# Predicting Compound-Protein Interactions From Machine Learning Perspective

PhD Candidate: Songpeng Zu Advisor: Shao Li

FIT 1-108, Tsinghua University

May 27, 2015

#### Outline

- Quantitatively Predicting Compound-Protein Interactions by Multi-Task Learning
- ► Inference on Chemogenomic Features from Drug-Target Interactions
- ► Application on the Modification of Natural Products.

### Part I

Quantitatively Predicting
Compound-Protein Interactions by
Multi-Task Learning

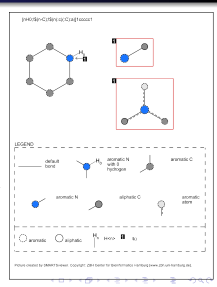
### Background

#### Methods for predicting compound-protein interactions (CPIs):

- Structure-based molecular dynamics
  - depend on proteins' 3D structures.
- Ligand-based method
  - can be independent of proteins' 3D structures.
  - large-scale known CPIs data
  - mainly dependent on machine learning approaches.

## Machine Learning on CPIs

- Compounds represented by topological fingerprints. The similar as proteins.
- CPIs recorded as binary variable or continuous variables.
- Classification or regression models then are used.



## Modeling on a single protein

Keiser M.J. *et al.*, *Nature* 2009, developed the SEA method to predict drugs' new molecular targets.

- ► Each target represented by its set of known ligands.
- Drugs computationally screen against a panel of proteins by comparing the similarity of ligands against these proteins.
- ► The similarities expressed as E-values, adapting the BLAST algorithm.

## Modeling on a single protein

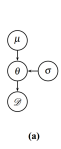
Besnard J. *et al.*, *Nature* 2012, used naive Bayesian model to predict compounds' polypharmacology profiling.

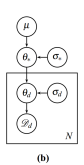
- ▶ 215,000 activity data including 133,061 compounds and 784 proteins were used.
- Every compounds represented by the binary vectors of ECFP6 representations.
- For every protein, a Laplacian-modified naive models was built for classification.

## Modeling on a protein family

Yabuuchi H. *et al.*, *Molecular Systems Biology* 2011 developed the CGBVS framework.

- ► 5207 CPIs data (including 317 GPCRs and 866 ligands)
- Compounds' structure and proteins' sequences converted into 929- and 400-dimensional vectors
- SVM then used.





### Machine Learning on CPIs

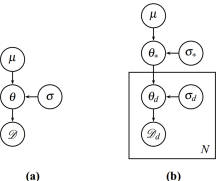
#### Current machine learning on predicting CPIs

- Modeling on a single protein
   More specificity Lots of data needed
- Modeling on a protein family
   Data sharing Less specificity

### Multi-Task Learning

#### Can we combine the two approaches?

- Learning different but similar tasks at the same time. (Finkel J.R. and Mannning C.D., 2009)
- Quantitative prediction.



### Hierarchical Bayesian Model

Suppose  $\mathcal{D}_j = \{\mathbf{X}_j, \mathbf{y}_j\}$ , j = 1, ..., m, and  $\mathbf{X}_j \in \mathbb{R}^{d \times n_j}$ . Then we have

$$\mathbf{y}_{j} \sim \mathcal{N}\left(\mathbf{X}_{j}^{T} \omega_{j}, \sigma_{y}^{2} \mathbf{I}\right) \tag{1}$$

Since different groups data may share similar features, we assume  $\omega_j$  have the same mean on the prior distribution.

$$\omega_j \sim \mathcal{N}\left(\omega_*, \sigma_j^2 \mathbf{I}\right)$$
 (2)

In which,

$$\omega_* \sim \mathcal{N}\left(\mu, \sigma_*^2 \mathbf{I}\right) \tag{3}$$

Suppose, for simplicity, that  $\mu = 0$ ,  $p(\sigma_v^2) \propto 1$ , and that  $\sigma_i^2$  and  $\sigma_*$  are fixed. Let  $\Theta = \{\omega_i, j = 1, ..., m, \omega_*, \sigma_v^2\}$ . We have

$$\mathcal{L}_{hier}(\mathcal{D};\Theta) = \mathcal{L}_{orig}(\mathcal{D}|\Theta) + logp(\Theta)$$

$$= \sum_{j} \left( logp(\mathcal{D}_{j}|\omega_{j}) - \frac{\parallel \omega_{j} - \omega_{*} \parallel^{2}}{2\sigma_{j}^{2}} \right) - \frac{\parallel \omega_{*} \parallel^{2}}{2\sigma_{*}^{2}}$$

$$- \sum_{j} \frac{d}{2} log(2\pi\sigma_{j}^{2}) - \frac{d}{2} log(2\pi\sigma_{*}^{2})$$

$$= \sum_{j} \left( -\frac{\parallel \mathbf{y}_{j} - \mathbf{X}_{j}^{T} \omega_{j} \parallel^{2}}{2\sigma_{y}^{2}} - \frac{\parallel \omega_{j} - \omega_{*} \parallel^{2}}{2\sigma_{j}^{2}} \right) - \frac{\parallel \omega_{*} \parallel^{2}}{2\sigma_{*}^{2}} - \sum_{j} \frac{n_{j}}{2} log(2\pi\sigma_{y}^{2})$$

$$- \sum_{j} \frac{d}{2} log(2\pi\sigma_{j}^{2}) - \frac{d}{2} log(2\pi\sigma_{*}^{2})$$

L-BFGS-B optimization method is then used following the gradient below.

$$\frac{\partial \mathcal{L}_{hier}(\mathcal{D};\Theta)}{\partial \omega_{j}} = -\frac{1}{2\sigma_{y}^{2}} \frac{\parallel \mathbf{y}_{j} - \mathbf{X}_{j}^{T} \omega_{j} \parallel^{2}}{\partial \omega_{j}} - \frac{1}{2\sigma_{j}^{2}} \frac{\parallel \omega_{j} - \omega_{*} \parallel^{2}}{\partial \omega_{j}}$$

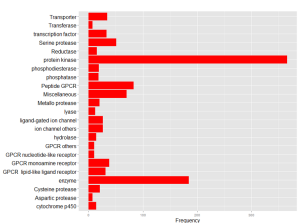
$$= \frac{\mathbf{X}_{j} \mathbf{y}_{j}}{\sigma_{y}^{2}} + \frac{\omega_{*}}{\sigma_{j}^{2}} - \left(\frac{\mathbf{X}_{j} \mathbf{X}_{j}^{T}}{\sigma_{y}^{2}} + \frac{1}{\sigma_{j}^{2}} \mathbf{I}\right) \omega_{j} \tag{5}$$

$$\frac{\partial \mathcal{L}_{hier}(\mathcal{D}; \Theta)}{\partial \omega_*} = -\sum_j \frac{\omega_* - \omega_j}{\sigma_j^2} - \frac{\omega_*}{\sigma_*^2} \tag{6}$$

$$\frac{\partial \mathcal{L}_{hier}(\mathcal{D}; \Theta)}{\partial \sigma_{y}^{2}} = \frac{\sum_{j} \| \mathbf{y}_{j} - \mathbf{X}_{j}^{T} \omega_{j} \|^{2}}{2(\sigma_{y}^{2})^{2}} - \frac{n}{2\sigma_{y}^{2}}$$
(7)

where n is the total number of samples in all the groups.

- ▶ 210,000 CPIs including more than 1,000 proteins from 20 protein families, and 150,000 compounds.
- ▶ 22 physicochemical properties and 881 chemical substructures as the compounds' features.

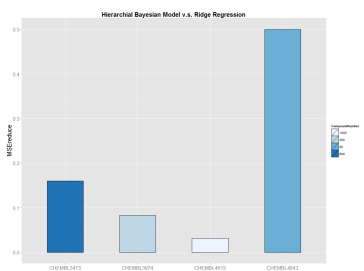


### **Feature Selection**

The protein family of Peptide GPCR including 85 proteins as examples.

- ▶ Based on the definitions of chemical fingerprints, SUB1-SUB115, SUB264-SUB327 are removed.
- Chemical fingerprints with too low or high frequencies are removed.
- Non-parametric dynamic slicing method for marginal feature selection.
- ▶ 284 features are finally kept.

## Comparison with Ridge Regression



### Discussion

- More computational tests
- ► The relationship between proteins' pharmacological and genomic information
- ▶ Deficiency:
  - High dimension v.s. Sparsity
  - Linear v.s. Nonlinear

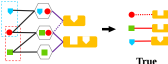
### Part II

Inference on Chemogenomic Features from Drug-Target Interactions

## Background

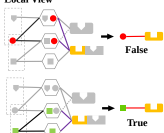
#### A. Goal





C. Global View: GIFT

#### **Local View**





Known interacting pairs

...... Known non-interacting pairs

Predicted interacting pairs

Predicted non-interacting pairs

#### Definition

- $O_{ij}$  The observations of drug and protein interactions.
- $YP_{ij}$  The binary variable of drug i and protein j interactions.
- $D_{mn}^{(ij)}$  The interaction result of substructure m from drug i and domain n from protein j

$$\theta_{mn}$$
  $\theta_{mn} = Pr(ZD_{mn} = 1)$ 

fn fn = 
$$Pr(O_{ij} = 0 | YP_{ij} = 1)$$

$$\mathsf{fp} \ \mathsf{fp} = Pr(O_{ij} = 1 | \mathit{YP}_{ij} = 0)$$

## **Assumption**

Consistency

$$\theta_{mn} = Pr(D_{mn}^{(ij)} = 1) \tag{8}$$

Independence

$$Pr(YP_{ij} = 1|\theta) = 1 - \prod_{D_{mn}^{(ij)}} (1 - \theta_{mn})$$
(9)

## **EM Algorithm**

▶ Then the log likelihood function is followed:

$$l(\theta) = \log\left(\Pr(O|\theta)\right) \tag{10}$$

$$Pr(O_{ij} = 1|\theta) = (1 - fn)Pr(YP_{ij} = 1|\theta) + fp \cdot Pr(YP_{ij} = 0|\theta)$$
(11)

▶ The EM Algorithm is used to get the MLE estimation due to the missing data of  $D_{mn}$ .

### **EM Algorithm**

► E Step:

$$E(D_{mn}^{(ij)}|O,\theta^{(t-1)}) = \frac{\theta_{mn}^{(t-1)}(1-fn)^{O_{ij}}fn^{1-O_{ij}}}{Pr(O_{ij}|\theta^{(t-1)})}$$
(12)

M Step:

$$\theta_{mn}^{(t)} = \frac{1}{N_{mn}} \sum_{i,j:Zm \in Y_i, Dn \in P_j} E(D_{mn}^{(ij)} | O_{ij}, \theta^{(t-1)})$$
(13)

### Variance Estimation

► The variance of the parameters are estimated by the observed Fisher information.

$$var(\hat{\theta}) = \frac{1}{I(\hat{\theta})}, I(\theta) = -\frac{d^2 log(Pr(O|\theta))}{d\theta^2}$$
 (14)

▶ In our model, the observed Fisher information is followed:

$$I(\theta_{mn}) = \sum_{i,j:Zm \in Y_i, Dn \in P_j} \delta_{mn}^{(i,j)^2} \left(\frac{O_{mn}^{(ij)}}{\mu_{mn}^{(ij)^2}} + \frac{1 - O_{mn}^{(ij)}}{(1 - \mu_{mn}^{(ij)})^2}\right)$$
(15)

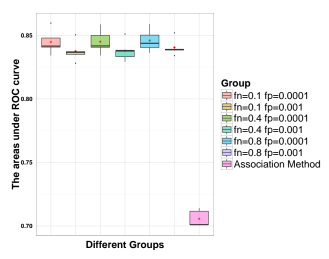
In which,

$$\delta_{mn}^{(ij)} = \frac{\mu_{mn}^{(ij)}}{\partial \theta_{mn}}, \mu_{mn}^{(ij)} = Pr(O_{mn}^{(ij)} = 1 | \theta)$$
 (16)

### **Data Source**

- ▶ 1862 drugs are represented by 881-dimensional chemical substructure binary vectors defined by the PubChem database.
- ▶ 1554 proteins are represented by 876-dimensional protein domain binary vectors from the Pfam database.
- ▶ 4809 interactions between drugs and proteins.

▶ Different combinations of fn and fp.



Comparison with other methods.

Ratio	GIFT	L1-Log	L1-SVM	SCCA
1	0.835	0.829	0.830	0.798
5	0.847	0.838	0.855	0.798

▶ Results of predictions on known drug-domain interactions.

Table 2. Representative results of the predictions on drug-domain interactions.

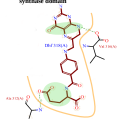
Protein	Drug	Domain	k value	Prediction
DNA (cytosine-5)-methyltransferase 1	S-Adenosylhomocysteine	C-5 cytosine-specific DNA methylase	1	TRUE
Alcohol dehydrogenase 1B	N-benzylformamide	Alcohol dehydrogenase GroES-like domain	0.58	TRUE
Androgen receptor	Flufenamic Acid	Ligand-binding domain of nuclear hormone receptor	0.9	TRUE
Ornithine carbamoyltransferase	N-(Phosphonoacetyl)-L-ornithine	Asp/Orn binding domain	0.51	TRUE
Progesterone receptor	Norethindrone	Ligand-binding domain of nuclear hormone receptor	1	TRUE
Rho-associated protein kinase 1	hydroxyfasudil	Protein kinase domain	0.94	TRUE
Tissue-type plasminogen activator	benzamidine	Trypsin	1	TRUE

k value is the proportion of the number of the binding positions in one domain over the total number of the binding positions. If k is no less than 0.5, the drug and domain interacts. TRUE means the predicted score of drug-domain interaction by GIFT is larger than zero.

#### A. Methotrexate against dihydrofolate reductase domain



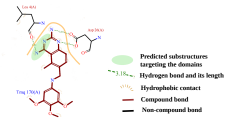
#### C. DHF against thymidylate synthase domain



#### Pemetrexed against dihydrofolate В. reductase domain



#### D. Trimetrexate against dihydrofolate reductase domain



### Discussion

- Here we propose an efficient method to extract meaningful chemogenomic features, and it also shows the power to predict drug-protein interactions.
- The predicted chemical substructures might be a useful source to design the compounds' analogs against a given protein or its domain.
- ▶ Large-scale compound-protein interactions are accumulated in the PDB database (known 3D structures), BindDB and ChEMBL database, which can be further studied by our method.

### Part III

Application on the Modification of Natural Products

## Lead Discovery From TCM Herbs

- ▶ Natural products important sources for drug discovery.
- By DrugCIPHER, several compounds from traditional Chinese Medicine (TCM) Herbs are predicted to have the antitumor activities.
- ► Many of them have been reported, but one compound called *Albiflorin* few researches.

- Our experiments: Albiflorin has the antitumor activities with low potency.
- ▶ Its biological mechanism is unknown.

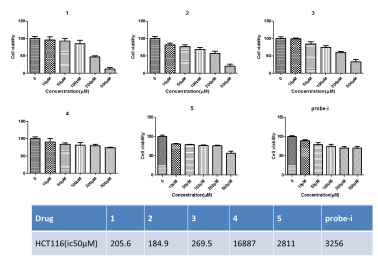
- ► *Albiflorin*, a typical example from natural products.
  - Low potencies or activities
  - Unknown targets
- ▶ Direct experiments difficult to discover the mechanisms.
  - Low potencies → false negative
  - Unknown targets → hard to design analogs
  - Complex structures

#### Method

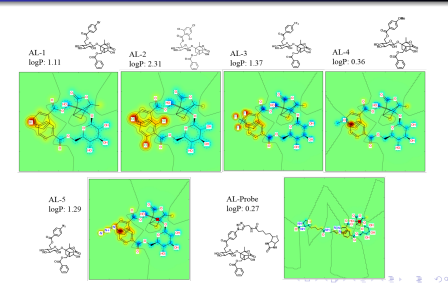
- Firstly, several analogs are designed based on the chemical experience as a starting point.
- ► Then MTT assays are used to test their biological activities on tumor growth.
- ▶ Next structure-activity relationship (SAR) analysis is performed to predicted its possible functional mechanism.
- Simulation and Filtering
  - Computational simulation of all the possible analogies.
  - Quantitatively predicting their targets.
- ► Experimental design and validation.



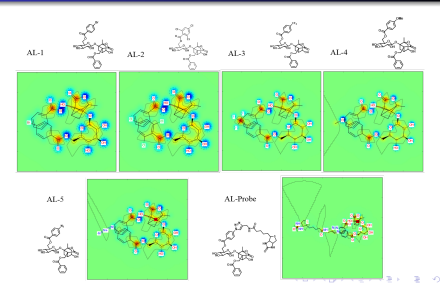
### MTT Assay



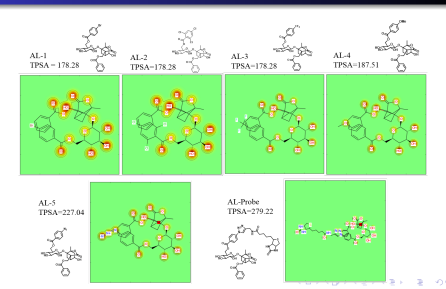
### LogP Analysis



### Partial Charge



### **TPSA**



### Discussion

Explore a new strategy to study natural products.

- Discovery by computational methods.
- Biological experiments validation.
- ► Computational Simulation and analysis all the possible analogs.
- ▶ Medicinal chemistry-based experiments validation.