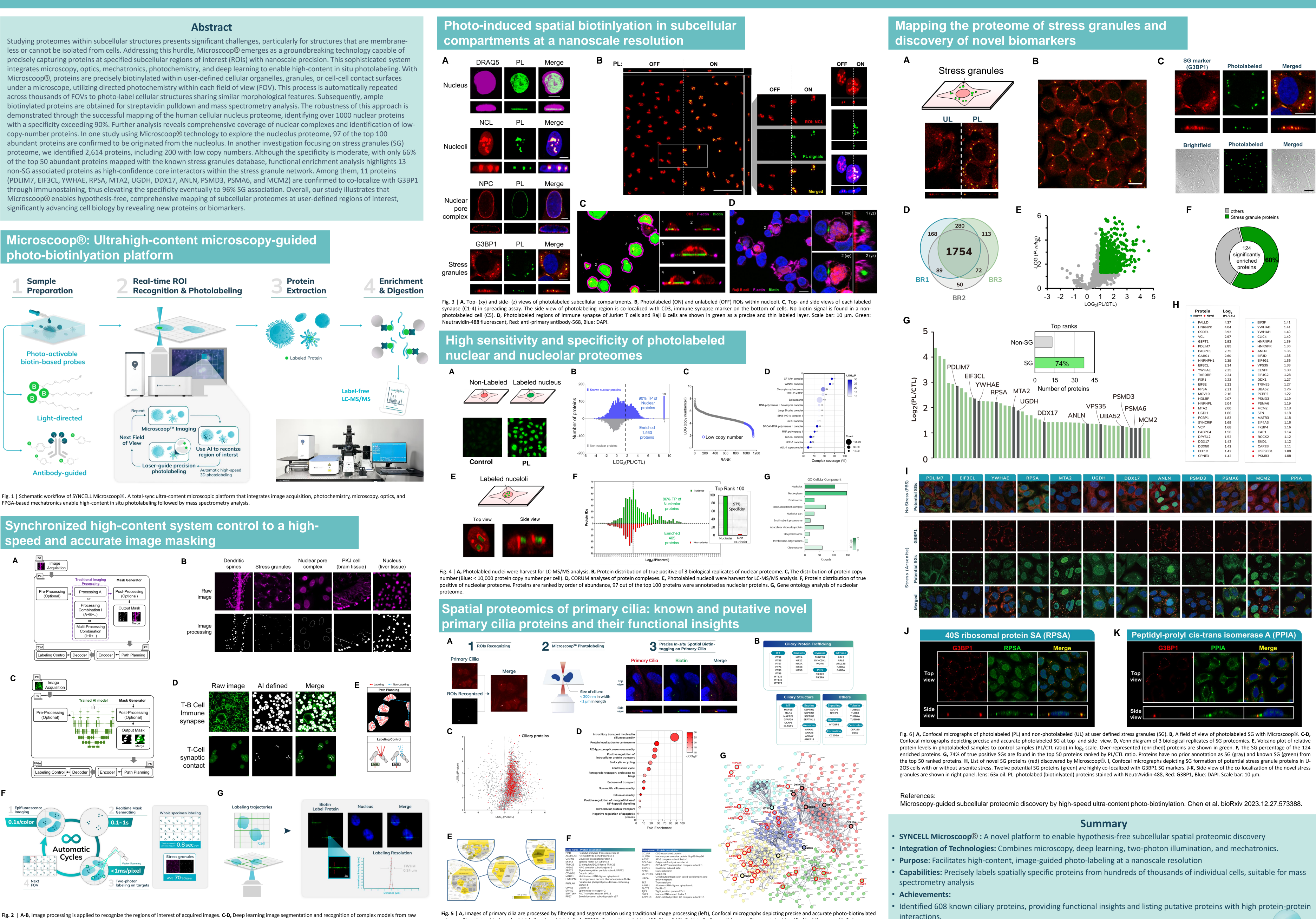
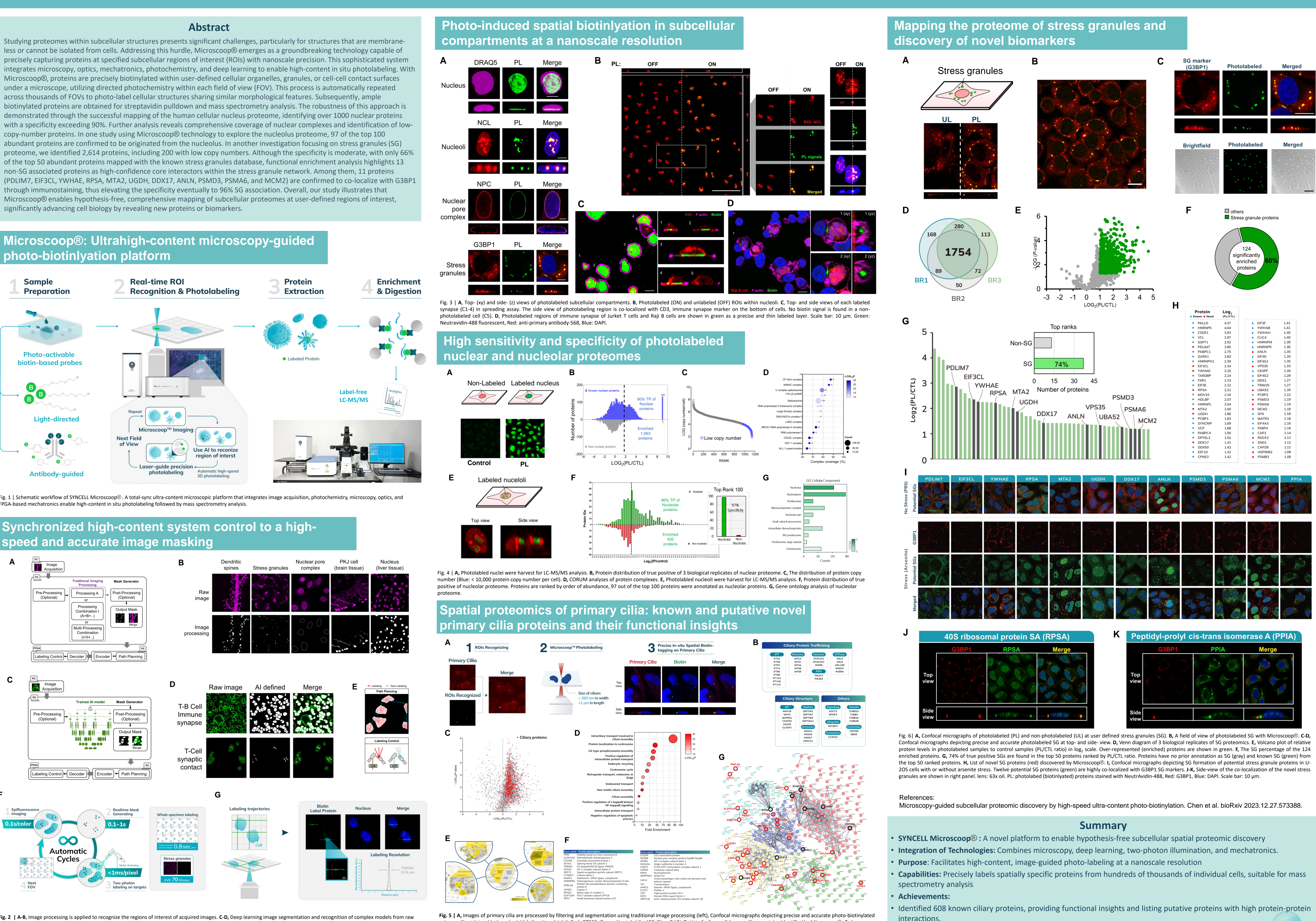


Total-sync ultra-content microscopic opto-biotinylation enables high-sensitivity hypothesis-free subcellular protein discovery

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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 ultrahighcontent targeted pohoto-biotinylation includes: (1) identifying and acquiring images of regions of interest by light microscope; (2) generating realtime patterns of ROIs; (3) illuminating the selected region within ROIs for protein 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 µm. 40x/0.95 NA objective.

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 Microscoop[®]. 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 Microscoop[®]. 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.

- interactions.
- Discovered dozen of novel stress granule proteins with high precision and accuracy



EIF3F	1.41	
(WHAB	1.41	
(WHAH	1.40	
CLIC4	1.40	
INRNPM	1.39	
INRNPR	1.36	
ANLN	1.35	
EIF3D	1.35	
EIF4G1	1.35	
/PS35	1.33	
CENPF	1.30	
EIF4G2	1.28	
DDX1	1.27	
rrim25	1.27	
JBA52	1.26	
PCBP2	1.22	
PSMD3	1.19	
PSMA6	1.19	
MCM2	1.18	
SFN	1.18	
MATR3	1.18	
EIF4A3	1.16	
FKBP4	1.16	
CAP1	1.14	
ROCK2	1.12	
SND1	1.12	
CAPZB	1.11	
HSP90B1	1.08	
PSMB3	1.08	
PSMB3	1.08	