

Kernel methods for *in silico* chemogenomics

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- Drug discovery
- Chemogenomics

2 Method

- Formalization
- Representation of pairs
- Kernel for ligands and targets

3 Results

4 Conclusion

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Classical approach

- Imitate traditional remedies.
- Accidental discoveries.

New trend

- ① Understand underlying biological process.
- ② Identify **targets** (typically proteins).
- ③ Identify **modulators** of these targets.

Typical targets

- G-protein-coupled-receptors (GPCR).
- Enzymes.
- Ion channels.

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- Need to test a huge number of candidate molecule against a target.
- *In silico* interaction prediction is therefore a key element.

Classical paradigms

- Ligand-based : compare candidate ligand to **known** ligands of the target (e.g. using machine learning).
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- Ligand-based : need to know (enough) ligands of a given target to produce (accurate) predictors.
- Structure-based :
 - Time-consuming.
 - Need to know the 3D structure of the target.

Chemogenomic framework

- Idea: mine the chemical space (small molecules) against the *whole* biological space (targets).
- Similar molecules bind similar targets.
- Advantage: ligand-based approaches on targets with no (or few) known ligands can take advantage of similar targets with known ligands.

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- Apply existing machine learning algorithms in this space.
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Single-target screening

- Target t with known ligands c_1, \dots, c_n .
- For each t , learn a function $f_t(c)$ from the c_i that predicts if unseen candidate c is a ligand of t .
- Linear case: given a description $\Phi(c)$ of the molecule,
 $f_t(c) = w_t^\top \Phi(c)$.

Chemogenomics setting

- Consider training **pairs** $(t, c)_i$ (known to interact or not to interact), represented by vectors $\Phi((t, c)_i)$.
- Learn a single function $f(t, c) = w^\top \Phi(t, c)$ in the joint space to predict if the candidate pair (t, c) interacts.
- How to choose the pair representation Φ ?

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Vector representation of pairs

Ligand representation

- A lot of existing work to represent a molecule t by a vector $\Phi_{ligand}(c) \in \mathbb{R}^{d_c}$.
- Physico-chemical, structural properties of the molecules.

Target representation

- Similarly, much work devoted to the construction of descriptors for a given protein t by a vector $\Phi_{target}(t) \in \mathbb{R}^{d_t}$.
- Properties of the sequence, structure of the protein.

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Vector representation of pairs

Pair representation

- Use products of features of c and features of t .
- Idea (binary case): indicate that both c and the t carry given features.
- May be strongly correlated with the fact that they interact.
- Set of all possible products of features of c and t is given by the tensor product:

$$\Phi(c, t) = \Phi_{ligand}(c) \otimes \Phi_{target}(t). \quad (1)$$

- **Potential issue:** $d_c \times d_t$ vector, may be prohibitively large.

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Kernels for ligand-target pairs

Kernel trick

- Need to compute efficiently the inner product between pairs.
- Classical property of tensor products :

$$\begin{aligned}\Phi(c, t)^\top \Phi(c', t') &= (\Phi_{lig}(c) \otimes \Phi_{tar}(t))^\top (\Phi_{lig}(c') \otimes \Phi_{tar}(t')) \\ &= \Phi_{lig}(c)^\top \Phi_{lig}(c') \times \Phi_{tar}(t)^\top \Phi_{tar}(t').\end{aligned}$$

More generally

Denoting

$$K_{lig}(c, c') = \Phi_{lig}(c)^\top \Phi_{lig}(c'), \quad K_{tar}(t, t') = \Phi_{tar}(t)^\top \Phi_{tar}(t'),$$

we obtain the inner product between tensor products by:

$$K((c, t), (c', t')) = K_{tar}(t, t') \times K_{lig}(c, c'). \quad (2)$$

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Kernel for ligands

- We chose the *Tanimoto kernel* (state-of-the-art performances in general).
- Characterizes the molecules by the occurrences of linear subgraphs of length 8 or less.

Kernels for targets

Dirac kernel

$$K_{dirac}(t, t') = \delta(t, t').$$

Equivalent to performing **independent** learning for each target.

Multitask kernel

$$K_{multitask}(a, a') = K_{dirac}(a, a') + 1.$$

Naive information sharing.

Sequence-based kernels

Classical **mismatch** and **local alignment** kernel on whole sequences.

Hierarchy kernel

Use KEGG hierarchy between the targets: number of common ancestors in the corresponding hierarchy plus one.

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Data generation

- KEGG data base (Kanehisa *et al.*, 2002).
- Ligand data for GPCR, enzymes and ion channels.
- For each positive pair, generate a negative ligand-target pair (same target, random ligand among existing ligands).

Final benchmark

- 2436 pairs for enzymes, 798 for GPCR and 2330 for ion channels.
- First experiment: 10-fold cross validation (assess the incidence of using ligands from other targets on the accuracy of the learned classifier for a given target).
- Second experiment: for each t learn a classifier using only interactions that do not involve t and test on the points that involve t (simulate the behavior when making predictions for orphan targets).

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First experiment

$K_{tar} \setminus$ Target	Enzymes	GPCR	Channels
Dirac	0.536 ± 0.005	0.682 ± 0.022	0.701 ± 0.017
multitask	0.874 ± 0.008	0.595 ± 0.030	0.797 ± 0.017
hierarchy	0.877 ± 0.008	0.817 ± 0.025	0.857 ± 0.015
local alignment	0.544 ± 0.007	0.696 ± 0.033	0.824 ± 0.015

Prediction accuracy for the first protocol on each dataset with various target kernels.

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Second experiment

$K_{tar} \setminus$ Target	Enzymes	GPCR	Channels
Dirac	0.500 ± 0.000	0.500 ± 0.000	0.500 ± 0.000
multitask	0.856 ± 0.009	0.477 ± 0.025	0.636 ± 0.021
hierarchy	0.862 ± 0.009	0.776 ± 0.026	0.805 ± 0.018
local alignment	0.521 ± 0.004	0.647 ± 0.030	0.722 ± 0.019

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Still possible to obtain reasonable results when no ligand is known for the target.

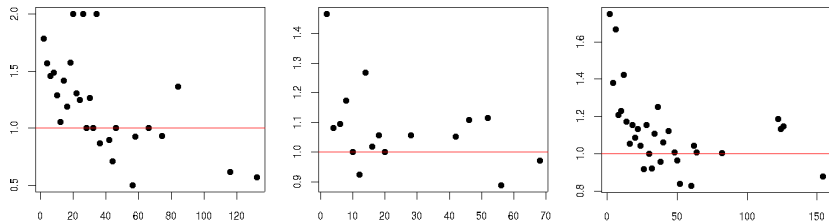
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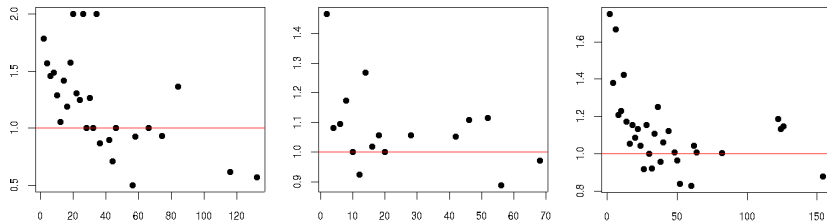
General behavior



Relative improvement of the *hierarchy* kernel against the *Dirac* kernel as a function of the number of known ligands for enzymes, GPCR and ion channel datasets.

- Strong improvement when few training points available.
- After a certain point, using similar targets can deteriorates accuracy.
- Suggests that the method could be improved by learning for each target independently how much information should be shared.

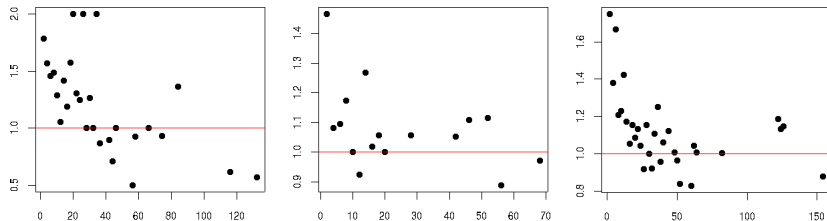
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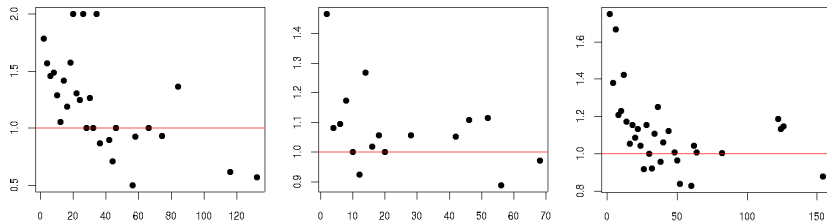
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Conclusion

- General method to predict interaction between any chemical compound and any biological target.
- Using target kernels allowing to share information across the targets improves the prediction.
- Accuracy improvement depends on the number of known ligands.
- Possible improvements:
 - Other kernels (using molecule 3D information or sequence/structure of targets).
 - Adapt the amount of information sharing to each target.
 - Other regularizations.

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