



INTRODUCTION

Despite the progress of optical imaging techniques to improve spatial resolution and detection accuracy for biological applications, the number of markers that can be visualized and analyzed simultaneously has largely been restricted by the limited range of the visible light spectrum. Here we present a novel multiparametric imaging technology, termed CODEX (CO-Detection by IndEXing) that combines highthroughput and high-content methodologies to label dozens of biomarkers simultaneously in a single sample, and detect and resolve their relative expression, abundance, and spatial relationships.



INSTRUMENT / MICROSCOPE INTEGRATION



Figure 3. Seamless microscope integration. The CODEX fluidics device integrates into microscope stages through a custom stage insert. The CODEX Driver Software is compatible with multiple microscope brands/types, including Keyence BZ-X710/800, Leica DMi8 & Zeiss Axio-Observer.

Multiparametric Proteomic Profiling Via Imaging Dozens of Biomarkers Simultaneously

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and deep analysis of many different cell types and their interactions, including B cells (CD19, CD20), T-cells (CD3), Helper T cells (CD4), Cytotoxic T cells (CD8), Dendritic Cells (CD123), Plasma Cells (CD38), NK cells (CD56), Granulocytes (CD15), and many more.

CD34 CD4 E-Cadherin **CD3** Vimentin Ki67 **CD20** cluster analysis, which identified 10 distinct clusters corresponding to 10 different cellular phenotypes. HOECHST3 vs HOECHST4 Profile_Homogeneity vs size 12464.24 11331.13 10198.03 7931.79 0 6798.6 5665. 4532.4 3399.3 2266.23 1133.11-Profile_Homogeneity HOECHST3 identification of cytotoxic T-cells (CD3+, CD8+), from which cell counts were determined Abundance of each cell type was counted relative to the maximum abundance of each cell type across bins. HelperT cel Cytotoxic T cells Proliferating cells Cytotoxic T cells

Figure 10. Cell-cell interaction neighborhood analysis. Cell phenotype specific clusters were subjected to cell-cell interaction analysis. Color in heatmap indicates intensity of log-odds ratio of interaction (frequency of cell-cell interactions relative expected frequency of random interactions). One interaction is defined as a less than 7 um distance between cells.

DEEP MULTIPARAMETRIC ANALYSIS OF MELANOMA SAMPLE

Figure 7. Cluster analysis of image data. Image data was subjected to segmentation to distinguish single cells and generate FCS data. This was subsequently analyzed via decision tree based

Figure 8. Cell feature based gating. Quantitative FCS data was gated by various cell features, including size, homogeneity, and specific biomarkers. Sequential gating allowed for accurate

Figure 9. Spatial analysis of immune cells. Relative density of three immune cell types were analyzed across the red rectangular region. The rectangular region was broken up into 25 bins.

