



Identification of the novel immune synapse-localized proteome for immuno-oncology using Microscop™-induced targeted photo-biotinylation



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Abstract

Although numerous immune synapse (IS) protein species have been identified, many IS-localized protein species remain unknown. Understanding the proteome of the IS between a target cell and a lymphocyte is crucial for advancing immuno-oncology. However, the low abundance of ISs and the absence of a definitive enrichment marker have hindered efficient proteomic profiling. In this study, we utilized Microscop™, an innovative system that integrates microscopy, machine learning, and photochemical labeling to enable precise and spatially specific enrichment of IS proteins, thereby facilitating proteomic discovery for the IS. We employed Raji B cells as antigen-presenting cells (APCs) and induced IS formation with Jurkat T cells. The system first employed immunofluorescence imaging of CD3, a common IS marker in Jurkat T cells, and utilized a convolutional neural network-based deep learning algorithm to recognize IS formation from CMTX-stained Raji B cells. Our automated system successfully achieved spatially targeted biotin-tagging of proteins at ISs through multiple rounds of imaging, deep learning-driven pattern generation, and photochemical labeling. Subsequent streptavidin pull-down and mass spectrometry analysis enabled the identification of IS-specific proteins. Remarkably, our spatial proteomic approach led to the isolation and identification of hundreds of different species at the IS interface, including proteins associated with key components of T-cell receptor (TCR) signaling pathways such as the TCR/CD3 complex, Src and Tec family tyrosine kinases, and pivotal NF- κ B signaling proteins. Additionally, we identified a significant enrichment of proteins not previously associated with the IS. Our study not only illuminates previously unknown aspects of immune regulation at the IS interface but also has significant implications for cancer research, particularly in understanding and manipulating immune responses for therapeutic purposes.

Microscop™: Ultrahigh-content microscopy-guided photo-biotinylation platform

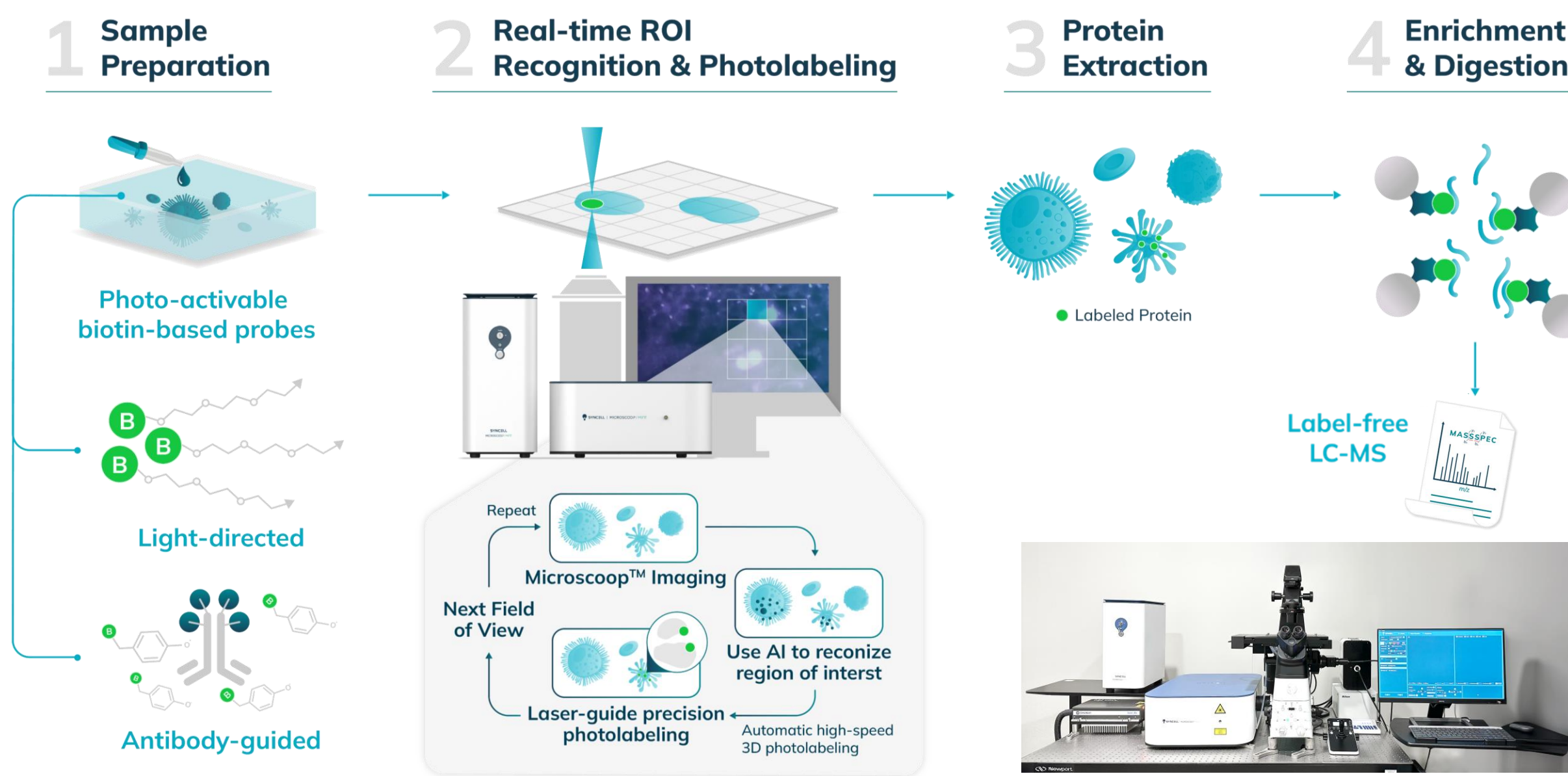


Fig. 1 | Schematic workflow of SYNCELL Microscop™. A total-syn ultra-content microscopic platform that integrates image acquisition, photochemistry, microscopy, optics, and FPGA-based mechatronics enable high-content in situ photolabeling followed by mass spectrometry analysis.

Synchronized system for high-speed and accurate image masking

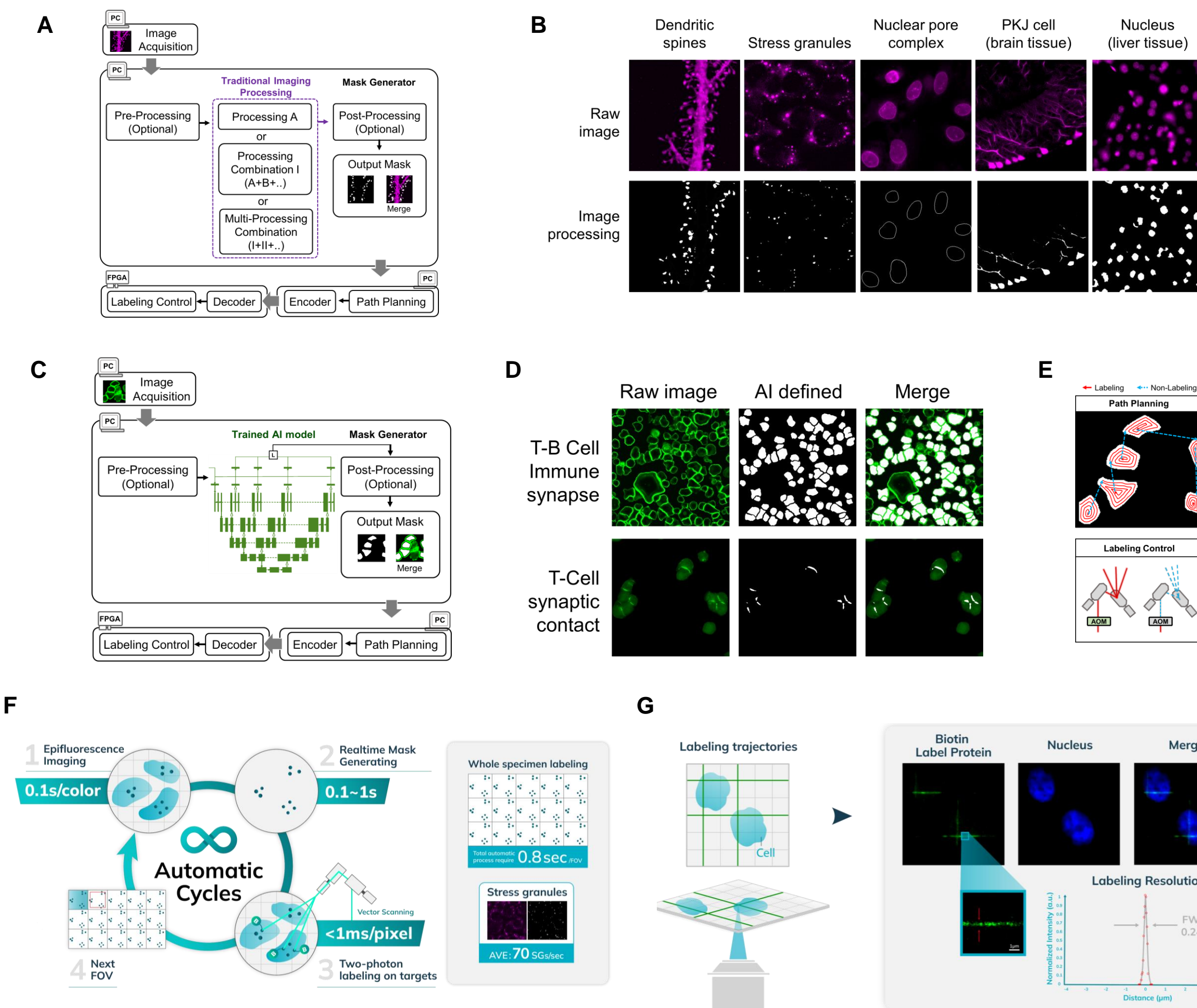


Fig. 2 | A-B. Image processing is applied to recognize the regions of interest of acquired images. C-D. Deep learning image segmentation and recognition of complex models from raw images. E. Algorithm generates the labeling path and the non-labeling path of an input mask, and labeling control of the galvanometer system and the AOM. F. Workflow for ultrahigh-content targeted photo-biotinylation includes: (1) identifying and acquiring images of regions of interest by light microscope; (2) generating real-time patterns of ROIs; (3) illuminating the selected region within ROIs for photo-biotinylation; (4) moving the stage to the next FOV; and repeating steps 1-4 for each FOV until all FOVs have been processed. G. Resolution of photo-biotinylation. A line "cross" pattern is photolabeled on fixed U-2OS cells, and the biotinylated molecules are shown in green. DAPI: Blue, scale bar: 10 μ m. 40x/0.95 NA objective.

Photo-induced spatial biotinylation in subcellular compartments at a nanoscale resolution and discovery of novel proteins

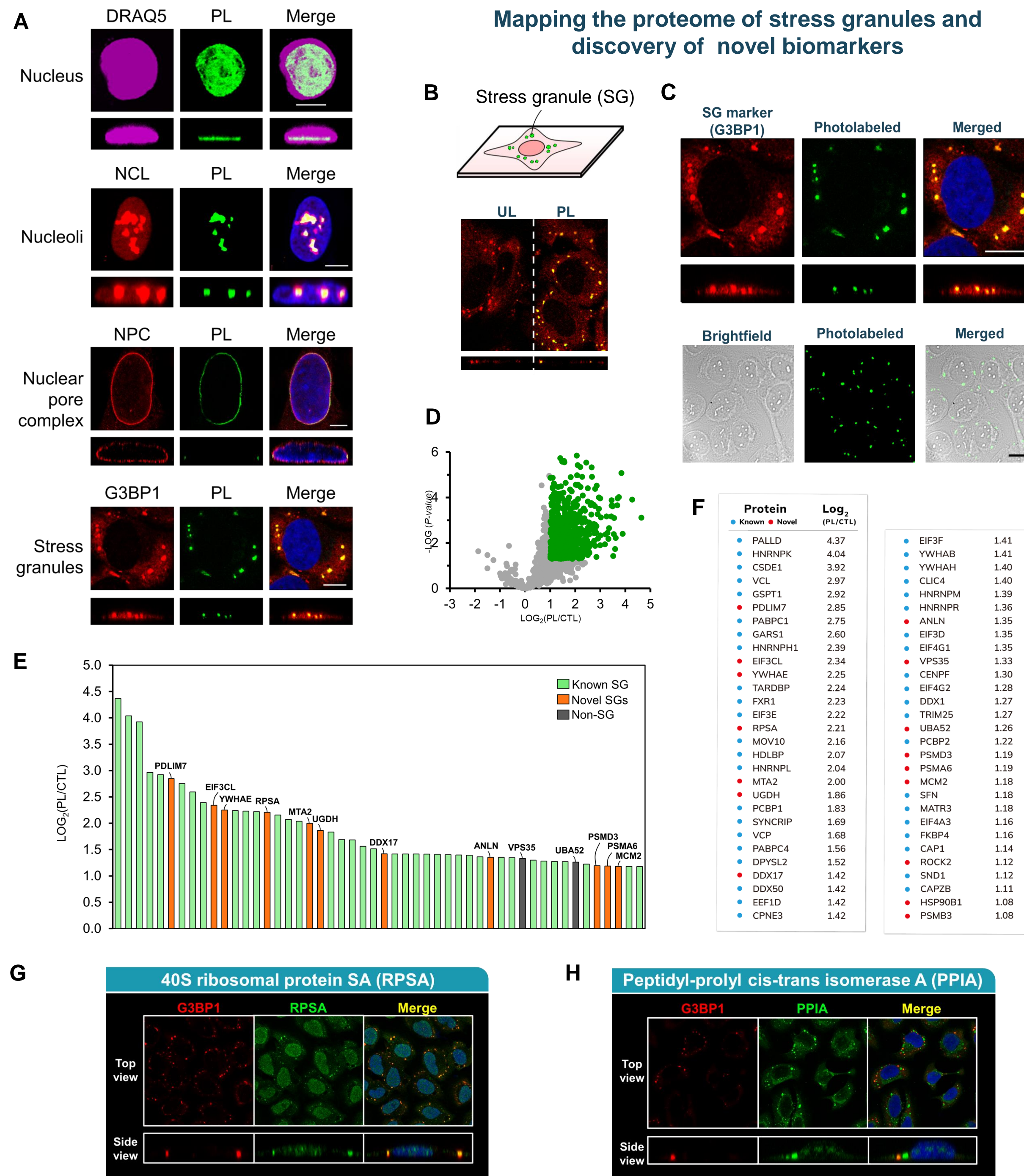


Fig. 3 | A. Top- (xy) and side- (yz) views of photolabeled subcellular compartments. B. Confocal micrographs of photolabeled (PL) and non-photolabeled (UL) at user defined stress granules (SG). C. Confocal micrographs depicting precise and accurate photolabeled SG at top- and side-view. D. Volcano plot of relative protein levels in photolabeled samples to control samples (PL/CTL ratio) in log₂ scale. Over-represented (enriched) proteins are shown in green. E. 74% of true positive SGs are found in the top 50 proteins ranked by PL/CTL ratio. Novel SG proteins (orange) and known SG proteins (green) from the top 50 ranked proteins. F. List of novel SG proteins (red) discovered by Microscop™. G-H. Confocal micrographs depicting SG formation of potential stress granule proteins in U-2OS cells with or without arsenite stress. Twelve potential SG proteins (green) are highly co-localized with G3BP1 SG markers. Lens: 63x oil. PL: photolabeled (biotinylated) proteins stained with NeutrAvidin-488, Red: G3BP1, Blue: DAPI. Scale bar: 10 μ m.

Spatial proteomics of primary cilia: known and putative novel primary cilia proteins and their functional insights

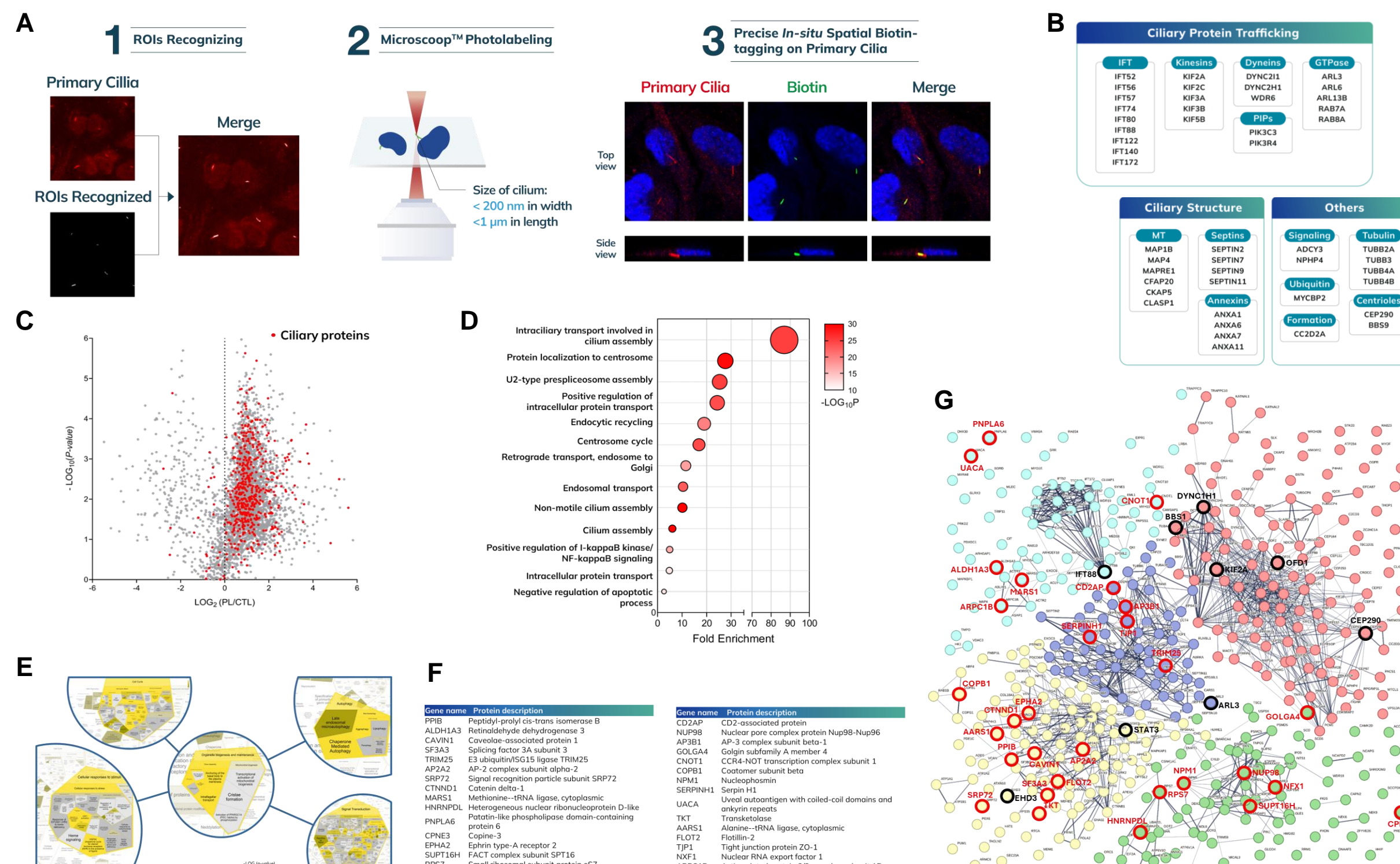


Fig. 4 | A. Images of primary cilia are processed by filtering and segmentation using traditional image processing (left). Confocal micrographs depicting precise and accurate photo-biotinylated primary cilia at lateral (xy)- and axial (z) directions (right). Red: GT335, Green: NeutrAvidin-488, Blue: DAPI. B. List of a few well-known ciliary proteins identified by Microscop™. C. A distribution of overall protein abundances by the ratio of photolabeled (PL) sample to those in a control (CTL) sample annotated as PL/CTL ratio. Ciliary proteins (red) are enriched in the PL group compared to the CTL sample. D. The top 100 enriched proteins were subjected to Gene ontology to reveal cilia related biological process. E. 427 enriched ciliary proteins were subjected to Reactome to reveal cilia related pathways. F. The list of the top 30 non-ciliary proteins (putative ciliary proteins) enriched by Microscop™. G. The 30 putative ciliary protein and 427 enriched ciliary proteins were subjected to STRING to reveal protein-protein interaction networks, where the 30 putative ciliary proteins (F) are indicated in red.

Mapping the immune synapse-localized proteome and immuno-oncologic protein interactions

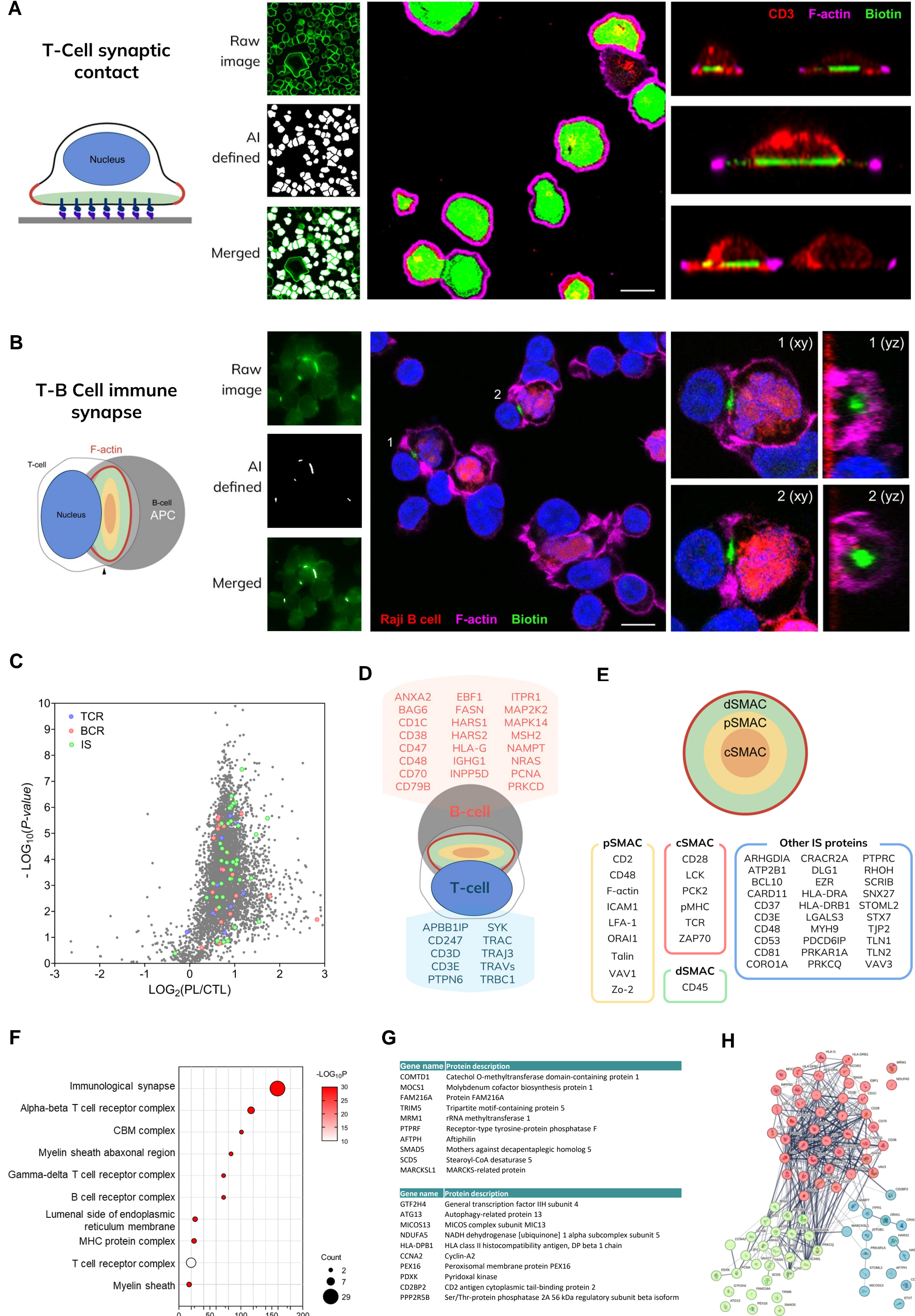


Fig. 5 | A. T-cell synaptic contact model (left). Machine learning recognized the region of IS in the spreading assay shown in white (center). IS proteins in the spreading assay were photo-labeled (biotin): green, indicating a precise labeling on IS. The side view of labeling region is co-localized with CD3, a well-known marker of IS on the bottom of cells (right). B. T-B cell conjugation model (left). Machine learning indicates the region of IS in the T-B cell conjugation shown in white (center). AI-guided the recognition of IS in the T-B conjugation, and proteins on ISs were photo-labeled shown in green, localized within the actin ring, indicating the precise labeling on IS. C. Distribution of overall protein abundance according to the photolabeled/control (PL/CTL) ratio for immune synapse proteome study; TCR: T-cell receptor, BCR: B-cell receptor, IS: Immune synapse. D. The TCR and BCR proteins identified by SYNCELL Microscop™. E. The classical immunological synapse clusters (cSMAC, pSMAC, IS) and other immune synapse proteins identified by SYNCELL Microscop™. F. 69 enriched immunological proteins were subjected to GeneOntology to reveal compartments. G. The list of the top 20 non-immune synapse proteins (putative immune synapse proteins) enriched by Microscop™. H. The 20 putative immune synapse proteins and 69 enriched immunological proteins were subjected to STRING to reveal protein-protein interaction networks. Scale bar=10 μ m.

References:
Microscopy-guided subcellular proteomic discovery by high-speed ultra-content photo-biotinylation. Chen et al. bioRxiv 2023.12.27.573388.

Summary

- SYNCELL Microscop™: A novel platform for hypothesis-free subcellular spatial proteomic discovery
- Integration of Technologies: Combines microscopy, deep learning, photochemistry, two-photon illumination, and mechatronics
- Purpose: Facilitates high-content, image-guided photo-labeling at a nanoscale resolution
- Capabilities: Precisely labels spatially specific proteins from hundreds of thousands of individual cells, suitable for mass spectrometry analysis
- Achievements:
 - Discovered dozen of novel stress granule proteins and identified 608 known ciliary proteins, providing functional insights and listing putative proteins with high protein-protein interactions
 - Precision photo-biotinylation of millions of immune synapses, followed by LC-MS/MS analysis, resulted in the identification of a highly enriched set of immune synapse proteins. Notably, this approach also revealed potentially novel proteins with strong correlations to immunological functions.