Shared Nearest Neighbors Approach and Interactive Browser for Network Analysis of a Comprehensive Non–Small-Cell Lung Cancer Data Set

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* Equal contribution

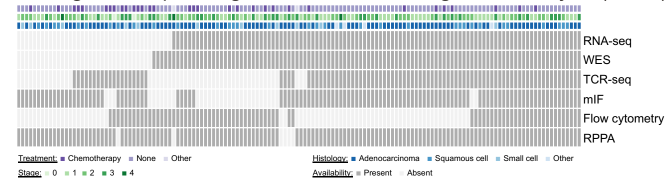
Schmidt, et al. JCO Clinical Cancer Informatics no. 6 (2022) e2200040.

Presenter (STS) has no disclosures.

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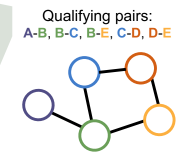
Immunogenomi**C** pr**O**filing of **N**on-small cell lung cancer Project (ICON)



Shared Nearest Neighbors (SNN) approach for multi-platform networks

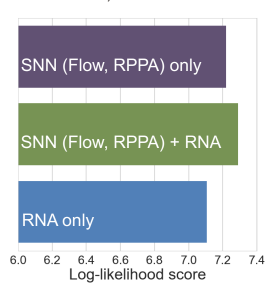
Rank	Gene A	Gene B	:		
1	Protein 1	Protein 2			
2	Protein 2	Protein 4			
3	Protein 3	Protein 1			
4	Protein 4	Protein 6			
5	Protein 5	Protein 5			

(3.11.1) 5.66.61.10.								
	1	8.0	0.6	0.2	0.4			
	0.8	1	0.8	0.4	0.8			
	0.6	0.8	1	1	0.4			
	0.2	0.4	1	1	0.8			
	0.4	8.0	0.4	8.0	1			
	٨	R	C	ח	E			

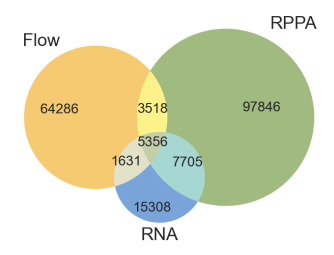


ICON data network highlights new connections based on interplay between integrated platforms

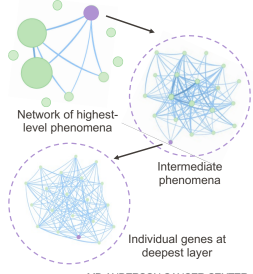
Selection of SNN with RNA-based edges Improved performance over SNN, RNA alone



SNN platforms provide more edges per node Overlaps in network edges from each platform shown



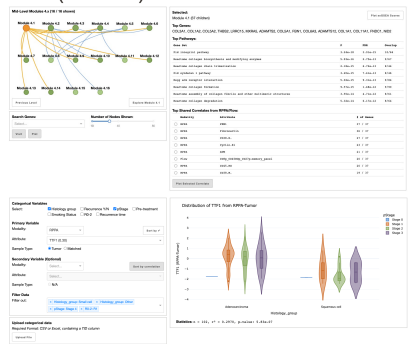
Community detection identifies modules
10 top-level and 93 mid-level modules with 10+ genes



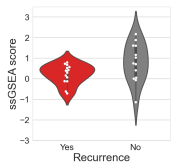
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ICON data browser enables interactive network exploration for insights into tumor characteristics of interest

Network panel (top) and plotting/modeling panel (bottom) of the ICON data browser



Score distributions for highlighted modules relevant to selected features



Recurrence in Stage 2+, non-LUSC tumors

TP53 oncogenotype in LUSC tumors



To learn more, please see our recent publication in JCO CCI.

Ongoing: Expansion of approach to NEOSTAR to enable multi-cohort, multi-platform integration

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