’ cross-validation is shown in Extended Data Fig. 3. In short, the metabolite production from MALDI-MSI data was taken out from one kidney slice (sample 1, Extended Data Fig. 3a) or two different kidney slices (sample 3, Extended Data Fig. 3h) and use samples 2 and 4 to impute the metabolite production in sample 1 and in sample 3, respectively, as described above. Next, Spearman’s correlation was calculated between the imputed and detected values in sample 1 and 3. Values higher than 0.8 were graded as a very strong positive correlation. Values between 0.6 to 0.8 were a strong positive correlation, and between 0.4 to 0.6 a moderate positive correlation.

Statistics and reproducibility. No statistical methods were used to predetermine sample sizes, but our sample sizes are similar to those reported in previous publications^{9,16}. All the experiments and data analysis were performed in triplicate (3 animals per group). No animals or data points were excluded from the analysis. All the data are presented as mean \pm s.d., unless indicated otherwise. The average fractional contribution values of identified clusters were used for statistical analysis. AUC was used for comparison of the changes following timecourse of isotope tracer incubation up to 2 hours between different groups. Data normality and equal variances were tested using the Shapiro–Wilk test. Differences between groups were assessed by paired two-tailed Student’s *t*-test, two-tailed Student’s *t*-test, when not normally distributed, or by two-tailed *F*-test. $P < 0.05$ was considered statistically significant.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The exported and processed MSI data for this study were deposited in FigShare at <https://doi.org/10.6084/m9.figshare.20227419.v1>. Owing to the large size of all raw data, parts of the raw MSI data are deposited to provide the necessary information of ¹³C-labeled metabolites and spectrum quality. For full raw MALDI-MSI data related to this study, please contact G. W. (g.wang@lumc.nl) or B. H. (b.p.a.m.heijs@lumc.nl). Upon reasonable request, data will be made available. Source data are provided with this paper.

Code availability

The codes used in this article is available in Github (<https://github.com/GangqiWang/scDYSMO>).

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Author contributions

G. W. designed research study, conducted experiments, processed data, and wrote the manuscript. B. H. provided the technical guidance of MALDI-MSI experiments and data processing and wrote the manuscript. S.K. and M.G. provided expert knowledge on metabolomics and lipidomics analysis, read the manuscript, and provided helpful comments. R.B., A.K., and L.A.K.v.d.P. performed the animal experiments, read the manuscript, and provided helpful comments. A.M. provided professional instruction of single-cell data analysis, read the manuscript, and provided helpful comments. R.G.J.R. and C.W.v.d.B. helped with the experiments, read the manuscript, and provided helpful comments. P.C. and S.J.D. read the manuscript and provided helpful comments. B.M.v.d.B. and T.J.R. organized funding, designed research study, and wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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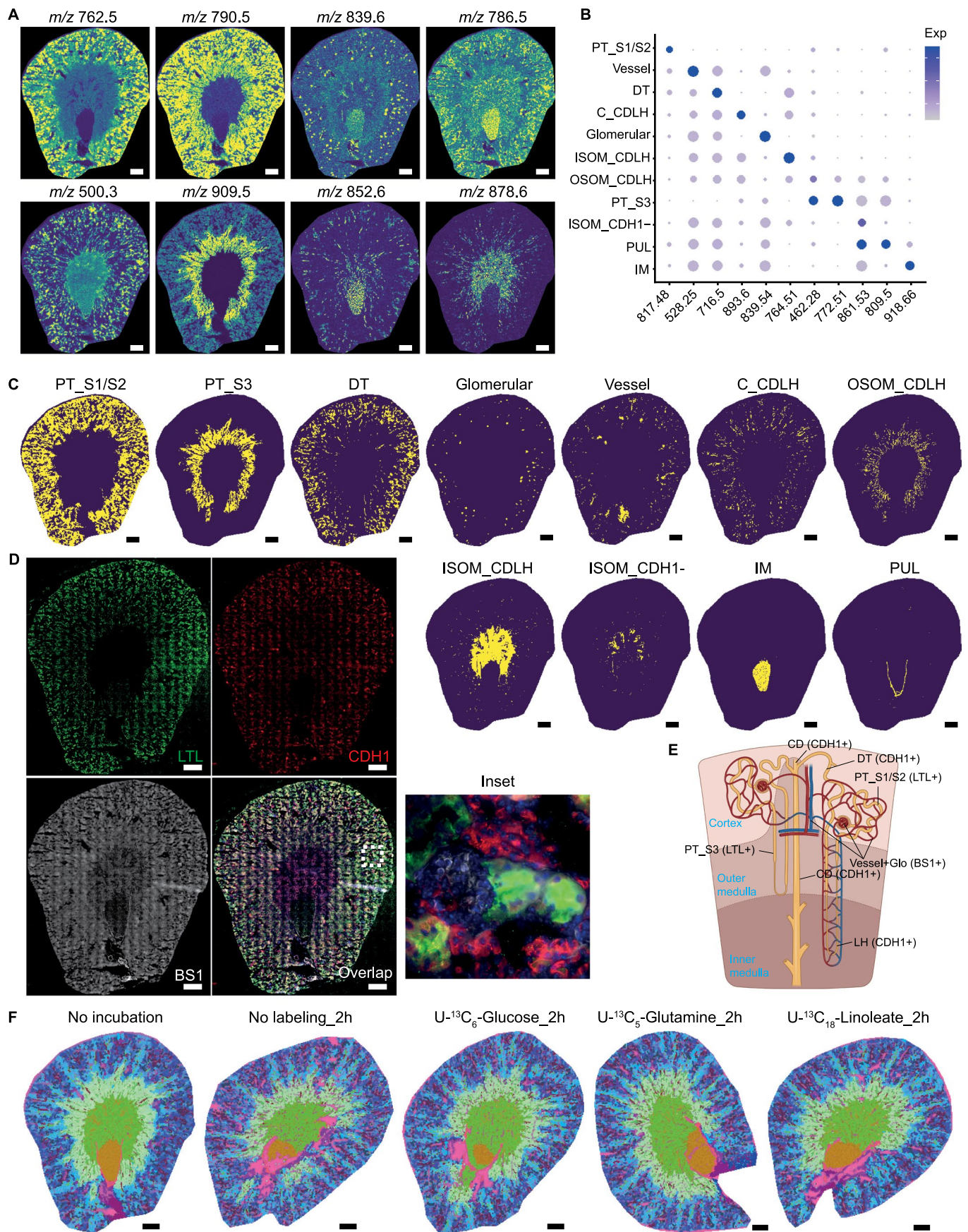
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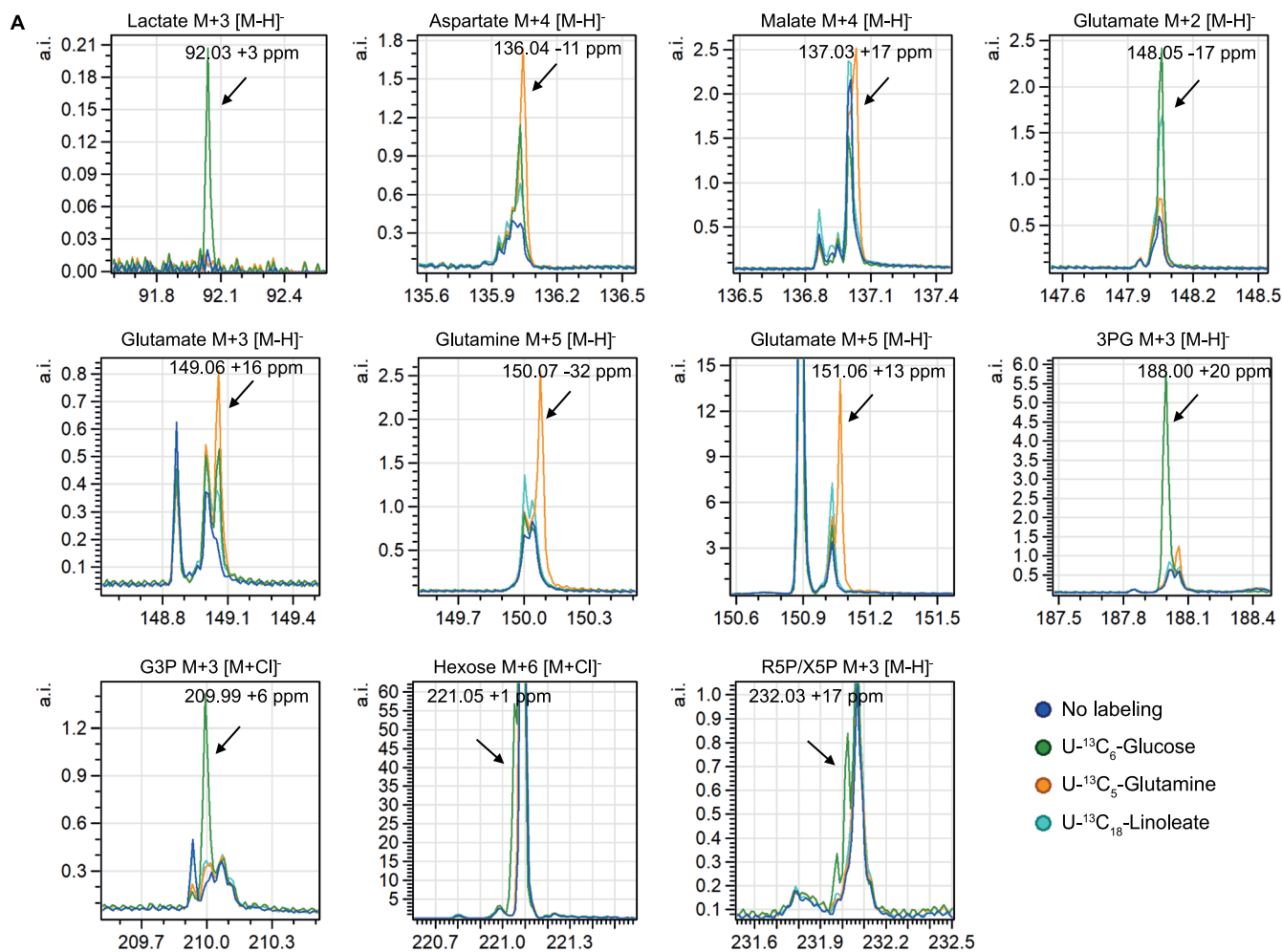
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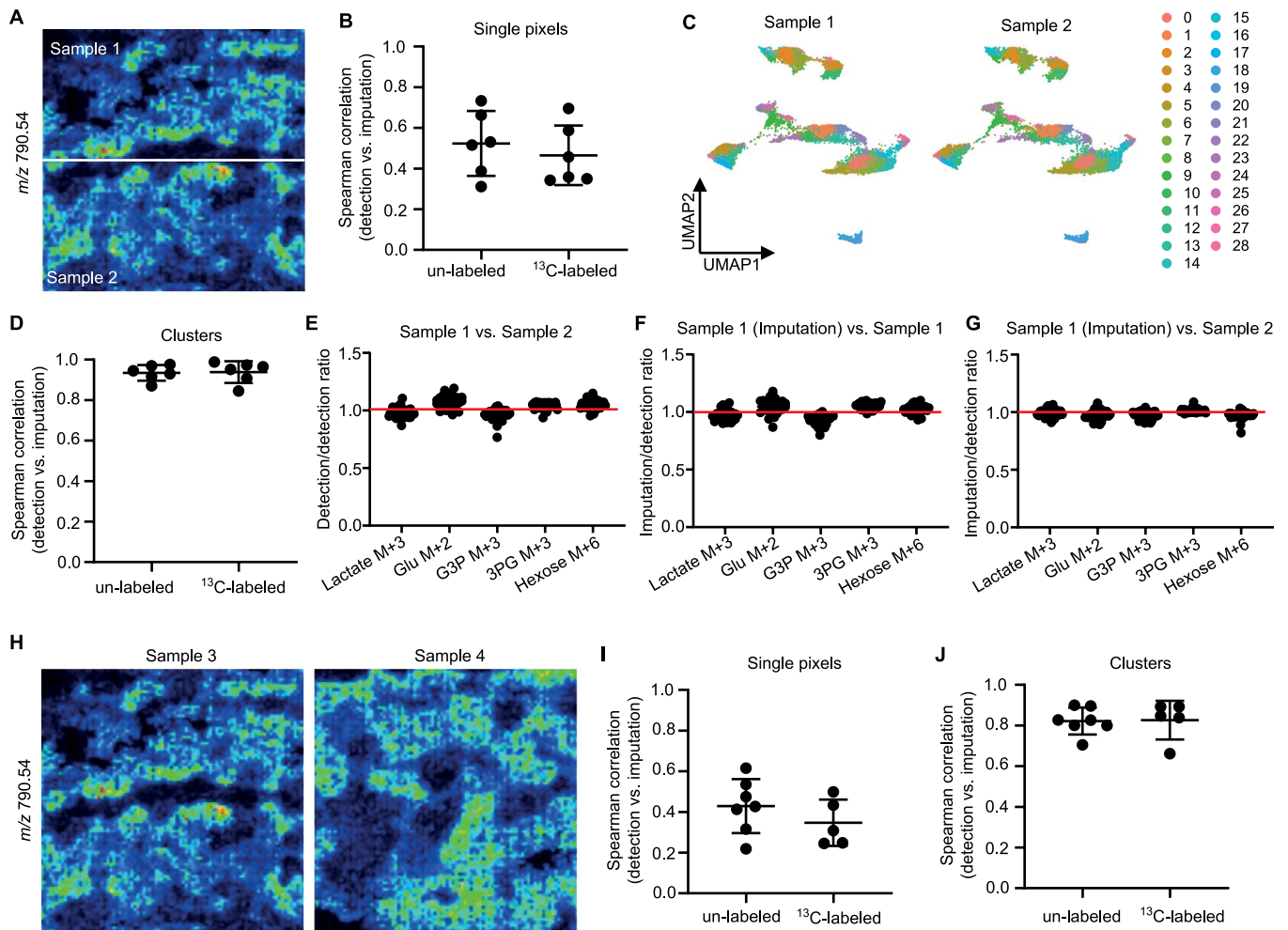


Extended Data Fig. 1 | See next page for caption.

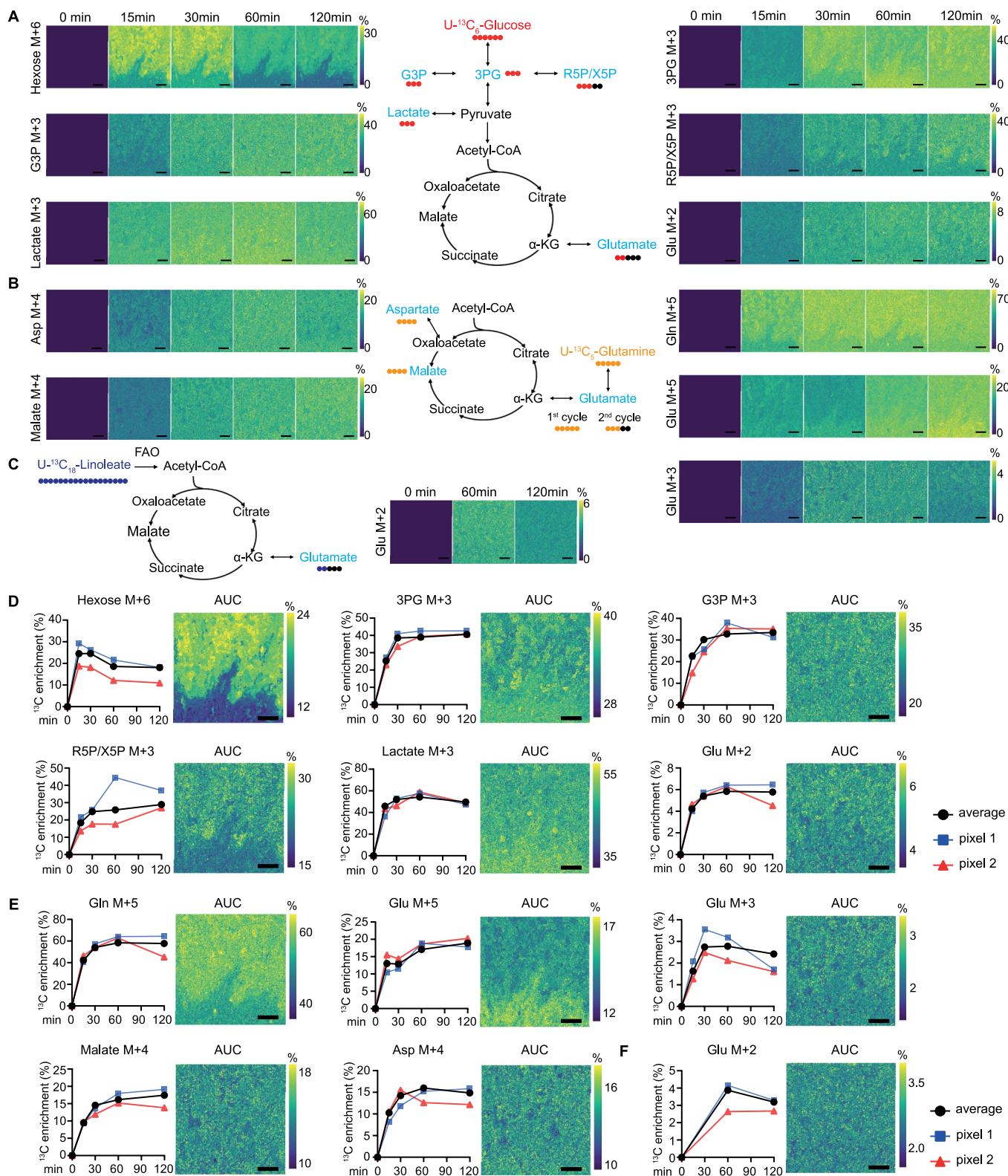
Extended Data Fig. 1 | Lipid heterogeneity and stability in mouse kidneys. **a**, Lipid species distribution in a mouse kidney tissue sample as recorded by MALDI-MSI at $20 \times 20 \mu\text{m}^2$ pixel-size. **b**, Dot plot displaying lipid expression of cluster-enriched signatures. **c**, Distribution of different renal cell clusters on tissue as identified in Fig. 2a. **d**, Immunofluorescent staining (*Lotus Tetragonolobus* Lectin (LTL, green), E-cadherin (CDH1, red) and BS1-lectin (gray)) after MALDI-MSI measurement. **e**, Nephron anatomy showing the identification of renal cell types based on immunofluorescent staining. **f**, Molecular histology of kidneys generated from integrated three-dimensional UMAP analysis of different datasets based on lipid profiles showing the specificity and stability of lipids profiles after introducing ^{13}C -labeled nutrients for 2 hours. All scale bars = $500 \mu\text{m}$. Abbreviations: PT_S1/S2, cortical proximal tubular segments 1/2; PT_S3, outer stripe of outer medulla proximal tubular segment; DT, distal tubule; CDLH, collecting duct and loop of Henle; IM, inner medulla; ISOM, inner stripe of outer medulla; OSOM, outer stripe of outer medulla; PUL, pelvic urothelial lining.



Extended Data Fig. 2 | Mass spectrum of measured ¹³C-labeled Metabolites. Mass spectra obtained from ¹³C-labeled- and control samples measured by MALDI-MSI at 5 × 5 μm² pixel size. Graphs show presence of various ¹³C-labeled metabolites.

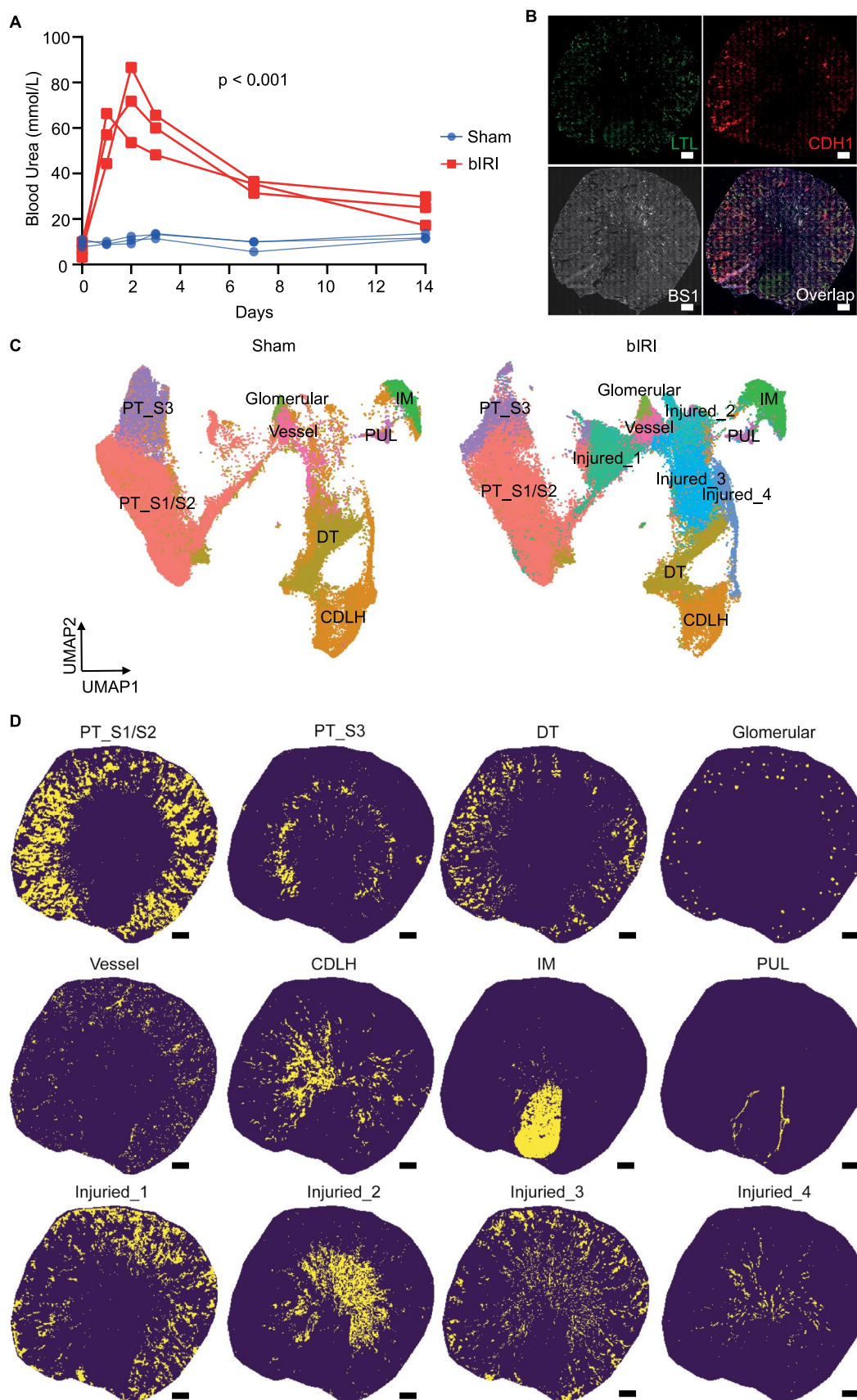


Extended Data Fig. 3 | Validation of the imputation performance - 'leave-one factor-out' cross-validation. **a**, MALDI-MSI data from a single kidney measurement, which was incubated with $U\text{-}^{13}\text{C}_6\text{-glucose}$ for 2 h, was bisected (into sample 1 and sample 2) and used to test the accuracy of the imputation. m/z features corresponding to metabolites were taken out from sample 1. Based on these lipid profiles, metabolite abundance of sample 2 was used to impute their abundance in sample 1. **b**, Spearman's correlation analysis on the imputed and detected metabolites in kidney sample 1 based on all the pixels. Dots represent different metabolites. **c**, UMAP analysis on integrated MALDI-MSI data of the 2 samples and resulting 29 clusters. **d**, Spearman's correlation analysis on the average value of each cluster between the imputed and detected metabolites abundance in kidney sample 1. **e**, The ratio of the ^{13}C enrichment calculated from detected metabolites abundance in the 29 clusters between sample 1 and sample 2. Dots represent different clusters. **f**, The ratio of the ^{13}C enrichment calculated from the imputed and detected metabolites abundance in the 29 cell clusters of kidney sample 1. Dots represent different clusters. **g**, The ratio of the ^{13}C enrichment calculated from the imputed metabolites abundance of sample 1 from sample 2 and detected metabolites abundance of kidney sample 2 in the 29 cell clusters. Dots represent different clusters. **h**, MALDI-MSI data from measurements of two different kidneys, which were incubated with $U\text{-}^{13}\text{C}_6\text{-glucose}$ for 2 h, were used to test the performance of the imputation between different measurements. **i**, Spearman's correlation analysis on the imputed and detected metabolites production based on all the pixels. Dots represent different metabolites. **j**, Spearman's correlation analysis on the average value of each cluster between the imputed and detected metabolites abundance. Dots represent different metabolites. All the performance test were done on 3 different imputations from different biological independent samples.

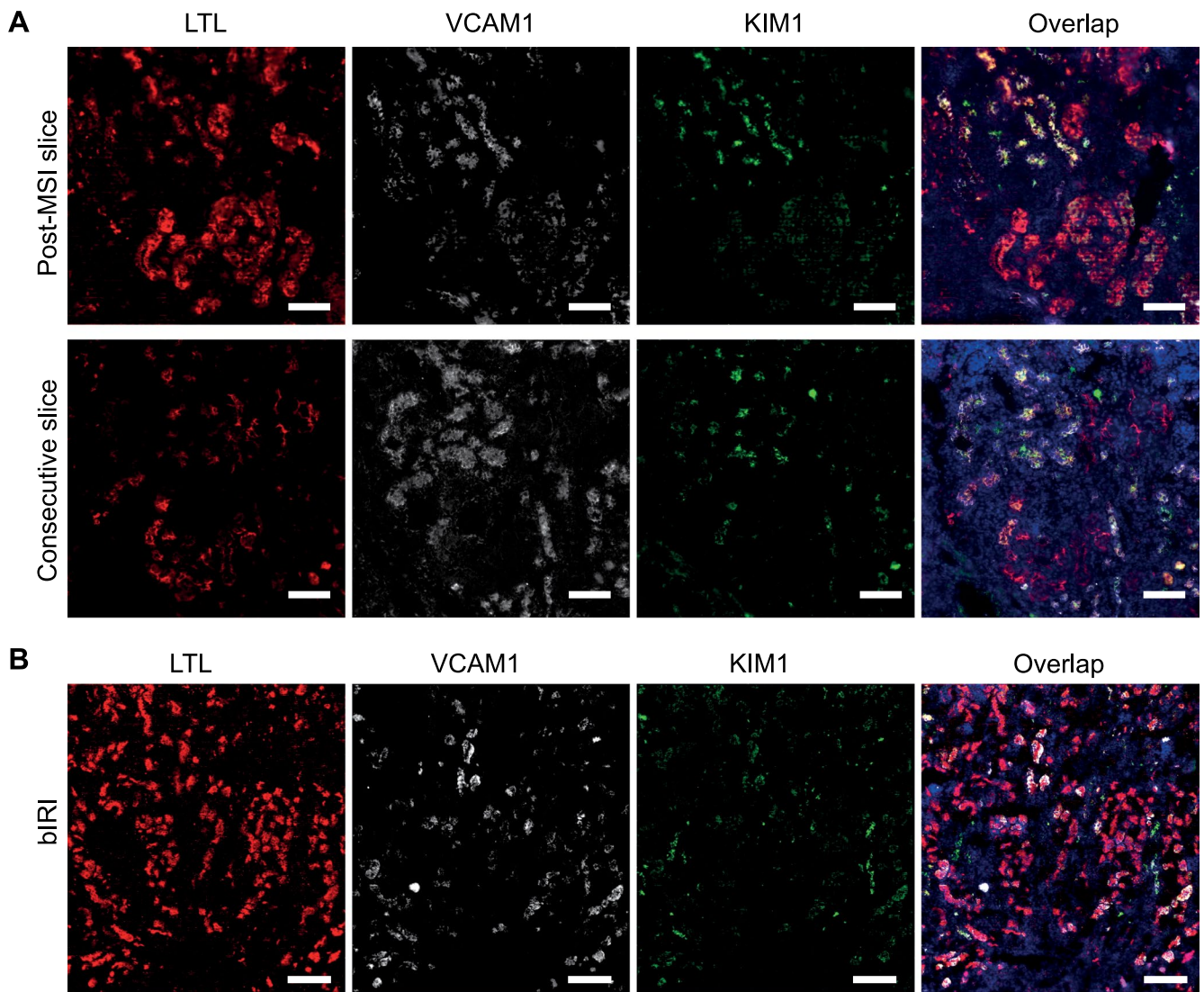


Extended Data Fig. 4 | See next page for caption.

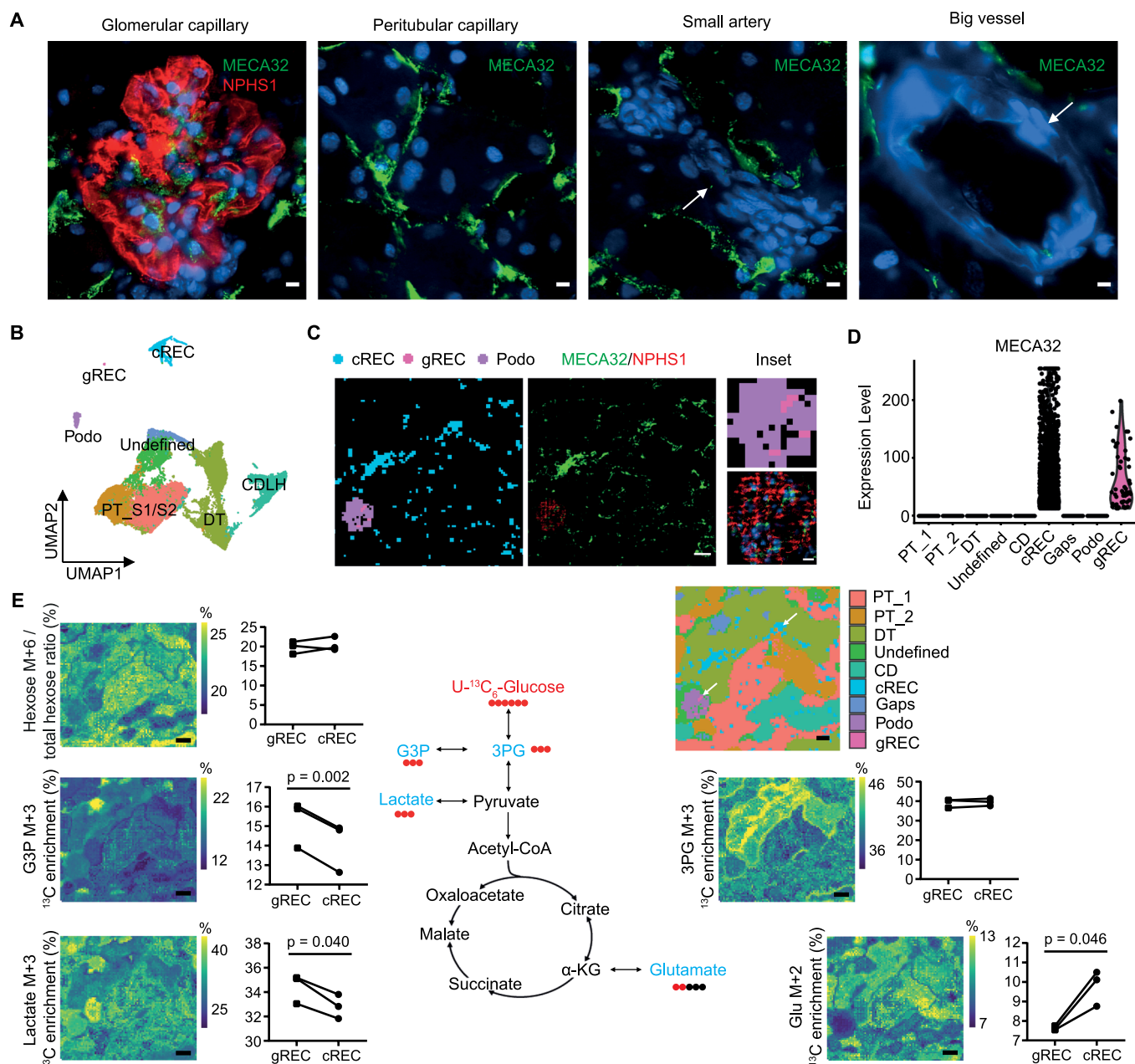
Extended Data Fig. 4 | Metabolic dynamics measurements on mouse kidney. **a**, Pseudo-images showing the ^{13}C enrichment of isotopologues of different metabolites over a time course (up to 2 h) of incubation with $\text{U-}^{13}\text{C}_6$ -glucose. Color scale representing the ^{13}C enrichment (%). Molecular histology and immunofluorescent staining on post-MALDI-MSI tissue are shown in Fig. 3a. **b**, Pseudo-images showing the ^{13}C enrichment of isotopologues of different metabolites during incubation with $\text{U-}^{13}\text{C}_5$ -glutamine for 2 h. Color scale representing the ^{13}C enrichment (%). **c**, Pseudo-images showing the ^{13}C enrichment of Glu M + 2 isotopologues during incubation with $\text{U-}^{13}\text{C}_{18}$ -linoleate for 2 h. Color scale representing the ^{13}C enrichment (%). **d-f**, Graphs showing the curve of ^{13}C enrichment of isotopologues of metabolites over a time course (up to 2 h) with incubation of $\text{U-}^{13}\text{C}_6$ -glucose (d), $\text{U-}^{13}\text{C}_5$ -glutamine (e) and $\text{U-}^{13}\text{C}_{18}$ -linoleate (f). Average values of all pixels and values from two representative pixels are shown. Images show the average ^{13}C enrichment (area under curve (AUC) normalized to total time) of isotopologues measured at different timepoints. All scale bars = 200 μm . Abbreviations: 3PG, 3-phosphoglycerate; G3P, glycerol 3-phosphate; α -KG, alpha-ketoglutaric acid.



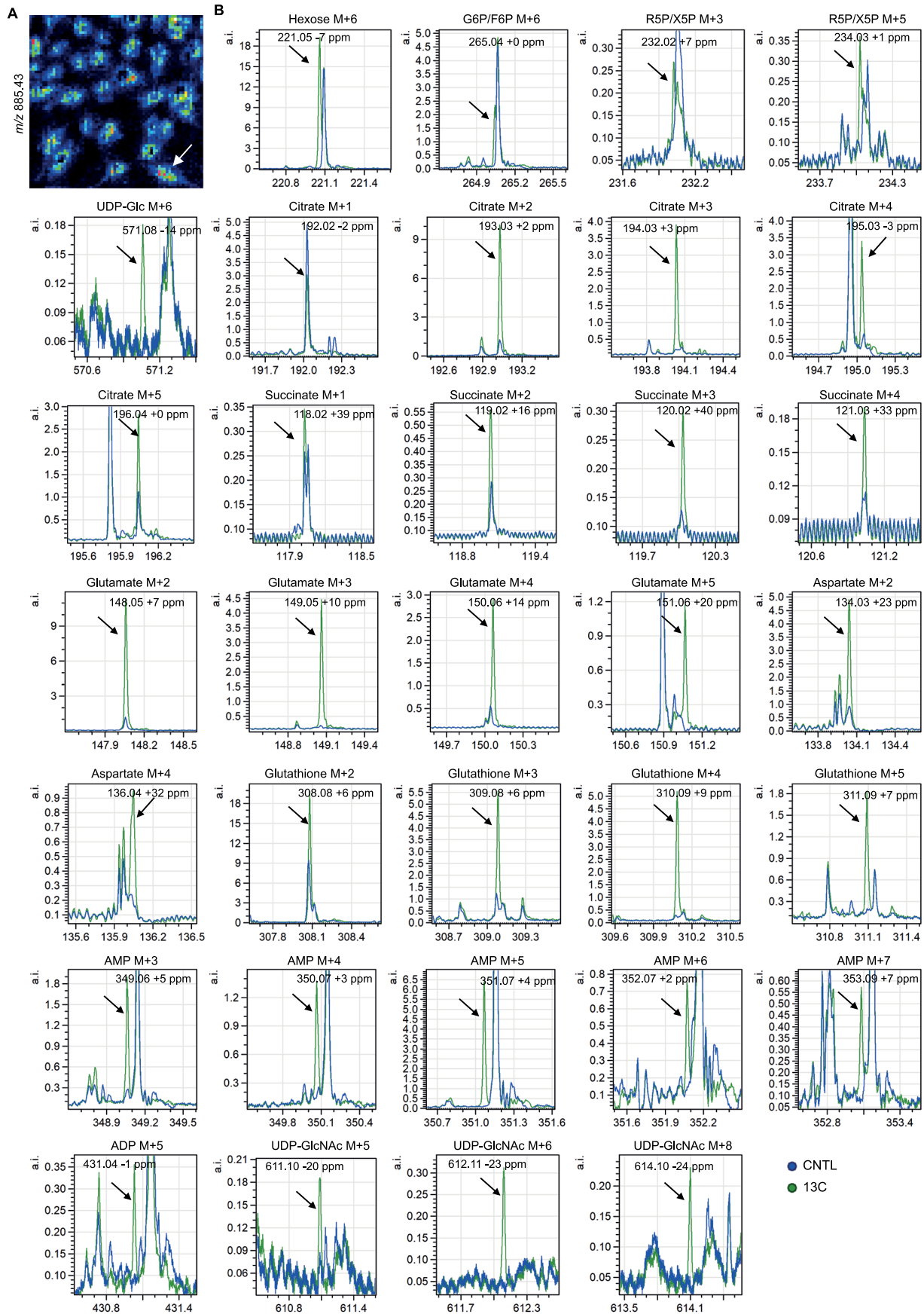
Extended Data Fig. 5 | Lipid heterogeneity in mouse IRI kidneys. **a**, Comparison of blood urea levels at different timepoints after IRI surgery ($n=3$ per group). Two tailed T test was performed on AUC. **b**, Immunofluorescent staining (*Lotus Tetragonolobus* Lectin (LTL, green), E-cadherin (CDH1, red) and BS1-lectin (gray)) after MALDI-MSI measurement of mouse IRI kidney. **c**, UMAP analysis on integrated lipidomics data of sham ($n=3$) and bIRI ($n=3$) mouse kidney samples. **d**, Distribution of different renal cell clusters on tissue as identified in Fig. 2b. All scale bars = 500 μm .



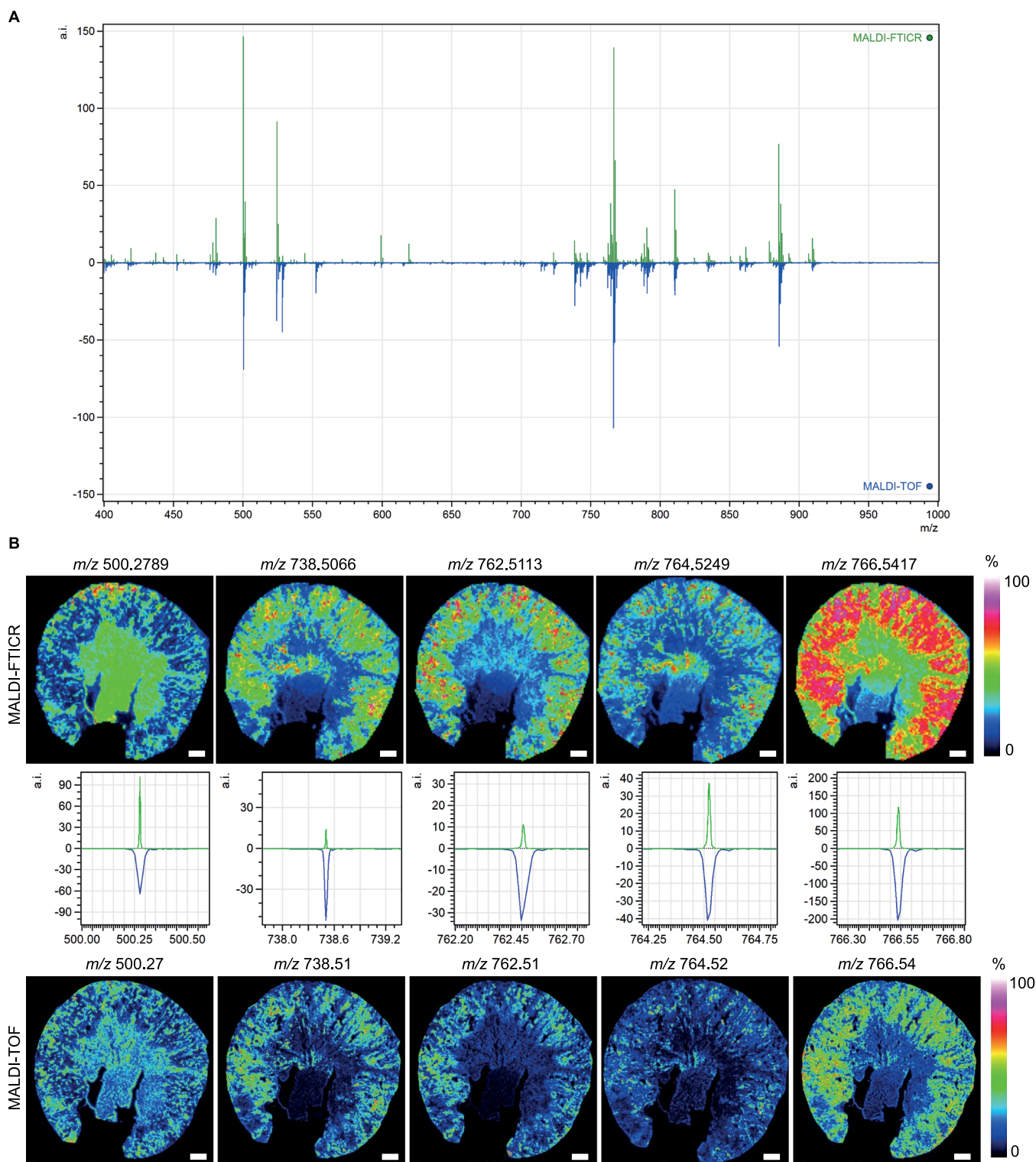
Extended Data Fig. 6 | Immunofluorescence staining on mouse IRI kidneys. a, Validation of immunofluorescence staining of LTL, VCAM1 and KIM1 on the measured area of post-MSI slice compared to consecutive slice without MALDI matrix. Scale bar = 100 μ m. **b**, Immunofluorescence staining of LTL, VCAM1 and KIM1 on bIRI kidney, same as the image shown in Fig. 4b. Scale bar = 200 μ m.



Extended Data Fig. 7 | Dynamic metabolic measurements on kidney endothelial cells. **a**, MECA32 staining of different vessels in mouse kidney to show specific capillary endothelial cell staining (Scale bar = 5 μm). Arrows depict the vessels with negative MECA32 staining. **b**, Metabolic heterogeneity within cortical cells in kidney, visualized in a two-dimensional UMAP plot of combined MALDI-MSI data and immunofluorescence staining at 5 \times 5 μm^2 pixel-size. Glomerular endothelial cells (gREC) and peritubular endothelial cells (cREC) were identified using the post-MALDI-MSI anti-MECA32 antibody staining and podocytes (Podo) with anti-NPHS1 antibody staining. Proximal tubules (PT), distal tubules (DT) and collecting ducts (CD) were identified based on their respective lipid profiles obtained from measurement shown in Extended Data Fig. 1b. **c**, Immunofluorescent (MECA32 and NPHS1, middle panel; scale bar = 40 μm) staining on post-MALDI-MSI tissue to determine gREC, cREC and podocyte distribution (left panel). Pixel colors used are similar to the given UMAP cell-cluster colors in A. Inset view of detailed glomerular area (right panels, scale bar = 5 μm). **d**, MECA32 expression in the observed cell types. **e**, Dynamic metabolic measurements using U- $^{13}\text{C}_6$ -glucose. Graphs show the ^{13}C enrichment of isotopologues of glucose, glycerol 3-phosphate (G3P), 3-phosphoglycerate (3PG), lactate and glutamate over time in gREC and cREC. Two tailed paired t-test was performed (n = 3). Spatial UMAP shows the metabolic cellular architecture of the tissue (arrows depict cREC and gREC). Pixel colors used are similar to the given UMAP cell-cluster colors in B. Scale bar = 200 μm .



Extended Data Fig. 8 | High spatial resolution MALDI-MSI measurement on cultured endothelial cell. a, MALDI-MSI data from measurements of cultured endothelial cells, which were incubated with U-¹³C₆-glucose for 24 h. Arrow depicts a single endothelial cell. **b**, Mass spectra obtained from ¹³C-labeled- and control samples measured by MALDI-MSI at 5 × 5 μm² pixel size to show the presence of ¹³C-labeled metabolites.



Extended Data Fig. 9 | Validation of MALDI-FTICR and MALDI-TOF data. a, Comparison of the average mass spectrum of an area with $50 \times 50 \mu\text{m}^2$ pixel-size (top, green) acquired by MALDI-FTICR and the average spectrum of $20 \times 20 \mu\text{m}^2$ pixel-size (bottom, blue) acquired by MALDI-TOF.

b, Comparison of the lipids distribution recorded by MALDI-FTICR at $50 \times 50 \mu\text{m}^2$ pixel-size and MALDI-TOF at $20 \times 20 \mu\text{m}^2$ pixel-size on same kidney. Scale bar = $500 \mu\text{m}$.

Extended Data Table 1 | Example of lipids in mouse kidney assigned by the mass measurements using both MALDI-TOF/TOF and MALDI-FTICR MS in negative ion mode

m/z from MALDI-TOF/TOF	m/z from MALDI-FTICR	Ion	Abbreviation*	Theoretical m/z	Formula	ppm***
419.2556	419.2578	[M-H]-	CPA(18:0)	419.2568	C21H41O6P	2.39
437.2674	437.2683	[M-H]-	LPA(18:0)	437.2674	C21H43O7P	2.06
452.2761	452.2789	[M-H]-	LPA(16:0)	452.2783	C21H44NO7P	1.33
453.2796	453.2826	[M-H]-	LPA(16:0)**	453.2817	C21H44NO7P	1.99
457.2335	457.2372	[M-H]-	LPA(20:4)	457.2361	C23H39O7P	2.41
462.2984	462.2998	[M-H]-	GPE(18:2)	462.299	C23H46NO6P	1.73
464.3133	464.3135	[M-H]-	LPE(18:0)	464.3146	C23H48NO6P	2.37
476.2749	476.2791	[M-H]-	LPE(18:2)	476.2783	C23H44NO7P	1.68
478.2925	478.2948	[M-H]-	LPE(18:1)	478.2939	C23H46NO7P	1.88
480.3085	480.3105	[M-H]-	LPE(18:0)/LPC(15:0)	480.3096	C23H48NO7P	1.87
481.3106	481.3139	[M-H]-	LPE(18:0)/LPC(15:0)**	481.313	C23H48NO7P	1.87
483.27	483.2718	[M-H]-	LPG(16:0)	483.2728	C22H45O9P	2.07
498.2584	498.2603	[M-H]-	LPE(20:5)	498.2626	C25H42NO7P	4.62
500.2746	500.2789	[M-H]-	LPE(20:4)	500.2783	C25H44NO7P	1.20
501.2767	501.2827	[M-H]-	LPE(20:4)**	501.2817	C25H44NO7P	1.99
524.2776	524.279	[M-H]-	LPE(22:6)	524.2783	C27H44NO7P	1.34
525.2781	525.2824	[M-H]-	LPE(22:6)**	525.2817	C27H44NO7P	1.33
526.2878	526.2913	[M-H]-	LPE(22:5)	526.2939	C27H46NO7P	4.94
556.3013	556.3049	[M-H]-	PC(20:4)	556.3045	C28H48NO8P	0.72
571.287	571.2892	[M-H]-	LPI(16:0)	571.2889	C25H49O12P	0.53
599.323	599.3189	[M-H]-	LPI(18:0)	599.3202	C27H53O12P	2.17
616.4742	616.4737	[M-H]-	CerP(d34:1)	616.4711	C34H68NO6P	4.22
619.2904	619.289	[M-H]-	LPI(20:4)/PI(20:4)	619.2889	C29H49O12P	0.16
647.4666	647.4669	[M-H]-	PA(32:0)	647.4657	C35H69O8P	1.85
671.4623	671.4665	[M-H]-	PA(34:2)	671.4657	C37H69O8P	1.19
673.48	673.479	[M-H]-	PA(34:1)	673.4814	C37H71O8P	3.56
685.5342	685.5264	[M-H]-	SM(d33:2)	685.529	C38H75N2O6P	3.79
687.5494	687.5448	[M-H]-	SM(d33:1)	687.5446	C38H77N2O6P	0.29
688.555	688.5492	[M-H]-	SM(d33:1)**	688.548	C38H77N2O6P	1.74
696.4726	696.4718	[M+Cl]-	LPC(28:1)/PE(p-31:0)	696.474	C36H72NO7P	3.16
697.4789	697.4837	[M-H]-	PA(36:3)	697.4814	C39H71O8P	3.30
699.4979	699.4971	[M-H]-	PA(36:2)	699.497	C39H73O8P	0.14
701.5135	701.5095	[M-H]-	PA(36:1)	701.5127	C39H75O8P	4.56
702.5229	702.5169	[M-H]-	PA(36:1)**	702.5161	C39H75O8P	1.14

705.4889	705.4856	[M-H]-	Cer(d17:1/LTE4)	705.4882	C40H70N2O6S	3.69
714.5084	714.5112	[M-H]-	PE-NMe(33:0)/PE-NMe2(32:2)/PE(34:2)	714.5079	C39H74NO8P	4.62
715.5134	715.5099	[M-H]-	PE-NMe(33:0)/PE-NMe2(32:2)/PE(34:2)**	715.5113	C39H74NO8P	1.96
716.5222	716.5246	[M-H]-	PE-NMe(33:1)/PE-NMe2(32:1)/PE(34:1)/PC(31:1)	716.5236	C39H74NO8P	1.40
717.5238	717.529	[M-H]-	PE-NMe(33:1)/PE-NMe2(32:1)/PE(34:1)/PC(31:1)**	717.527	C39H74NO8P	2.79
719.4979	719.4659	[M-H]-	PA(38:6)	719.4657	C41H69O8P	0.28
722.511	722.4785	[M-H]-	PE-NMe(34:5)/PE(35:5)/PC(32:5)	722.4766	C40H70NO8P	2.63
723.5082	723.4997	[M-H]-	PA(38:4)	723.497	C41H73O8P	3.73
724.5126	724.5033	[M-H]-	PA(38:4)**	724.5004	C41H73O8P	4.00
725.5126	725.5097	[M-H]-	PA(38:3)	725.5127	C41H75O8P	4.14
736.4927	736.4941	[M-H]-	PE-NMe(35:5)/PE-NMe2(34:5)/PE(33:5)	736.4923	C41H72NO8P	2.44
738.5087	738.5066	[M-H]-	PE-NMe(35:4)/PE-NMe2(34:4)/PE(36:4)/PC(33:4)	738.5079	C41H74NO8P	1.76
739.5137	739.5099	[M-H]-	PE-NMe(35:4)/PE-NMe2(34:4)/PE(36:4)/PC(33:4)**	739.5113	C41H74NO8P	1.89
740.5201	740.523	[M-H]-	PE-NMe(35:3)/PE-NMe2(34:3)/PE(36:3)/PC(33:3)	740.5236	C41H76NO8P	0.81
741.5281	741.5246	[M-H]-	PE-NMe(35:3)/PE-NMe2(34:3)/PE(36:3)/PC(33:3)**	741.527	C41H76NO8P	3.24
742.5375	742.5362	[M-H]-	PE-NMe(35:2)/PE-NMe2(34:2)/PE(36:2)/PC(33:2)	742.5392	C41H78NO8P	4.04
743.5432	743.5433	[M-H]-	PE-NMe(35:2)/PE-NMe2(34:2)/PE(36:2)/PC(33:2)**	743.5426	C41H78NO8P	0.94
744.5546	744.5531	[M-H]-	PE-NMe(35:1)/PE-NMe2(34:1)/PE(36:1)/PC(33:1)	744.5549	C41H80NO8P	2.42
745.5023	745.4795	[M-H]-	PA(40:7)	745.4814	C43H71O8P	2.55
745.5593	745.5514	[M-H]-	SM(d35:2-2OH)	745.5501	C40H79N2O8P	1.74
747.5163	747.5197	[M-H]-	PG(34:1)	747.5182	C40H77O10P	2.01
749.531	749.513	[M-H]-	PA(40:5)	749.5127	C43H75O8P	0.40
762.5107	762.5113	[M-H]-	PE-NMe(37:6)/PE-NMe2(36:6)/PE(38:6)	762.5079	C43H74NO8P	4.46
763.5118	763.513	[M-H]-	PE-NMe(37:6)/PE-NMe2(36:6)/PE(38:6)**	763.5113	C43H74NO8P	2.23
764.5222	764.5249	[M-H]-	PE-NMe(37:5)/PE-NMe2(36:5)/PE(38:5)/PC(35:5)	764.5236	C43H76NO8P	1.70
765.5262	765.5244	[M-H]-	PE-NMe(37:5)/PE-NMe2(36:5)/PE(38:5)/PC(35:5)**	765.527	C43H76NO8P	3.40
766.5381	766.5417	[M-H]-	PE-NMe(37:4)/PE-NMe2(36:4)/PE(38:4)/PC(35:4)	766.5392	C43H78NO8P	3.26

771.6455	771.6381	[M-H]-	SM(d39:1)	771.6386	C44H89N2O6P	0.65
776.5052	776.5062	[M-H]-	PE(15:0)/PGF1alpha)/PE(PGF1alpha/15:0)	776.5083	C40H76NO11P	2.70
777.5085	777.5059	[M-H]-	PA(41:5-O)	777.5076	C44H75O9P	2.19
778.5211	778.5161	[M-H]-	3-O-Sulfogalactosylceramide (d34:1)	778.5145	C40H77NO11S	2.06
779.5233	779.5211	[M-H]-	PA(41:5-OH)	779.5232	C44H77O9P	2.69
782.4979	782.4961	[M-H]-	PS(36:4)	782.4978	C42H74NO10P	2.17
786.5251	786.533	[M-H]-	PS(36:2)	786.5291	C42H78NO10P	4.96
787.528	787.5347	[M-H]-	PS(36:2)**	787.5325	C42H78NO10P	0.79
790.5427	790.539	[M-H]-	PE-NMe2(38:6)/PE(40:6)/PC(37:6)	790.5392	C45H78NO8P	0.25
791.5437	791.5428	[M-H]-	PE-NMe2(38:6)/PE(40:6)/PC(37:6)	791.5426	C45H78NO8P	0.25
797.6611	797.6526	[M-H]-	SM(d41:2)	797.6542	C46H91N2O6P	2.01
799.6774	799.6662	[M-H]-	SM(d41:1)	799.6698	C46H93N2O6P	4.50
800.6828	800.6767	[M-H]-	SM(d41:1)**	800.6732	C46H93N2O6P	4.37
807.4975	807.5021	[M-H]-	PI(32:1)	807.5029	C41H77O13P	0.99
808.509	808.5157	[M-H]-	PS(38:5)	808.514	C44H77NO10P	2.10
811.5326	811.5292	[M-H]-	PS(38:4)**	811.5325	C44H78NO10P	4.07
816.5329	816.5184	[M-H]-	PC(38:7-O)	816.5185	C46H76NO9P	0.12
826.5036	826.5064	[M-H]-	PE(42:9+=O)	826.5028	C47H74NO9P	4.36
832.5148	832.5151	[M-H]-	PS(40:7)	832.5134	C46H76NO10P	2.04
833.5215	833.5206	[M-H]-	PI(34:2)	833.5186	C43H79O13P	2.40
834.5275	834.5285	[M-H]-	PS(40:6)	834.5291	C46H78NO10P	0.72
836.5398	836.5425	[M-H]-	PS(40:5)	836.5447	C46H80NO10P	2.63
837.5507	837.5487	[M-H]-	PI(34:0)	837.5499	C43H83O13P	1.43
838.557	838.5572	[M-H]-	PS(40:4)	838.5604	C46H82NO10P	3.82
850.5783	850.5715	[M+Cl]-	PE(40:1-O)	850.5734	C45H86NO9P	2.23
857.5225	857.5165	[M-H]-	PI(36:4)	857.5186	C45H79O13P	2.45
858.5242	858.5261	[M-H]-	PI(36:4)**	858.522	C45H79O13P	4.78
860.5374	860.5428	[M-H]-	PS(42:7)	860.5447	C48H80NO10P	2.21
861.553	861.5499	[M-H]-	PI(36:2)	861.5499	C45H83O13P	0.00
862.556	862.5594	[M-H]-	PS(42:6)	862.5604	C48H82NO10P	1.16
863.5647	863.5616	[M-H]-	PI(36:1)	863.5655	C45H85O13P	4.52
864.5811	864.5758	[M-H]-	PS(42:5)	864.576	C48H84NO10P	0.23
865.486	865.4863	[M-H]-	PG(40:8-3OH)	865.4873	C46H75O13P	1.16
876.6256	876.6264	[M+Cl]-	PE-NMe2(42:2)/PC(40:2)	876.6255	C48H92NO8P	1.03
880.6143	880.6113	[M-H]-	PE(44:5-2OH)	880.6073	C49H88NO10P	4.54

883.5364	883.5303	[M-H]-	PI(38:5)	883.5277	C47H81O13P	2.94
884.5409	884.5414	[M-H]-	PS(44:9)	884.5447	C50H80NO10P	3.73
887.5659	887.5682	[M-H]-	PI(38:3)	887.5655	C47H85O13P	3.04
888.5697	888.5681	[M-H]-	PI(38:3)**	888.5689	C47H85O13P	0.90
890.6429	890.6362	[M-H]-	3-O-Sulfogalactosylceramide (d42:1)	890.6397	C47H87O13P	3.93
891.6471	891.643	[M-H]-	3-O-Sulfogalactosylceramide (d42:1)**	891.6431	C47H87O13P	0.11
892.6249	892.6213	[M-H]-	PC(40:1-O)	892.6204	C48H92NO9P	1.01
893.6281	893.6223	[M-H]-	PC(40:1-O)**	893.6238	C48H92NO9P	1.68
894.6286	894.6255	[M-H]-	PS(44:4)	894.623	C50H90NO10P	2.79
904.6242	904.6245	[M-H]-	PS(42:1-2OH)	904.6284	C48H92NO12P	4.31
909.553	909.553	[M-H]-	PI(40:6)	909.5499	C49H83O13P	3.41
910.557	910.5494	[M-H]-	PI(40:6)**	910.5533	C49H83O13P	4.28
913.5809	913.5843	[M-H]-	PI(40:4)	913.5812	C49H87O13P	3.39
918.6721	918.6767	[M+Cl]-	PE-NMe2(44:2)/PE(46:2)	918.6724	C51H98NO8P	4.68
945.5148	945.5235	[M-H]-	PGP(i-42:5-O)	945.5264	C48H84O14P2	3.07
950.6394	950.6401	[M+Cl]-	PC(46:7)	950.6411	C54H94NO8P	1.05
952.6523	952.6554	[M-H]-	PC(46:6)	952.6568	C54H96NO8P	1.47

* only one representative lipid name was shown for some m/z features

** The second lipid isotope

***ppm calculated from MALDI-FTICR data

Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our [Editorial Policies](#) and the [Editorial Policy Checklist](#).

Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

n/a Confirmed

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| <input type="checkbox"/> | <input checked="" type="checkbox"/> | For null hypothesis testing, the test statistic (e.g. F , t , r) with confidence intervals, effect sizes, degrees of freedom and P value noted
<i>Give P values as exact values whenever suitable.</i> |
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| <input type="checkbox"/> | <input checked="" type="checkbox"/> | Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated |

Our web collection on [statistics for biologists](#) contains articles on many of the points above.

Software and code

Policy information about [availability of computer code](#)

Data collection flexControl (Version 4.0, Bruker Daltonics); ftmsControl (Version 2.1.0, Bruker Daltonics);

Data analysis flexImaging 5.0 (Bruker Daltonics); SCILS Lab 2016b (SCILS, Bruker Daltonics); CaseViewer (version 2.4, 3DHISTECH Ltd); Matlab R2019a; R (version 4.0); Seurat 3.0; Matlab R2019a; Monocle 3; RStudio (version 1.4.1717); pheatmap (version 1.0.12); IsoCorrector (version 1.14.0)
The code used in this study is available in Github (<https://github.com/Gangqiwang/scDYMO>).

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](#) for further information.

Data

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All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our [policy](#)

The exported and processed MSI data for this study were deposited in FigShare at <https://doi.org/10.6084/m9.figshare.20227419.v1>. Due to the large size of all raw data, parts of the raw MSI data are deposited to provide the necessary information of ¹³C-labeled metabolites and spectrum quality. For full availability of raw MALDI-MSI data related to this study, please contact Gangqi Wang (g.wang@lumc.nl) or Bram Heijs (b.p.a.m.heijs@lumc.nl) upon reasonable request data will be made available. Source data are provided with this article.

Human Metabolome Database (<https://hmdb.ca/>) was used for lipid and metabolite annotation.

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	For overall metabolic changes post mortem material of 12-week-old male C57BL/6J mice (n = 3) culled as breeding surplus was used. For bilateral ischemia and reperfusion injury experiments (bIRI) we used 12-week-old male constitutional renin reporter (B6.Ren1cCre/TdTomato/J) mice. 6 mice were divided into 2 groups randomly (n = 3/group). No statistical methods were used to pre-determine sample sizes, but our sample sizes are similar to those reported in previous publications.
Data exclusions	No data was excluded from the analyses.
Replication	All the experiments and data analysis were performed on 3 biological replicates (3 animals per group).
Randomization	For bIRI experiment, 6 mice were divided into 2 groups randomly (n = 3/group).
Blinding	This is not relevant in this study, since there was no group allocation.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

n/a	Involved in the study
<input type="checkbox"/>	<input checked="" type="checkbox"/> Antibodies
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<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology and archaeology
<input type="checkbox"/>	<input checked="" type="checkbox"/> Animals and other organisms
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<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data
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Methods

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<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging

Antibodies

Antibodies used	Monoclonal Rat IgG2B anti-mouse KIM1 antibody (5 µg/ml, R&D Systems, Abingdon, UK, MAB1817), Rabbit monoclonal anti-VCAM1 antibody (1:250, abcam, ab134047); Mouse IgG2a anti-CDH1 antibody (1:300, BD Biosciences, 610181); Polyclonal Goat IgG anti-NPHS1 (2 µg/mL, R&D Systems, AF3159); Rat IgG2a anti-mouse pan-endothelial cell antigen (2 µg/mL, MECA32, BD Biosciences, 553849); donkey anti rat IgG AF488 (1:300, Invitrogen, A21208); donkey anti rabbit IgG AF647 (1:300, Invitrogen, A31573); donkey anti mouse IgG AF488 (1:300, Invitrogen, A21202); donkey anti sheep IgG AF568 (1:300, Invitrogen, A21099);
Validation	anti-mouse KIM1 antibody (R&D Systems, Abingdon, UK, MAB1817): source: Monoclonal Rat IgG2B Clone # 222414; Species Reactivity: mouse; Specificity: Detects mouse TIM 1/KIM-1/HAVCR in ELISAs and Western blots. This antibody does not cross-react with recombinant human (rh) TIM 1, rmTIM-2, or rmTIM-3; validation statements, relevant citation and other information can be found from the manufacturer's website: https://www.rndsystems.com/products/mouse-tim-1-kim-1-havcr-antibody-222414_mab1817 anti-VCAM1 antibody (abcam, ab134047): source: Rabbit monoclonal [EPR5047]; Species Reactivity: Mouse, Rat, Human; Application: WB, IP, IHC-P, Flow Cyt (Intra), ICC/IF, ELISA; validation statements, relevant citation and other information can be found from the manufacturer's website: https://www.abcam.com/VCAM1-antibody-EPR5047-ab134047.html?gclid=CjwKCAjwryUBhBSEiwAGN5OCK-ODP6CXh8uHi9qleNoHz46ftMFojX_C3Y3VRXYgTwehAZ76kSY8RoC3yoQAvD_BwE anti-CDH1 antibody (BD Biosciences, 610181): source: Mouse IgG2a, κ; species reactivity: Human (QC Testing), Mouse, Rat, Dog (Tested in Development); Application: Western blot (Routinely Tested), Immunofluorescence, Immunohistochemistry, Immunoprecipitation (Tested During Development); validation statements, relevant citation and other information can be found from the manufacturer's website: https://www.bdbiosciences.com/en-us/products/reagents/microscopy-imaging-reagents/immunofluorescence-reagents/purified-mouse-anti-e-cadherin.610181 anti-NPHS1 (R&D Systems, Abingdon, UK, AF3159): source: Polyclonal Goat IgG; species reactivity: Mouse; Application: Immunohistochemistry; validation statements, relevant citation and other information can be found from the manufacturer's

website: https://www.rndsystems.com/products/mouse-nephrin-antibody_af3159.
 anti-mouse pan-endothelial cell antigen (MECA32, BD Biosciences, 553849): source: Rat IgG2a, κ ; species reactivity: Mouse; Application: Flow cytometry (Routinely Tested), Immunohistochemistry-frozen, Immunoprecipitation, Western blot (Reported); validation statements, relevant citation and other information can be found from the manufacturer's website: <https://www.bdbiosciences.com/en-us/products/reagents/western-blotting-and-molecular-reagents/western-blot-reagents/purified-rat-anti-mouse-panendothelial-cell-antigen.553849>.
 donkey anti rat IgG AF488 (Invitrogen, A21208): <https://www.thermofisher.com/antibody/product/Donkey-anti-Rat-IgG-H-L-Highly-Cross-Adsorbed-Secondary-Antibody-Polyclonal/A-21208>
 donkey anti rabbit IgG AF647 (Invitrogen, A31573): <https://www.thermofisher.com/antibody/product/Donkey-anti-Rabbit-IgG-H-L-Highly-Cross-Adsorbed-Secondary-Antibody-Polyclonal/A-31573>
 donkey anti mouse IgG AF488 (Invitrogen, A21202): <https://www.thermofisher.com/antibody/product/Donkey-anti-Mouse-IgG-H-L-Highly-Cross-Adsorbed-Secondary-Antibody-Polyclonal/A-21202>
 donkey anti sheep IgG AF568 (Invitrogen, A21099): <https://www.thermofisher.com/antibody/product/Donkey-anti-Sheep-IgG-H-L-Highly-Cross-Adsorbed-Secondary-Antibody-Polyclonal/A-21099>

Animals and other organisms

Policy information about [studies involving animals](#); [ARRIVE guidelines](#) recommended for reporting animal research

Laboratory animals	12-week-old male C57BL/6J mice and renin reporter (B6.Ren1cCre/TdTomato/J) mice.
Wild animals	the study did not involve wild animals
Field-collected samples	the study did not involve samples collected from the field
Ethics oversight	We used post mortem material of 12-week-old male C57BL/6J mice culled as breeding surplus. Mice were kept and cared for in accordance with the Experiments on Animals Act (Wod, revision 2014, The Netherlands) and EU directive no.2010/63/EU. For bIRI study, animal experiments were approved by the Ethical Committee on Animal Care and Experimentation of the Leiden University Medical Center (permit no. AVD1160020171145).

Note that full information on the approval of the study protocol must also be provided in the manuscript.