The Conjunctive Disjunctive Node Kernel

Dinh Tran Van¹, Alessandro Sperduti¹ and Fabrizio Costa²

1- Department of Mathematics, Padova University Trieste, 63, 35121 Padova, Italy

2- Bioinformatics Group, Department of Computer Science, Freiburg University Georges-Kohler-Allee 106, 79110 Freiburg, Germany

Abstract. Gene-disease associations are inferred on the basis of similarities between genes. Biological relationships that are exploited to define similarities range from interacting proteins, proteins that participate in pathways and gene expression profiles. Though graph kernel methods have become a prominent approach for association prediction, most solutions are based on a notion of information diffusion that does not capture the specificity of different network parts. Here we propose a graph kernel method that explicitly models the configuration of each gene's context. An empirical evaluation on several biological databases show that our proposal is competitive w.r.t. state-of-the-art kernel approaches.

1 Introduction and Related Work

Predictive systems for gene-disease associations are often based on a notion of similarity between genes. A common strategy is to encode relations between genes as a network and use graph based techniques to make useful inferences. In the last decades, a number of graph kernel methods have been proposed that directly exploit transitive properties in biological networks. The prototypical method is the Diffusion kernel (DK) [2] inspired by the heat diffusion phenomenon. The key idea is to allow a given amount of heat placed on nodes to diffuse through the edges. The similarity between two nodes v_i, v_j is then defined as the amount of heat starting from v_i and reaching v_j within a given time interval. In DK the heat flow is proportional to the number of paths connecting two vertices, which introduces an undesired bias that penalize peripheral nodes w.r.t. central ones. This problem is tackled by a modified version of DK called Markov exponential diffusion kernel (MED) [3] where a Markov matrix replaces the Laplacian matrix. Another kernel called Markov diffusion kernel (MD) [4], exploits instead the notion of diffusion distances, a measure of similarity between patterns of heat diffusion. The Regularized Laplacian kernel (RL) [5] represents instead a normalized version of the random walk with restart model and defines the node similarity as the number of paths connecting two nodes with different lengths. All these approaches can be applied to dense networks with high degree nodes. A drawback of these approaches is however their relatively low discriminative capacity. This is in part due to the fact that information is processed in an additive and independent fashion which prevents them from accurately modeling the configuration of each gene's context. To address this issue here we propose to employ a decompositional graph kernel (DGK) [1] technique. To exploit its higher discriminative capacity we first decompose the network in a collection of connected sparse graphs and then we develop a suitable kernel, that we call Conjunctive Disjunctive Node Kernel (CDNK).

2 Method

2.1 Network Decomposition

Dense networks, or networks with nodes that have a high degree, cannot be meaningfully processed by decomposition kernels based on neighborhood matches since even the immediate context can be too large and hence unique.

Notation and Definitions A graph G = (V, E) is a structure that consists of two sets: a node set V and an edge set E. The notation V(G) and E(G) are used to refer to the node set and edge set of G. The distance between two vertices u and v, notated as $\mathcal{D}(u,v)$, is the length of the shortest path between them. The neighborhood with radius r of a vertex v is the set of vertices at a distance no greater than r from v and denoted by $N_r(v)$. The induced subgraph $\mathcal{N}(W)$ is a graph induced from G with the node set W and the edge set containing every edge in G whose endpoints are in W. The neighborhood subgraph with radius r of vertex v is the subgraph induced by the neighborhood with radius r of v and is denoted by \mathcal{N}_r^v . The degree of a node v is the cardinality of its neighborhood set with radius 1 and is denoted as d(v). A clique of the graph G is a fully connected subgraph of G.

In this section, we introduce a new method for graph decomposition. The method is based on the idea of kcore and clique decomposition discussed in [8], [9] respectively. Besides, we pose to use two forms of edge called Conjunctive and Disjunctive. In the method, when checking the node degree or clique, we only consider the conjunctive. Given a graph G, a degree threshold D and a clique size threshold C, in the following we describe two steps of the method: kcore decomposition and clique decomposition.

Kcore Decomposition: Kcore decomposition intend to decompose a given graph to have a graph without any node degree greater than D. Kcore contains an iterative process in which each round consists of a procedure to alternately extract a high degree and a low degree subgraph from the input graph. The high degree subgraph (G_H) is the induced subgraph on the set of nodes with degree bigger than D, meanwhile the low degree subgraph (G_L) is the one on the set of nodes with degree smaller then D. At the first round, we takes G as the input. At the step i, the input graph is $G_{H_{i-1}}$ taken from the output of the step (i-1). The iteration stops when the G_{H_i} of the output is empty. When the decomposition process is done, we form a decomposed graph by making the union of all G_{L_i} taken from all rounds. We then add the set of edges from G that are not present in any G_{L_i} as the disjunctive edges of the decomposed graph.

Clique Decomposition: The clique decomposition begins with finding the set of cliques L in G that have size no less than C. For each clique in L, first, we add a new node to G. We then connect the new node to clique's nodes with disjunctive edges and to all neighborhood of clique's nodes at distance 1 with

conjunctive edges. Finally, we remove all conjunctive edges that have end points at one of the clique nodes.

2.2 Node Labeling

The similarity notion defined by a decomposition approach is based on the count of common fragments. A node kernel for single network cannot therefore have unique gene identifiers as node labels, but rather one has to define a small finite alphabet. We use the *gene ontology* [10] to construct binary vectors representing the bag-of- words encoding for each gene. The resulting representations are then clustered to finally yield a discretized label for each gene.

2.3 The Neighborhood Subgraph Pairwise Distance Kernel

The NSPDK is an instance of convolution kernel which is designed for measuring the similarity between graphs.

Given a graph $G \in \mathcal{G}$ (\mathcal{G} is the graph domain) and two rooted graphs A_u, B_v , the relation $R_{r,d}(A_u, B_v, G)$ is defined to be true iff A_u and B_v are in $\{\mathcal{N}_r^v : v \in V(G)\}$, where A_u (B_v) needs to be isomorphic with some \mathcal{N}_r and $\mathcal{D}(u, v) = d$. We denote R^{-1} as the inverse relation that returns subgraphs of G, $R_{r,d}^{-1}(G) = \{A_u, B_v | R_{r,d}(A_u, B_v, G)\}$. The kernel $\kappa_{r,d}$ over $\mathcal{G} \times \mathcal{G}$ takes into account the number of identical neighboring graph pairs with radius r at distance d between two graph and is formulated as:

$$\kappa_{r,d}(G, G') = \sum_{\substack{A_v, B_u \in R_{r,d}^{-1}(G) \\ A'_{v'}, B'_{u'} \in R_{r,d}^{-1}(G')}} \sigma(A_v, A'_{v'}) \sigma(B_u, B'_{u'}),$$

where $\sigma(x,y)$ is the exact matching function that returns 1 if x is isomorphic to y and 0 otherwise. In order to solve the graph isomorphism problem, an efficient approximate algorithm is also proposed in [6]. Finally, the NSPDK is defined as $K(G,G') = \sum_{r} \sum_{d} \kappa_{r,d}(G,G')$. For efficiency issue, we limit the values of r and d with the upper bounds r^* and d^* , respectively.

2.4 The Conjunctive Disjunctive Node Kernel

In this section, we describe our proposed kernel, CDNK, which is a modification of NSPDK to measure the node similarity in the graph which consists of conjunctive and disjunctive edges. In our kernel, we consider only conjunctive edges when computing the distance between nodes and extracting neighborhood subgraphs. We define two relations: the conjunctive relation $R_{r,d}^{\wedge}(A_u, B_v, G)$ to be true iff (i) A_u and B_v are in $\{\mathcal{N}_v^v: v \in V(G)\}$, where A_u (B_v) needs to be isomorphic with some \mathcal{N}_r , (ii) $\mathcal{D}(u,v)=d$; and the disjunctive relation $R_{r,d}^{\vee}(A_u, B_v, G)$ to be true iff (i) A_u and B_v are in $\{\mathcal{N}_v^v: v \in V(G)\}$, where A_u (B_v) needs to be isomorphic with some \mathcal{N}_r , (ii) there exists a vertex w such that $\mathcal{D}(u,w)=d$, (iii) (w,v) is a disjunctive edge.

We define $\kappa_{r,d}$, an instance of the DGK on the relation $R_{r,d}^{\wedge}$ and $R_{r,d}^{\vee}$ as

$$\kappa_{r,d}(u,v) = \sum_{\substack{A_u, \ A'_{u'} \in (R_{r,d}^{\wedge}^{-1}(G) \cup R_{r,d}^{\vee}^{-1}(G)) \\ B_v, \ B'_{v'} \in (R_{r,d}^{\wedge}^{-1}(G) \cup R_{r,d}^{\vee}^{-1}(G))}} \mathbf{1}_{A_u \cong B_v} \cdot \mathbf{1}_{A'_{u'} \cong B'_{v'}},$$

where $\mathbf{1}_{A\cong B}$ is the indicator function that returns 1 if A is isomorphic to B and 0 otherwise. $\kappa_{r,d}$ counts the number of identical pairs of neighboring graphs of radius r at a distance d between two vertices. The CDNK is defined as $K(u,v) = \sum_{r} \sum_{d} \kappa_{r,d}(u,v)$.

3 Evaluation

We perform an empirical evaluation of the predictive performance of several kernel based methods on two of the datasets used in [3].

BioGPS: A gene co-expression network is constructed from BioGPS dataset, which contains 79 tissues in duplicates, measured with the Affymetrix U133A array. Pairwise Pearson correlation coefficients (PCC) are calculated and a pair of genes are linked by an edge if the PCC value is larger than 0.5.

Pathways: Pathway information is retrieved from KEGG, Reactome, PharmGKB and the Pathway Interaction Database. If a couple of proteins coparticipate in any pathway, the two corresponding genes are linked.

3.1 Evaluation Methods

We evaluate the performance of graph node kernels in the gene prioritization problem. Given a set of genes known to be associated to a given disease, gene prioritization is a task that aims to rank the candidate genes based on their probabilities to be related to that disease.

Similar to the evaluation process used in [3], we choose 12 diseases in which each one contains at least 30 confirmed genes. For each disease, we construct a positive set \mathcal{P} and a negative set \mathcal{N} . The set \mathcal{P} consists of all disease gene members. The set \mathcal{N} is built by randomly picking genes from known disease genes, genes associated at least to one disease class, but not related to the current class, such that $|\mathcal{N}| = \frac{1}{2}|\mathcal{P}|$. After that, leave-one-out cross validation is used to evaluate the performance of the algorithm. Each turn one gene is out to be the test gene and the rest are used to train the model using SVM. Starting from the output scores, we compute a decision score q_i for the test gene g_i as the top percentage value of score s_i among all scores. $q_i = \frac{|\{j|s_i \geq s_j\}|}{N}, i = 1, 2, \ldots, N$, where s_i , s_j are scores, N is the length of gene list. We collect all decision scores for every gene in the training set of a disease to form a global decision score list. The performance of the algorithm is measured by using AUC calculated that list.

3.2 Parameter Selection

In order to select the optimal parameter values for each kernel, we use one disease gene set for parameter selection and use Kfold with k equals to 3. We set the

	BioGPS					Pathways				
Disease	K1	K2	К3	K4	K5	K1	K2	K3	K4	K5
1	52/5	57/4	59/3	59/2	65/1	75/5	76/4	79/3	79/2	80/1
2	82/2	79/3	75/4	75/5	88/1	55/5	65/4	77/3	77/2	81/1
3	64/4	60/5	72/2	72/1	66/3	55/5	63/4	64/3	66/2	67/1
4	65/4	58/5	68/3	68/2	72/1	54/5	65/4	74/1	74/2	66/3
5	64/5	64/4	67/2	66/3	76/1	53/5	56/4	63/2	63/3	68/1
6	75/2	70/5	71/4	71/3	79/1	83/5	93/4	97/2	97/1	93/3
7	73/3	67/5	75/2	76/1	69/4	85/5	88/4	89/2	90/1	89/3
8	74/5	77/1	76/3	76/2	75/4	54/5	66/4	72/3	72/2	73/1
9	72/1	66/5	68/3	70/2	67/4	53/5	65/2	64/4	64/3	81/1
10	54/3	50/5	56/2	51/4	78/1	69/3	65/5	74/2	74/1	67/4
11	58/4	51/5	59/3	59/2	72/1	54/5	69/4	75/2	74/3	77/1
\overline{AUC}	66.6	63.5	67.8	67.5	73.3	62.7	70.1	75.3	75.5	76.5
\overline{Rank}	3.5	4.3	2.8	2.5	2.0	4.8	3.9	2.5	2.0	1.8

Table 1: The performance of kernels on different genetic diseases using BioGPS and Pathway dataset. Each element in the table shows the AUC in percentage and the order of kernel comparing to the rest (AUC/Rank). K1 = DK, K2 = MD, K3 = MED, K4 = RL, K5 = CDNK.

values for diffusion parameter in DK and MED as $\{10^{-3}, 10^{-3}, 10^{-2}, 10^{-1}\}$, for time steps in MD as $\{1, 10, 100\}$ and for RL parameter as $\{1, 4, 7\}$. For CDNK, we set values for degree threshold in $\{10, 15, 20\}$, clique size threshold in $\{4, 5\}$, maximum radius in $\{1, 2\}$, maximum distance in $\{2, 3, 4\}$. Finally, the C of SVM is set as $\{10^{-5}, 10^{-4}, 10^{-3}, 10^{-2}, 10^{-1}, 1, 10, 10^{2}\}$.

4 Results and Discussion

Table 1 shows the AUC performance of the models trained by using different graph node kernels on 11 genetic diseases using BioGPS and Pathways datasets. In the table, the best result on each disease is marked in bold. By observing the results, we note that the kernel CDNK perform the best results comparing with other considered kernels. Particularly, the CDNK is ranked at the first order in seven out of 11 diseases on both datasets. It also illustrates the highest results in average AUC and rank with 73.3/2.0, 76.5/1.8, and the AUC difference with the second best ones are 5.5% and 1% on BioGPS and Pathways, respectively. The MED and RL show similar and moderate results with small gap between them. Last, DK and MD demonstrate modest performance in average comparing with other adopted kernels. They are ranked in the last position in many diseases, especially 7 times out of 11 for MD in BioGPS and 10 out of 11 for DK in Pathways. While DK shows better performance than MD in BioGPS, it presents

worse in Pathways. In conclusion, CDNK outperforms all employed graph node kernels in term of both average rank and AUC measure.

The CDNK shows the state of the art results. However, in the case that the input graph has high average node degree and they are uniformly distributed, the decomposed graph is too sparse and it can lead our kernel to the poor performance.

5 Conclusions

We have shown how decomposing a network in a set of connected sparse graphs allows us to take advantage of the greater discriminative power of CDNK, a novel decomposition kernel, and achieve state- of-the-art results. In future work we will investigate how to extend the CDNK approach to gene-disease association problems exploiting multiple heterogeneous information sources.

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