A KDD APPROACH FOR DESIGNING FILTERING STRATEGIES TO IMPROVE VIRTUAL SCREENING

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Abstract: Virtual screening has become an essential step in the early drug discovery process. Generally speaking, it consists in using computational techniques for selecting compounds from chemical libraries in order to identify drug-like molecules acting on a biological target of therapeutic interest. In the present study we consider virtual screening as a particular form of the KDD (Knowledge Discovery from Databases) approach. The knowledge to be discovered concerns the way a compound can be considered as a consistent ligand for a given target. The data from which this knowledge has to be discovered derive from diverse sources such as chemical, structural, and biological data related to ligands and their cognate targets. More precisely, we aim to extract filters from chemical libraries and protein-ligand interactions. In this context, the three basic steps of a KDD process have to be implemented. Firstly, a model-driven data integration step is applied to appropriate heterogeneous data found in public databases. This facilitates subsequent extraction of various datasets for mining. In a second step, mining algorithms are applied to the datasets and finally the most accurate knowledge units are eventually proposed as new filters. We present here this KDD approach and the experimental results we obtained with a set of ligands of the hormone receptor LXR.

1 INTRODUCTION

In silico drug discovery covers diverse computational techniques for capturing, integrating and analyzing biological and chemical data from diverse sources. Many programs address the issue of identifying drug-like molecules by calculating the docking energies of ligands bound to biological targets. Indeed, virtual screening is recognized today as a very promising process in early drug discovery process because it provides an excellent cost-to-efficiency ratio (Jorgensen 2004; Köppen 2009). However high-throughput virtual screening methods are still under-exploited due to the computing cost of the current docking programs. One way to overcome such limitations is to couple multiple techniques in a funnel-like filtering process in which fast selection methods are used first for discriminating candidates that can be quickly recognized as consistent for being passed to the next step of the funnel. Filters that can be used for this first fast selection step are classically grouped into two categories. On one hand, the structure-based methods involve computing either geometrical matching between target and ligand, or a combination of features characterizing the binding mode of ligand to target (pharmacophore, Finn et al. 1998). These methods require that the 3D structure of the target is known. On the other hand, the ligand-based methods rely on a representative set of reference structures, known to be biologically active on the target, and compute structure-activity relationships based on various molecular descriptors. Both categories (structure-based and ligand-based) of methods result in a ranked list of screened compounds.

Actually, the design of a virtual screening filter can be considered as a particular case of the KDD (Knowledge Discovery from Databases) approach. The knowledge to be discovered concerns the discrimination between good and bad ligands for a given target, i.e. a classification problem. The data to be mined for knowledge extraction are chemical, structural and biological data related to the ligands and their cognate targets. Indeed the powerful KDD
paradigm (Fayyad *et al.* 1996) provides a consistent way to address a virtual screening issue. It stresses the importance of data integration as a first preparation step and allows diverse mining algorithms to be applied on several selected subsets of the integrated data. Knowledge units can be extracted from these datasets to derive activity prediction models. Once validated, such prediction models can be used as a novel type of virtual screening filters. Since they are produced along a KDD process, these filters will be called here “knowledge-based” filters.

The KDD approach presented in this paper concerns the definition of new virtual screening filters in a drug discovery context. Special emphasis is brought to the data integration step since the ligand descriptor space is huge and complex. Current programs are able to rapidly calculate hundreds of molecular descriptors corresponding to 1D, 2D and 3D physico-chemical descriptors. In most data analysis contexts, data integration efforts yield a simple matrix of data because most data mining algorithms accept as input unique tables where the data are represented as objects displaying specific values for given properties. However, a single table representation hardly reflects the complexity of biological and chemical data related to Protein-Ligand Interaction (PLI) data. Our approach is thus rather based on an entity-relationship data model. An integrated database is then produced from which various sets of data can be easily extracted for mining as in Karp *et al.* (2008). Interestingly, this architecture revealed to be useful for solving the multiple-instance learning problem that arises when considering simultaneously the descriptors of the ligands and their 3D conformations.

The proposed KDD methodology has been tested on three targets corresponding to three distinct 3D conformers of the same protein. The challenge addressed here is to combine various sets of ligand descriptors, pertaining to both structure-based and ligand-based methods. Section 2 describes the biological background of this study; the proposed KDD approach is presented in section 3; section 4 reports on the results of the conducted experiment. The last section concludes on the advantages and perspectives of this approach.

## 2 PREDICTION OF LIGAND ACTIVITY FOR DRUG DISCOVERY

Several programs exist for both ligand- and structure-based screening methods (Kirchmair *et al.* 2008) and recent developments confirm that combining results from different methods leads to better docking performance (Feher 2006). Several combination methods have been proposed among which the recent VSM-G approach that designs the hit identification process as a funnel of several progressive screening programs (Beautrait *et al.* 2008). It is composed of a rigid geometrical docking program (SHEF, Cai *et al.* 2008) followed by a flexible docking program (GOLD, Jones *et al.*, 1997), both programs acting obviously as structure-based filters. Since the number of false positive hits is still very high, one direction for improving VSM-G is to develop knowledge-based filters. This should reduce the number of false positive hits that are finally retained.

The KDD approach presented in this paper is tested with a collection of molecules known for their activity towards a particular biological target, the Liver X Receptor (LXR). The LXR receptor is an attractive target for the development of new therapeutic agents in the treatment of cardiovascular-related diseases (Lala, 2005). Reports on structural characterization of the LXR receptor reveal a great plasticity of the ligand binding pocket, which is able to accommodate ligands with different shapes and sizes (Farnegardh *et al.* 2003). We consider in this study three distinct 3D conformations of the LXR target (codes: 1P8D, 1PQ6, and 1PQ9) obtained by X-ray crystallography.

## 3 MODEL-DRIVEN DATA INTEGRATION AND MINING

Our methodology is composed of four main steps: (i) building a data model for PLI data taking into account user requirements and existing resources; (ii) specifying a workflow for collecting data from the different resources leading to the specification of specific wrappers for populating a relational DB; (iii) writing queries on the data model for each identified user requirement; (iv) applying a mining program to the retrieved dataset. The last two steps can be iterated upon analysis of the extracted knowledge units.

Figure 1 presents the entity-relationship data model of the PLI database. The model contains five entities namely Protein, Ligand, P_L_Interaction, Protein Conformer, and Ligand Conformer, connected with relevant relationships. A protein is described by several attributes (e.g. Name,
Sequence, Size) available in the UniProtKB protein knowledge base. A set of physical and chemical attributes are computed by specific programs from each ligand with respect to its chemical formula. A protein may have a known interaction with a ligand. Each PLI is documented either in the PDB (Protein Data Bank; Berman et al. 2000), Pubmed or IntAct databases by a set of characteristics (e.g. EC50, Kd). The Protein_Conformer and Ligand_Conformer entities contain topological descriptors of 3D conformations for each protein and each ligand. These include Spherical Harmonic (SH) coefficients which describe the shapes of the target binding site and of the ligand for easy comparison (Cai et al., 2008).

The overall KDD strategy is figured out in Figure 2. On the left, the original resources for the data relative to PLI are represented together with the main data flows for collecting relevant data concerning a list of targets of interest and a list of drug-like molecules. This leads to instantiate the PLI database (Figure 2, centre) for a given virtual screening problem. Once the PLI database is ready, the users can retrieve various datasets in order to design knowledge-based filters for virtual screening (Figure 2, right). At this stage of the work, the SQL view definition mechanism may constitute a powerful way for retrieving data sets to be mined. A typical dataset is composed of various ligands (set of objects) with their values for different descriptors (set of attributes), including a class attribute (active / inactive, or binding / not binding). Mining algorithms can then exploit such datasets in order to produce prediction models such as decision trees (DTs). Interestingly, the KDD process adapted to the virtual screening problem facilitates the exploration of the ligand descriptor space by selecting various descriptor sets and by evaluating the quality of the subsequent prediction models.

4 EXPERIMENTAL RESULTS
4.1 Instanciating the PLI Database

The PLI database was constructed according to a relational data model straight derived from the entity-relationship model shown in Figure 1. Data related to the three LXR conformers were imported from the PDB entries named 1P8D, 1PQ6 and 1PQ9 and used to fill the Protein_Conformer table. In particular, the structural descriptors of the binding pocket of each LXR conformer, including their SH coefficients, were computed and inserted in this table. A total of 222 LXR ligands were retrieved from the literature (Spencer et al., 2001; Bennett et al., 2008; Janowski et al., 1999) and inserted in the Ligand table. Their activity towards the LXR target was stored in the Protein_Ligand_Interaction table. The distinction between active and inactive ligands was based here on the transactivation (EC50) value found in the papers cited above. It was arbitrarily assumed that an active molecule is any molecule for which the transactivation value has been found lower than a given threshold of 1 μM (micromole per liter). This criterion yielded 157 active versus 65 inactive ligands in the database. About 20 possible conformers were generated for each ligand by a specific program (OpenEye Suite) in order to fill the Ligand_Conformer table with computed 3D structural descriptors of ligand conformers.
4.2 Datasets

Three sets of descriptors were considered for all the 222 ligands from the database. (i) The **SAR** descriptor set includes the classical ligand descriptors used for Structure-Activity Relationship analysis (Winkler 2002). This set corresponds to twenty-two attributes of the Ligand table. (ii) The **CONF** descriptor set includes six attributes corresponding to six 3D structural descriptors of ligand conformers stored in the Ligand_Conformer table. (iii) The **SAR-CONF** descriptor set is the union of the **SAR** and **CONF** descriptor sets (28 attributes). A class attribute (active/inactive) is added to each descriptor set.

In the **CONF** and **SAR-CONF** datasets several 3D conformers are associated with the same ligand. This leads to a multiple-instance learning problem (Maron & Lozano-Perez 1998) since the ligand conformers can be considered as distinct instances of the ligand, sharing common ligand properties (**SAR** descriptors and activity) but having specific conformer descriptors (**CONF** descriptors). To solve this problem we decided to select for each ligand the best-matching conformer towards each of the three LXR target conformers. This selection was based on the highest similarity score calculated with the SHEF program (Cai et al. 2008) between the SH coefficients of the ligand conformer on one hand, and of the binding pocket of the LXR conformer on the other hand. Finally three single-instance **CONF** (respectively **SAR-CONF**) descriptor sets were obtained, one for each LXR conformer.

4.3 Construction of Decision Trees

The mining experiments reported in this paper were carried out with the Weka machine learning program (Witten & Frank 2005) which includes an implementation of the J48 version of the C4.5 program for building Decision Trees (DTs) relying on the divide and conquer principle. The J48 program was run with the default parameters. The DTs were evaluated by a 10-fold stratified validation.

There are at least two reasons for using a DT-type mining algorithm for this experimentation. Firstly, we want to produce explicit activity prediction models in which the discriminative descriptors are made available to the domain experts. Secondly, the values taken by the descriptors in the datasets are not binary but rather numeric, which excludes in a first approach any symbolic data mining algorithm such as those searching for frequent itemsets or association rules.

4.4 Evaluation of the Prediction Models and Discussion

Applying the J48 program on the **SAR** and **SAR-CONF** datasets, using as class attribute the active/inactive attribute defined in section 4.1, failed to produce any consistent DT (no descriptor in the DT). The results simply lead to predicting the major class in all cases, resulting in an estimation of the maximal percentage of incorrectly classified instances of 32%. Conversely, DTs were obtained with the **CONF** datasets for each LXR conformer using the same active/inactive class attribute. The observed performances are presented in Table 1.

<table>
<thead>
<tr>
<th>LXR conformer</th>
<th>DT&lt;sub&gt;CONF&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>number of descriptors in the DT</td>
<td>2</td>
</tr>
<tr>
<td>number of false negatives / false positives</td>
<td>4 / 65</td>
</tr>
<tr>
<td>percentage of incorrectly classified instances</td>
<td>31%</td>
</tr>
<tr>
<td>weighted average of true positive rates</td>
<td>0.69</td>
</tr>
</tbody>
</table>

The accuracy of the prediction is very low for the three DTs. About 30% of the instances are incorrectly classified, which is very close to the maximal percentage of incorrectly classified instances. The number of false positive instances is high (61 to 65). Obviously, these results show that the considered descriptor sets cannot accurately predict ligand activity towards any of the three LXR conformers.

Since it is generally assumed that the activity is related to the binding, we decided to explore the capacity of the various descriptor sets to discriminate between binding and not binding ligands. Indeed the ultimate screening filter used in the VSM-G funnel is a flexible docking program that evaluates a docking score taking into account the flexibility of both target and ligand conformers. This flexible docking step requires powerful computing capacities to be conducted on large sets of molecules (about one hour is required on one processor core for docking one molecule on one target which means about 3 days for one thousand of molecules on a cluster of 16 bi-quad nodes). We therefore used the same datasets and simply
Table 2: Performance of the DTs predicting the docking of a ligand conformer with a given LXR conformer. Abbreviations are the same as in Table 1.

<table>
<thead>
<tr>
<th>LXR conformer</th>
<th>DT SAR</th>
<th>DT CONF</th>
<th>DT SAR-CONF</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1P8D</td>
<td>1PQ6</td>
<td>1PQ9</td>
</tr>
<tr>
<td>201/21</td>
<td>201/21</td>
<td>201/21</td>
<td></td>
</tr>
<tr>
<td>115/106</td>
<td>115/106</td>
<td>115/106</td>
<td></td>
</tr>
<tr>
<td>#docking / #not docking (in the dataset)</td>
<td>22</td>
<td>22</td>
<td>22</td>
</tr>
<tr>
<td>#descriptors (in the dataset)</td>
<td>6</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>#descriptors in the DT</td>
<td>1</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>#FN / #FP</td>
<td>9 / 9</td>
<td>6 / 11</td>
<td>3 / 12</td>
</tr>
<tr>
<td>Incorrectly classified instances</td>
<td>8%</td>
<td>7.6%</td>
<td>6.7%</td>
</tr>
<tr>
<td>Weighted average of TP rates</td>
<td>0.9</td>
<td>0.92</td>
<td>0.91</td>
</tr>
</tbody>
</table>

replaced the active/inactive class attribute with a docking/not docking class attribute. This information was produced for each ligand towards each LXR conformer with the Glide software (Halgren et al. 2004). The docking score was converted to a binary class attribute based on a docking score threshold.

The results are summarized in Table 2. The DT S, DT CONF, and DT SAR,CONF decision trees correspond to the SAR, CONF, and SAR-CONF description sets respectively. It appears clearly that the accuracy of docking prediction is globally much more satisfying than the accuracy of activity prediction was (Table 1). Less than 10% of the instances are incorrectly classified and the number of false positives is much lower (9 to 16). The accuracy figures of the DTs obtained with the three types of descriptor sets towards the three LXR conformers are very close one to the other. A possible comparison criterion is the number of attributes used in each DT, assuming that more efficient DTs use less attributes for the same accuracy. With such an hypothesis, the DT CONF perform better than the DT S and DT SAR,CONF for the three LXR conformers. For illustration, Figure 3 shows the docking DT CONF obtained for the 1PQ6 LXR conformer. The contribution of all these suggested filters has now to be evaluated upon screening a large molecule database against the considered targets. In particular it will be interesting to compare the efficiency of the VSM-G screening funnel with and without these additional filters.

The discrepancy observed between the activity and the docking prediction models raises the question of the differences that exist between binding and activity. Indeed, a retrospective analysis of the 222 molecules of our dataset reveals that for each target conformer (i) some active ligands are found unable to dock and (ii) some inactive ligands are docked. This apparent paradox can be explained by the fact that activity information is captured from functional biological tests in which the protein can adopt different conformations in addition to the three ones tested in the present study. Moreover, functional tests are designed for active compounds and cannot distinguish between binding and not binding inactive compounds.

**5 CONCLUSIONS AND PERSPECTIVES**

Our methodology for data integration and mining includes the rigorous construction of an integrated database in which data are collected from various resources. Careful design of such a database facilitates data preparation and selection upstream various data mining procedures when searching for significant hidden patterns. Moreover, it may help solving the multiple-instance learning problem by providing rapid access to the information required for converting a dataset into a single-instance one.

We have illustrated our approach with PLI data in a specific context of drug discovery. We have shown how the KDD methodology enables an actual exploratory data mining approach, leading to the choice of the best prediction models given three
types of descriptor sets. In our case, the suggested KDD approach succeeded in unifying the ligand- and structure-based approaches for virtual screening. The prediction models based on the CONF descriptor set can now be tested as knowledge-based filters in the VSM-G screening funnel upstream the flexible docking step in order to reduce the number of molecules to test with the docking software.

We see two main directions for future work. Firstly, we plan to use relational data mining methods for mining relational data and producing more expressive regularities (Finn et al., 1998; Dzeroski & Lavrac, 2001; Page & Craven, 2003). This would allow taking into account the chemical groups composing a ligand as well as atom-specific attributes. Secondly, we want to explore various definitions of ligand activity together with sets of relational descriptors for producing improved activity prediction models.

REFERENCES


