Combining bottom up and top down approaches for Drug-Target-Interaction prediction

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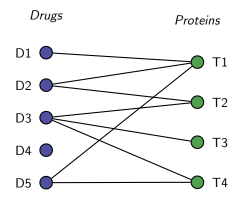
Outline

1. Problem Description

- 2. Recent approaches
- 3. A medium approach to DTI prediction Filtering possible targets

Problem Description

Prediction over bipartite graph:



Classification of recent approaches¹²

	Drugs	Protein
bottom-up	▶ GCN over molecules▶ drug similarity	 secondary structure prediction contact prediction convolution over amino acid sequences
top-down	network approachesdrug similarity	▶ protein similarity

¹Chen Wang et al., Briefings in Bioinformatics, 2018

²Yu Ding et al., Briefings in Bioinformatics, 2019

Problems of recent approaches

Main issues:

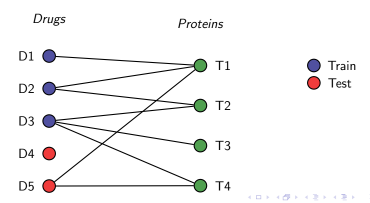
- Lack ability to generalize or are unable to spot small differences
- Usually only top-down or bottom-up
- Not making use of interaction networks



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How to bring both approaches together?

Algorithm

- For each drug, filter possible targets according to protein based measurement
- 2. Bring those together with the interaction networks for the prediction

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Two hypotheses to be tested:

- Interaction only network sufficient?
- Does filter method increase performance?

Filtering possible targets

How to build such a filter?

STITCH database got

- ▶ 390 000 chemicals, and
- ▶ 3.6 million proteins
- ► from over 2000 organisms



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For each drug, find a motif in the target amino acid sequences and query that against all possible proteins and use the results as features for the prediction



How to find motifs?

- 1. Build multi sequence alignment with alignment tool of choice over non-human proteins
 - FAMSA, MAFFT, Kalign, MSA Props, Muscle, ...
- 2. Build representation of that alignment
 - Hidden Markov model with HMMER
 - PWM
- 3. Query representation against database



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This is quite expensive to compute ... (8000 hours for alignments of 614 drugs + IBEX queue times for such big jobs)



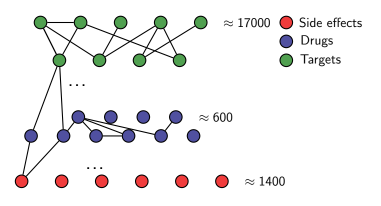
Other features for the model

- Protein-protein-interaction network from STRING (only human)
- ▶ Drug-drug-interaction network from Boyce³ and Drugbank
- Side-effect data for drugs from SIDER, annotated with MedDRA hierarchy (semantic similarity)



³Boyce et al., 2015

Features in context



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How to learn a sufficient embedding?

- Use Filtered results as node features
- Run different models for classification of proteins/nodes in the graph:
 - GraphSage
 - HinSage
 - Attri2Vec
 - Graph Attention Network (GAT)
 - ► Graph Convolutional Network (GCN) for node classification
 - GCNs for graph classification of induced subgraph
 - SimpleGCN
 - Personalized Propagation of Neural Predictions (PPNP)
 - Approximate Personalized Propagation of Neural Predictions (APPNP)



Filtering possible targets

First results

Used data			layer sizes	model	Acc	AUC	
PPI	DDI	SE	HMM				
1	1	0	0	641	graphsage	88.97	66.20
1	1	0	0	641	graphsage	80.28	70.51
1	1	0	0	641	graphsage	87.44	67.80
1	1	0	0	641	graphsage	86.46	68.39
1	1	0	0	641	graphsage	88.84	66.62
1	1	1	0	641	graphsage	93.34	62.42
1	1	1	0	641	graphsage	73.20	62.53
1	1	1	0	641	graphsage	93.27	64.81
1	1	1	0	641	graphsage	74.28	59.18
1	1	1	0	641	graphsage	72.40	63.93
1	1	1	0	641	gcn	88.61	67.27
1	1	1	0	641	gat	82.65	71.69
1	1	1	0	641	hinsage	83.18	70.33
1	1	1	0	641	ppnp	89.80	67.78
1	1	1	0	641	appnp	84.35	71.82
1	1	1	0	641	sgc	86.50	68.45

Future Work

- ▶ Prove improvement of generalization
- ► Test performance against recent approaches
- Tweak model properly
- If performance and generalization is good enough:
 - ► Test some predicted interactions in vitro
 - Use DTI for Side-effect-phenotype mapping



Full citations

- Yu Ding, Hong Wang, Hewei Zheng, Lianzong Wang, Guosi Zhang, Jiaxin Yang, Xiaoyan Lu, Yu Bai, Haotian Zhang, Jing Li, Wenyan Gao, Fukun Chen, Shui Hu, Jingqi Wu, Liangde Xu, Evaluation of drug efficacy based on the spatial position comparison of drug-target interaction centers, Briefings in Bioinformatics, , bbz024, https://doi.org/10.1093/bib/bbz024
- Chen Wang, Lukasz Kurgan, Review and comparative assessment of similarity-based methods for prediction of drug-protein interactions in the druggable human proteome, Briefings in Bioinformatics, , bby069, https://doi.org/10.1093/bib/bby069
- Toward a complete dataset of drug-drug interaction information from publicly available sources,
 - SerkanAyvazaJohnHornbOktieHassanzadehcQianZhudJohannStaneNicholas P.TatonettifSantiagoVilarfMathiasBrochhausengMatthiasSamwaldhMajidRastegar-MojaradiMichelDumontierjRichard D.Boyce, 2015

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