A tutorial for using OmniPath App in Cytoscape

supplementary material for

F. Ceccarelli, D. Turei, A. Gabor and J. Saez-Rodriguez

Bringing data from curated pathway resources to Cytoscape with OmniPath







Institute for Computational Biomedicine Heidelberg University & RWTH Aachen

Case study 2: Training network model to data



Intro

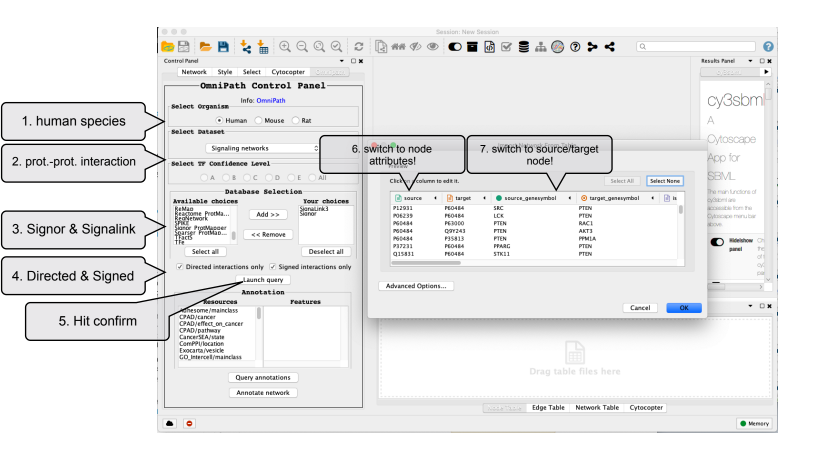
- Here we combine Omnipath App with PathLinker and Cytocopter
- Problem statement:
 - we want to investigate the signal propagation in the MAP-kinase pathway upon treatment with EGF and TNFa stimulation and PI3K and RAF1 inhibition
 - We have measurements for the following nodes:
 - RAF1, MAPK3, JUN, GSK3b, MAPK14, NFkB1
- Pipeline:
 - we query signaling interactions from Omnipath App
 - select a subnetwork by finding shortest path between the perturbed node and measured phosphoproteins. We use PathLinker for this job
 - Turn the network into a Boolean logic model and train the model to the data using Cytocopter.

Requirements

- Install
 - Omnipath App (<u>https://apps.cytoscape.org/apps/omnipath</u>)
 - Cytocoptr (<u>https://apps.cytoscape.org/apps/cytocopter</u>)
 - PathLinker (https://apps.cytoscape.org/apps/pathlinker)
- user data is provided on Github
 - (<u>https://github.com/saezlab/Omnipath_Cytoscape/tree/master/tutorial</u>)

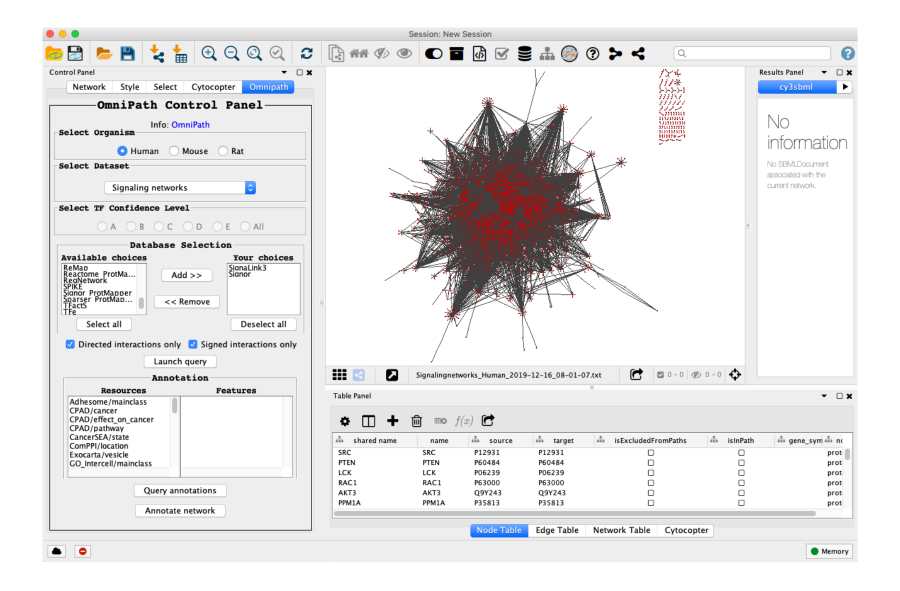


Import signaling network from Omnipath



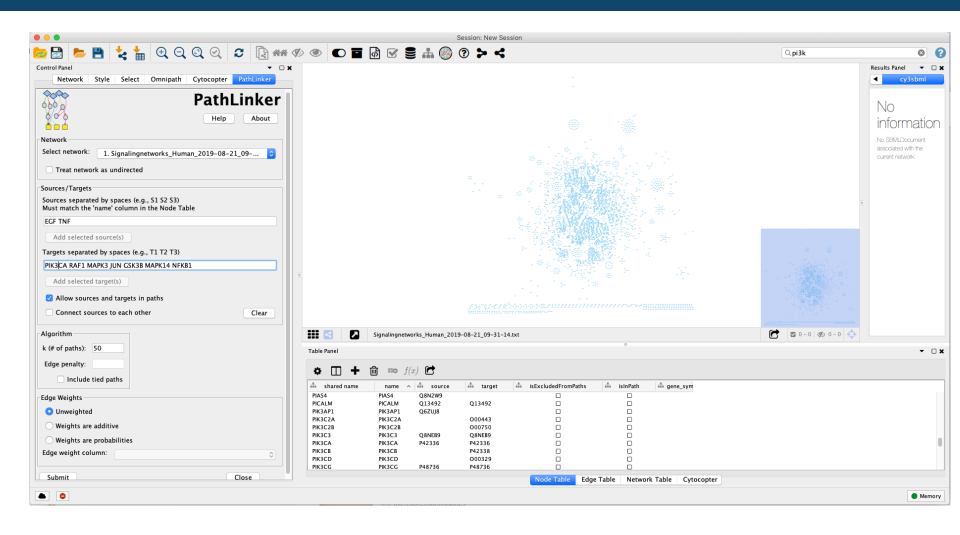


Imported network with organic layout





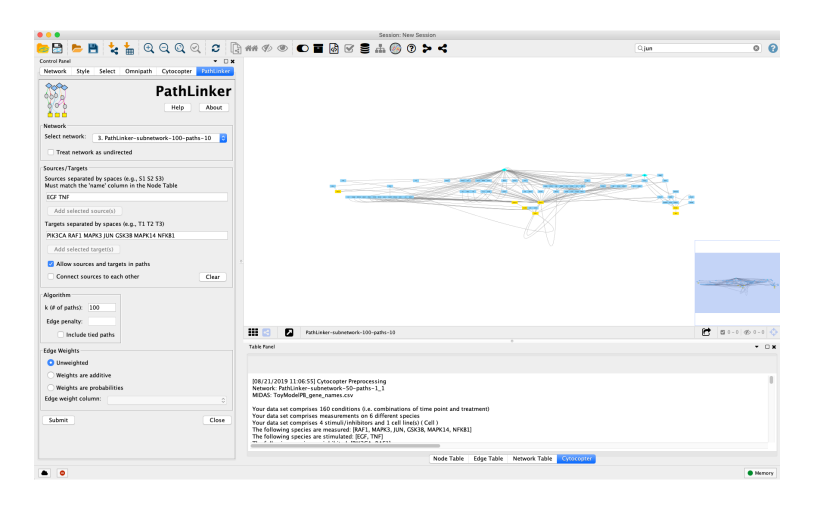
PathLinker: find relevant subnetwork



Sources: EGF and TNFa stimulation nodes Targets: measred proteins



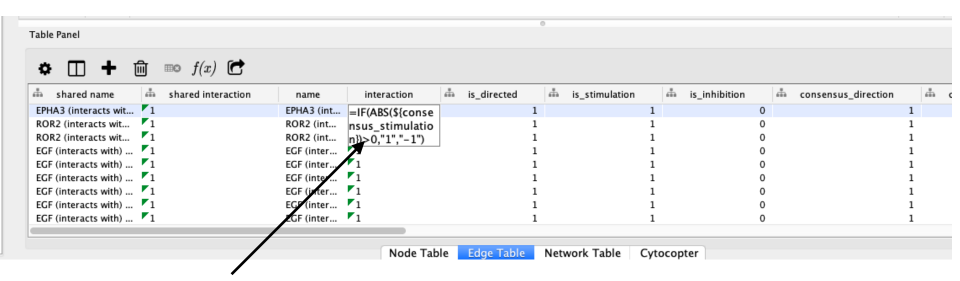
Results of PathLinker from Omnipath



This network is translated to a Boolean model in the next step using Cytocopter.



Define interaction type for the Boolean model



Cytocopter requires that the sign of each edge is defined in the edge table / interaction column:

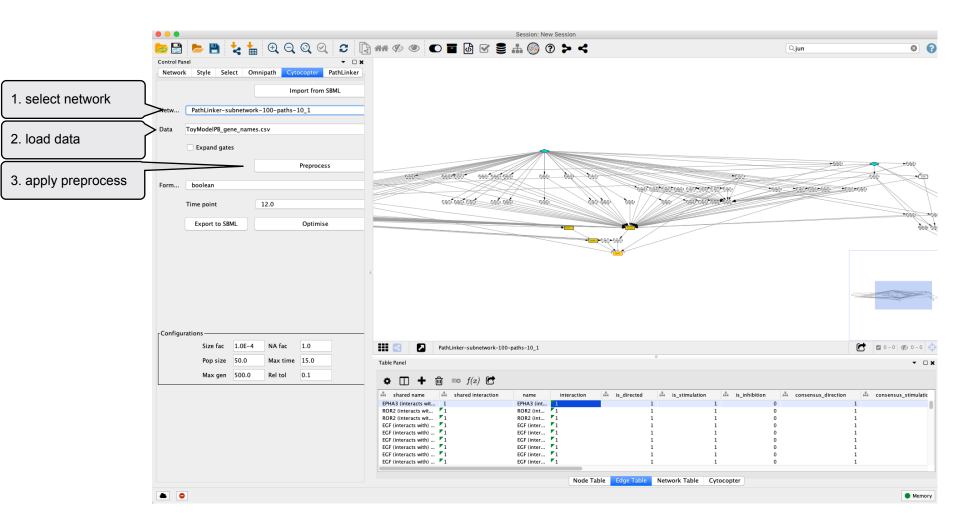
- →1 for activation
- ⇒-1 for inhibition

Therefore, interaction field has to be updated,

Then right click on the cell and select "apply to the entire column".

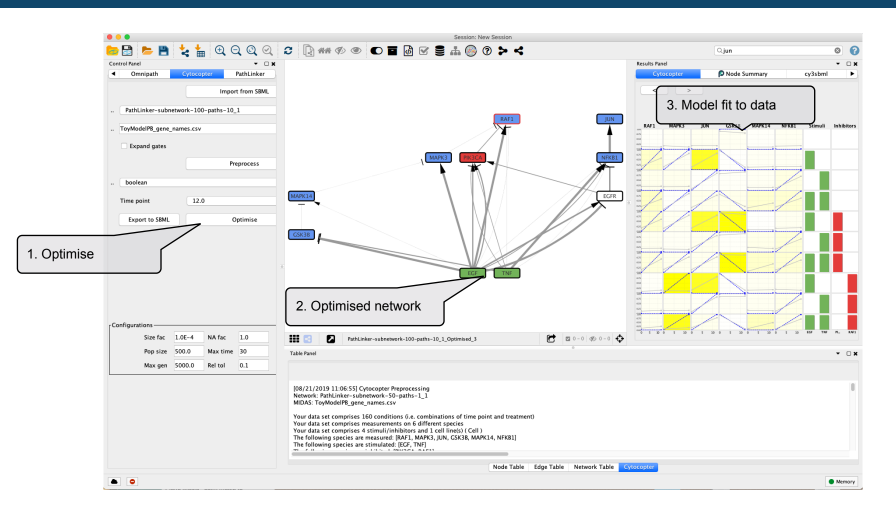


Setup the Boolean model in Cytocopter





Build model and optimise the network



Optimisation finds a set of models that fits the data equally good. The strength of the edges is proportional to the number of models where the edge present. Light edge shows unlikely interactions.