Supporting Information

Coupling microdroplet-based sample preparation, multiplexed isobaric labeling, and nanoflow peptide fractionation for deep proteome profiling of tissue microenvironment

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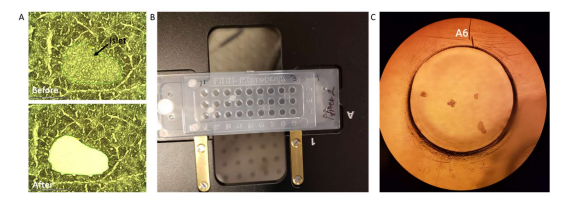
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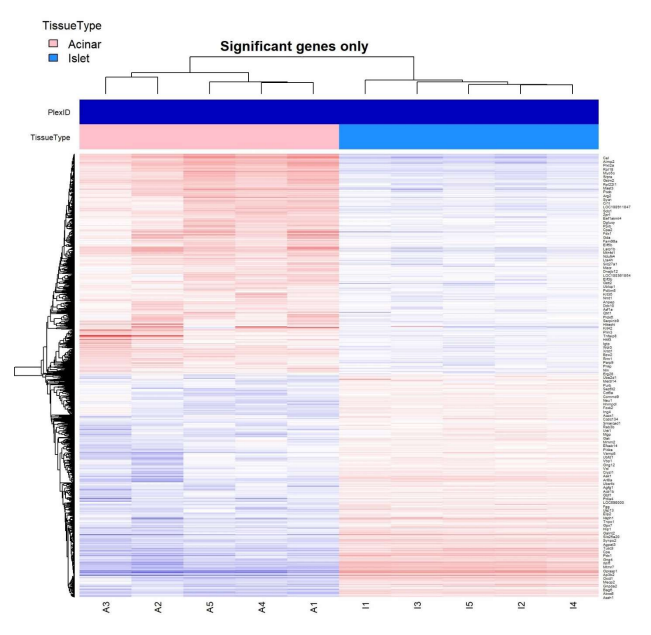
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Supporting figure 1. Images of the dissection and collection of pancreas tissue voxels into the microPOTS chip.

Supporting Figure 2: The heat map visualization of distinct cluster of all significant genes, indicating different biological functions of the two tissue types (islet and acinar).



Supporting figure 1. Images of the dissection and collection of pancreas tissue voxels into the microPOTS chip. A) Islet region before and after laser-microdissection B) Microchip with collected pancreas tissue samples. Microwells were preloaded with DMSO that served as a capturing medium. C) Islet tissue voxels collected into microwell A6, observed under the Zeiss LCM microscope.



Supporting Figure 2: The heat map visualization of distinct cluster of all significant genes, indicating different biological functions of the two tissue types (islet and acinar).