Marginalized Denoising for Link Prediction and Multi-label Learning

Zheng Chen\textsuperscript{1,2}, Minmin Chen\textsuperscript{3}, Kilian Q. Weinberger\textsuperscript{2}, Weixiong Zhang\textsuperscript{2,1}

\textsuperscript{1}Institute for Systems Biology, Jianghan University, Wuhan, Hubei 430056, China.
\textsuperscript{2}Department of Computer Science and Engineering, Washington University in St. Louis, St. Louis, MO 63130, USA.
\textsuperscript{3}Criteo Lab, Palo Alto, CA, USA
zheng.chen@wustl.edu, m.chen@criteo.com, \{kilian, weixiong.zhang\}@wustl.edu

Abstract

Link prediction and multi-label learning on graphs are two important but challenging machine learning problems that have broad applications in diverse fields. Not only are the two problems inherently correlated and often appear concurrently, they are also exacerbated by incomplete data. We develop a novel algorithm to solve these two problems jointly under a unified framework, which helps reduce the impact of graph noise and benefits both tasks individually. We reduce multi-label learning problem into an additional link prediction task and solve both problems with marginalized denoising, which we co-regularize with Laplacian smoothing. This approach combines both learning tasks into a single convex objective function, which we optimize efficiently with iterative closed-form updates. The resulting approach performs significantly better than prior work on several important real-world applications with great consistency.

Introduction

We study two of the most prominent machine learning tasks on graphs: link prediction (LP) and multi-label learning (MLL). On an abstract level the two problems are similar: LP aims to impute missing links and MLL aims to impute missing labels. In real world applications, the two problems do indeed often appear hand-in-hand. Consider as an example protein-protein interaction (PPI) networks, which represent proteins bound to form protein complexes so as to exert their functions. Here, inferring interactions among proteins is an LP problem (Jansen et al. 2003) and predicting protein functions (labels) (Kourmpetis et al. 2010). Similarly in social networks, adding missing friendship edges will help predict interests of users, which are often shared amongst friends. Likewise, interests of users can be utilized to predict hidden links of friendship.

The most obvious approach to exploit such reciprocal relationship between LP and MLL is to alternately apply LP and MLL algorithms and use the output of one method to facilitate the other (Bilgic, Namata, and Getoor 2007). However, it is non-trivial to determine how frequently and to what extend information should be exchanged between LP and MLL. Most importantly, the alternating iterations between the two tasks are not guaranteed to converge.

We introduce a highly novel approach to combine the MLL and LP problem into a single joint objective with the recently introduced marginalized denoising framework (Chen et al. 2012; Chen, Zheng, and Weinberger 2013; Chen and Zhang 2014). We phrase both problems as instances of graph denoising and co-regularize the predictions with Laplacian graph regularization. Our resulting algorithm, which we refer to as Marginalized Label and Link Prediction (MLLP), is jointly convex and can be optimized very efficiently. Different from prior work (Wang, Huang, and Ding 2011), which essentially uses LP formulation to solve a MLL problem, we incorporate both problems into one single loss function and let them benefit from each other as information naturally flows both ways. We evaluated MLLP on three real-world applications - one social network and two PPI networks. MLLP outperforms every competing prior work consistently across all our benchmark data sets.
We consider the problems of LP and MLL on a graph \( \mathcal{G} = (\mathbf{G}, \mathbf{Y}) \) of \( n \) nodes in a transductive learning setting. \( \mathbf{G} \in \{0,1\}^{n \times n} \) denotes a partially observed relational graph between nodes, where \( G_{ij} = 1 \) iff we observe a link between nodes \( i \) and \( j \), and \( G_{ij} = 0 \), otherwise. For simplicity, we consider binary graphs throughout the paper, but in practice our method can be easily extended to weighted graphs, where \( G_{ij} \) takes real-valued edge weights. \( \mathbf{Y} \in \{0,1\}^{K \times n} \) represents partially observed labels of nodes, where \( K \) is the total number of labels and \( Y_{ki} = 1 \) iff node \( i \) is observed to be associated with the \( k \)th label, and \( Y_{ki} = 0 \), otherwise. Note that each node can be associated with more than one label. We denote the \( l \)th column of \( \mathbf{Y} \), i.e. the label vector of node \( i \), as \( y_i \), and without loss of generality, we assume the labels of the first \( l \) nodes, \( y_1, \ldots, y_l \), are given (albeit incomplete) and all the other labels are unknown.

Multi-label Learning on Relational Graphs

We first focus on the problem of inferring a more complete label matrix \( \hat{\mathbf{Y}} \in \mathbb{R}^{K \times n} \) from a partially observed label matrix \( \mathbf{Y} \). (The matrix \( \hat{\mathbf{Y}} \) can subsequently be thresholded to yield hard label assignments.) Given a partially observed adjacency matrix \( \mathbf{G} \), we can formulate the problem as a classical graph-based semi-supervised learning problem (Zhou and Schölkopf 2004):

\[
\arg\min_{\hat{\mathbf{Y}}} \text{Tr}(\hat{\mathbf{Y}} \mathbf{L} \hat{\mathbf{Y}}^\top) + \alpha \left\| \hat{\mathbf{Y}} - \mathbf{Y} \right\|_F^2, \tag{1}
\]

where \( \mathbf{L} = \mathbf{D} - \mathbf{G} \) is the graph Laplacian defined on graph \( \mathbf{G} \) and \( \mathbf{D} \) is a diagonal matrix with \( D_{ii} = \sum_j G_{ij} \). The first term of (1) measures the smoothness of the predicted scores \( \hat{\mathbf{Y}} \) on the graph structure and encourages the scores of adjacent nodes to be similar. The second term of (1) penalizes large deviations of the predictions from the given labels (and increases numerical stability). The hyperparameter \( \alpha > 0 \) adjusts the trade-off between these two objectives.

The optimization problem in (1) can be solved in a closed form as \( \hat{\mathbf{Y}} = \alpha \mathbf{Y} (\mathbf{L} + \alpha \mathbf{I})^{-1} \). In general, two different constants of \( \alpha \) for trade off can be used: one for the initially labeled nodes and the other for unlabeled nodes. This framework, in combination with alternative choices of the graph Laplacian matrix \( \mathbf{L} \), takes several existing classical graph-based semi-supervised learning algorithms as special cases (up to some scaling factors), e.g., the local and global consistent model of Zhou and Schölkopf (2003), and the random-walk with restart (RWR) model of Pan et al. (2004). In this study, we will follow the graph-based semi-supervised learning framework defined in (1) as the centerpiece for MLL.

Link Prediction on Bi-relational Graphs

We now consider the LP problem of inferring missing links in a partially observed graph \( \mathbf{G} \). Different from previous work, we also incorporate the known, albeit incomplete, labels \( \mathbf{Y} \) in the process.

Notation

Constructing a bi-relational graph

We first inject label information by augmenting the original graph \( \mathcal{G} \) with additional nodes and edges. Specifically, we introduce \( K \) new nodes, one per label. We refer to these newly added nodes as "label nodes" and the original nodes as "data nodes." In addition, we create a link between the \( i \)th data node and the \( k \)th label node if node \( i \) carries label \( k \), i.e. \( Y_{ki} = 1 \); see box 3 in Figure 1 for a schematic illustration. The resulting bi-relational graph characterizes relationships between two types of nodes, the data and label nodes, and we use \( \mathcal{G} \in \{0,1\}^{(n+K) \times (n+K)} \) to denote the link matrix of the new graph. \( \mathcal{G} \) can be decomposed as

\[
\mathcal{G} = \begin{bmatrix} \mathbf{G} & \mathbf{Y}^\top \\ \mathbf{Y} & \mathbf{H} \end{bmatrix}, \tag{2}
\]

where \( \mathbf{G} \) is the adjacency matrix of the original graph, \( \mathbf{Y} \) encodes the association between data nodes and labels (or label nodes), and \( \mathbf{H} \), which may be all-zeros or some prior that prescribes the similarities among labels. We use the cosine similarity among observed labels, as suggested before (Wang, Huang, and Ding 2011).

It is important to note that the bi-relational graph \( \mathcal{G} \) is considered to be only partially observed, because each of its components: \( \mathbf{G} \), \( \mathbf{Y} \), and \( \mathbf{H} \) may be incomplete. We will learn to predict missing links on the augmented bi-relational graph \( \mathcal{G} \) instead of the original graph \( \mathbf{G} \). Adding links between data and label nodes in \( \mathcal{G} \) is another form for completing \( \mathbf{Y} \), complementary to the graph Laplacian based method we introduced in the previous section. It also serves as a key component for linking LP and MLL.

Marginalized denoising model for link prediction

Different from most prior work, we use the recently developed marginalized denoising framework for link prediction (Chen et al. 2012; Chen, Zheng, and Weinberger 2013; Chen and Zhang 2014).

Consider the observed links in an "incomplete" bi-relational graph \( \mathcal{G}^* \) as randomly sampled from an (unknown) "complete" bi-relational graph \( \mathcal{G}^+ \). The goal for LP is to find a linear mapping \( \mathbf{W} \) to reverse the corruption process and approximately reconstructs the "complete" graph \( \mathcal{G}^+ \) from the "incomplete" graph \( \mathcal{G}^* \) such that \( \mathbf{W} \mathcal{G}^* \approx \mathcal{G}^+ \).

Because \( \mathcal{G}^* \) is unknown, we cannot learn \( \mathbf{W} \) directly. Instead, we construct an artificial training set of \( m \) further corrupted graphs of \( \mathcal{G}^1, \mathcal{G}^2, \ldots, \mathcal{G}^m \), by randomly removing edges with some probability \( p \in [0,1] \). We then train \( \mathbf{W} \) to reconstruct \( \mathcal{G}^* \) from these further corrupted graphs with

\[
\min_{\mathbf{W}} \frac{1}{m} \sum_{j=1}^m \left\| \mathcal{G} - \mathbf{W} \mathcal{G}^j \right\|_F^2 \tag{3}.
\]

If this corruption mechanism approximates how \( \mathcal{G}^* \) was generated from the "complete" graph \( \mathcal{G}^+ \), re-applying the learned mapping \( \mathbf{W} \) to \( \mathcal{G} \) will approximate the "uncorrupted" graph \( \mathcal{G}^* \). As far as the learning of \( \mathbf{W} \) is concerned,
it is generally better to set \( m \) as large as possible. We therefore consider the limiting case, \( m \to \infty \), so that (3) becomes

\[
\min_{\mathbf{W}} \mathbb{E} \left[ \left\| \mathbf{G} - \mathbf{W} \mathbf{G}^\top \right\|_F^2 \right]_{p(\mathbf{G} | \mathcal{Z})}. 
\]

This expectation is analytically computable in closed form (Chen et al. 2012). It is straightforward to show that (4) can be rewritten as

\[
\min_{\mathbf{W}} \text{Tr}(\mathbf{WQW}^\top - 2(1-p)\mathbf{WG}G^\top),
\]

where \( \mathbf{Q} = (1-p)^2 \mathbf{G}G^\top + p(1-p)\text{diag}(\mathbf{G}G^\top) \). Eq. (5) describes a parabola and its solution can be computed in closed form. Note that the expectation in (4) is still tractable even for more general corruption models (Chen et al. 2012; van der Maaten et al. 2013).

Because of the bi-relational nature of \( \mathbf{G} \), the mapping \( \mathbf{W} \) can be used for both label prediction and link prediction. To simplify notation, let \( \mathbf{W} = [\mathbf{W}_g \mathbf{W}_y] \), where the subscripts \( g \) and \( y \) denote the first \( n \) and last \( K \) \emph{rows} of \( \mathbf{W} \) respectively, and we can decompose \( \mathbf{G} = [\mathbf{G}_g \mathbf{G}_y]^\top \) (the first \( n \) and the last \( K \) \emph{columns}) accordingly. This approach can be used naturally for both link and label prediction.

**Link prediction.** By the definition of bi-relational graph \( \mathbf{G} \) in (2), the original graph \( \mathbf{G} \) corresponds to its upper-left corner. We can tease out this part of the interpolated link matrix as

\[
\mathbf{G}_{\text{dir}} = \mathbf{W}_g \mathbf{G}_g, \quad \mathbf{G}_{\text{undir}} = \frac{1}{2} (\mathbf{W}_g \mathbf{G}_g + \mathbf{G}_g^\top \mathbf{W}_g^\top),
\]

where \( \mathbf{G}_{\text{dir}} \) describes the solution in the scenario of directed graphs and \( \mathbf{G}_{\text{undir}} \) in the scenario of undirected graphs. (In the case of undirected graphs it is advantageous to symmetrize the output after the mapping with \( \mathbf{W} \).)

**Label prediction.** Imputed links between the data and label nodes can be interpreted as predictions of labels. Regardless of the nature of the input graph, we consider these links always undirected and we compute the predicted label matrix, \( \hat{\mathbf{Y}}' \), via link prediction as

\[
\hat{\mathbf{Y}}' = \frac{1}{2} (\mathbf{W}_y \mathbf{G}_g + \mathbf{G}_y^\top \mathbf{W}_g^\top).
\]

**Joint Objective**

In this section we combine the LP and MLL problems into a single joint objective, which allows information flow in both directions. The key idea, which forms the cornerstone of this study, is to establish the correspondence between LP and MLL such that these two tasks can exchange information under a unified framework. Note that the second term of the MLL objective (1), \( \| \hat{\mathbf{Y}} - \mathbf{Y} \|_F^2 \), penalizes large deviations of the learned scores \( \hat{\mathbf{Y}} \) from observed labels \( \mathbf{Y} \). These observed labels themselves, however, are notoriously incomplete. Instead of relying completely on the original observed labels, we encourage the two approaches, LP and MLL, to agree with each other with respect to their label predictions, \( i.e. \) we co-regularize the two predictions with the term \( \| \hat{\mathbf{Y}} - \mathbf{Y}' \|_F^2 \). We include the closed form solution for \( \hat{\mathbf{Y}}' \), eq. (7), in this co-regularizer and combine it with the two objectives (1) and (5) to obtain the final joint optimization:

\[
\text{Tr} \left( \hat{\mathbf{Y}}' \mathbf{Y}'^\top \right) + \alpha \left\| \hat{\mathbf{Y}} - \frac{1}{2} (\mathbf{W}_y \mathbf{G}_g + \mathbf{G}_y^\top \mathbf{W}_g^\top) \right\|_F^2 + \beta \text{Tr}(\mathbf{WQW}^\top - 2(1-p)\mathbf{WG}G^\top).
\]

This optimization is over \( \hat{\mathbf{Y}} \) and \( \mathbf{W} \). The constant \( \alpha \geq 0 \) trades-off the co-regularization terms and \( \beta > 0 \) balances the two tasks. Note that the last term is crucial for incorporating the original labels \( \mathbf{Y} \) in \( \mathbf{G} \). Without it, the optimization would yield a trivial all-zero solution for \( \mathbf{W} \) and \( \hat{\mathbf{Y}} \). Figure 1 sketches the intuition behind the joint optimization.

The objective in (8) is jointly convex and quadratic in \( \hat{\mathbf{Y}} \) and \( \mathbf{W} \). This means that we can obtain the global solution by solving both matrices in closed forms in an alternating fashion. Algorithm 1 summarizes the joint optimization algorithm with detailed alternating update rules. Note that the

![Figure 1: An illustration of MLLP. Each data node can be associated with one or more labels in a label set: Red, Green, and Blue. The original graph is augmented into a bi-relational graph, \( \overrightarrow{\mathbf{G}} \) (Box 3). The joint model (Box 2) encourages agreement between the predicted labels from MLL (Box 1) and the enriched labels from LP (Box 3).](image-url)
most computationally intensive operations, the matrix inversions, conveniently involve only with sparse matrices and are executed out the main loop.

Non-uniform co-regularization. In our formulation in (8), the observed labels only enter the optimization in LP (the third term). To propagate this information to the MLL objective (first term), it makes sense to co-regularize the two methods more for nodes that are accompanied with labels. In practice, we use different \( \alpha \)'s for nodes with and without labels \((\alpha_l, \alpha_u \text{ respectively})\). We have found empirically that it is best to set \( \alpha_l \gg \alpha_u \) when there are many unlabeled nodes.

Stacked link prediction. Chen et al. (2012) suggest to improve the performance of marginalized denoising in document representation through stacking. We observe a similar performance boost in our setting. In other words, after solving (8) we "enrich" the graph \( \tilde{G} \) with the mapping \( \tilde{W} \) and truncate all entries to be within \([0, 1]\).

We then use this enriched link matrix as the input for another round of optimization of (8). We repeat this process until the LP result stops to improve on a hold-out set.

Algorithm 1 Joint link prediction and multi-label learning

Input: Labels \( Y \), Graph \( G \), Parameters \( \alpha, \beta > 0 \), and level of corruption: \( 0 < p < 1 \)

Output: link prediction scores \( G \) and label prediction scores \( Y \).

Set \( H_1 = \frac{1}{|x||y|} \mathbb{1}_{x \mid y} \), \( \tilde{G} = [G, Y^T; Y, H] \), \( L \leftarrow \) graph Laplacian of \( G \), \( P = (1-p)WGG^T \). Initialize \( \tilde{Y} \) and \( \tilde{W} \) randomly.

Set \( Z_1 = (\frac{1}{2} \tilde{G}_g \tilde{G}_g + \beta Q)^{-1} \)

Set \( Z_2 = (\frac{1}{2} \tilde{G}_y \tilde{G}_y + \beta Q)^{-1} \)

Set \( Z_3 = (L + \alpha I)^{-1} \)

repeat

\[
W_y \leftarrow (\frac{1}{2}(Y - \tilde{G}_g^T \tilde{W}_y^T) \tilde{G}_y^T + \beta P_y)Z_1
\]

\[
W_g \leftarrow (\frac{1}{2}(Y^T - \tilde{G}_g \tilde{W}_y^T) \tilde{G}_y^T + \beta P_y)Z_2
\]

\[
\tilde{Y} \leftarrow \frac{1}{2} (W_y \tilde{G}_y + \tilde{G}_y W_y)Z_3
\]

until convergence

return \( \tilde{Y} \) and \( \tilde{G} \), as defined in (6)

Experimental Results

We evaluated MLLP, together with several other MLL and LP algorithms on three real-world datasets, two of which are from computational biology and the third is on social networks.

Datasets. 1) Yeast: we extracted the yeast PPI graph from HINT (Das and Yu 2012), which is a human-curated high-quality PPI dataset. The functional annotations (labels) of proteins were obtained from the SGD database. We kept 99 function labels that were associated with at least 20 proteins and removed proteins that were not annotated by any of these 99 functions. This resulted in a graph consisting of 2,925 nodes annotated with at least one label.

2) Human: the human PPI graph was also taken from HINT. The human proteins were annotated according to the Gene Ontology (Consortium and others 2000). We followed the same filtering steps as in the Yeast dataset, obtaining a PPI graph of 4,877 nodes and a set of 267 distinct labels.

3) LiveJournal: the whole dataset describes a social network from an online blogging community of more than 3 million users (Yang and Leskovec 2012). We randomly sampled a subgraph of the network consisting of a connected component of 11,921 users. Each user is associated with one or more user-defined groups. Given the observed network and the known groups associated with each user, the LP task is to predict if two users are likely to establish friendship, and the MLL task is to identify hidden user groups that a user is associated with.

Methods. We compared our method MLLP to several existing methods that solve the MLL and LP separately. Three state-of-the-art MLL algorithms that are designed to handle incomplete labels, i.e. MLR-GL (Bucak, Jin, and Jain 2011), I-BG (Wang, Huang, and Ding 2011), and ProtDM (Yu et al. 2013), were chosen. We also included three state-of-the-art LP methods: Kernel-based scores (Kernel) (Fouss et al. 2012), link propagation (LinkProp) (Kashima et al. 2009), and supervised link prediction (SLP) (Al Hasan et al. 2006).

To demonstrate the benefit of solving MLL and LP jointly, we included the marginalized denoising method (MDM) in Eq. (4) on the bi-relational graph, and the graph-based regularization framework (GR) in Eq. (1) as the baselines. We also include the alternating method (Alter) (Bilgic, Namata, and Getoor 2007) as a baseline. There are numerous choices of how the prediction results can be exchanged between MDM and GR at each iteration in Alter. We choose to add the 100 top-ranked edges predicted by MDM to the observed graph for GR and add 10 top-ranked instances for each label predicted by GR to the known labels for MDM because it leads to the best performance. We repeat this procedures for 20 times. We set the hyper-parameters of all methods using 5-fold cross-validation, except the corruption noise \( p \) of MLLP and MDM is set to a high value \( p = 0.95 \), throughout.

Experiment setup and evaluation metrics. Both MLL and LP are essentially tasks to infer missing information in a graph. To establish ground truth for comparing performance of the method considered, we removed labels and edges in three ways. 1) We randomly removed a fraction, \( \gamma \in \{0, 0.1, 0.2, 0.3, 0.4\} \), of the existing edges. 2) We randomly selected 50% of the nodes and remove all label information from them (these are the MLL test nodes). 3) To simulate various degrees of incomplete training information, we randomly removed a fraction \( \delta \in \{0, 0.1, 0.2, 0.3, 0.4\} \) of the label information \( Y \) of all remaining nodes. As there are five choices for \( \gamma \) and \( \delta \) we ultimately created 25 distinct setups for training. For both MLL and LP, we reported the AUC of the ROC curve (Huang and Ling 2005) of the re-constructed links and induced labels (on the testing nodes) averaged over five runs along with standard deviations.

Performance of link prediction. Fig. 2 summarizes the AUCs of the various methods for link prediction on the three datasets. In the first row, we plot the average AUCs as a function of the fraction of missing links, \( \gamma \). Here, we fixed the
fraction of missing training labels to $\delta = 0.2$. In the second row, we varied $\delta$ and set $\gamma = 0.2$. The baseline model, MDM, outperforms the three state-of-the-art methods, Kernel, LinkProp, and SLP, for all settings on the Yeast and Human datasets by a relatively large margin. It indicates that the marginalized denoising model is indeed promising in link prediction. Its performance on the LiveJournal dataset is not satisfactory though. We hypothesize that this may be due to the relatively high sparseness (lower graph density and node degree) of the graph, which makes it difficult to recover the hidden link correlations solely through corruption and denoising. The Alter method outperforms MDM when the fraction of missing labels ($\delta$) is small, but its performance decreases as $\delta$ increases. This is probably because when $\delta$ is large, the MLL part, GR, is more likely to make wrong predictions and thus makes the bi-relational graph of MDM unreliable. Our method MLLP, on the other hand, consistently outperforms the other methods on all three datasets. In particular, it boosts the performance of MDM significantly on the LiveJournal dataset, which indicates that solving the MLL and LP problems jointly significantly improves the performance of the LP accuracy. As expected, when the fraction of removed links $\gamma$ increases, the overall LP AUCs decreases. However, removing labels (increased $\delta$) has a less pronounced effect on the LP performance on the Yeast and Human datasets, probably because these two networks are relatively dense enough for the LP algorithms to infer links from the observed links alone.

**Performance of multi-label learning** Fig. 3 summarizes the MLL performance of various methods following the same setting as in Fig. 2. As shown, the baseline method GR for MLL is not as powerful as MDM for LP. It performs worse than most of the existing methods. The Alter improves upon GR in almost all settings. It is worth to note that as the fraction of missing links ($\gamma$) increases, the improvement of Alter becomes marginal. This is similar to the effect of $\delta$ on Alter in the LP task; i.e. the alternating method is prone to aggregating wrong predictions made by previous iterations when $\gamma$ is large. Our new method MLLP enhanced with the link prediction component, is able to considerably improve its performance. MLLP outperforms almost all other methods across different settings except for LiveJournal when
Related Works and Discussion

Link prediction. The existing works on LP can be roughly grouped into three categories. 1) Similarity based methods, e.g. Adamic-Adar Index (Adamic and Adar 2003), extract similarity scores between nodes based on graph topologies. The scores are later thresholded or ranked to form links. 2) Latent factor models explain the observed adjacency matrix using a small number of latent factors. The latent factors are usually learned through matrix factorization (Koren, Bell, and Volinsky 2009) or Bayesian models (Miller, Griffiths, and Jordan 2009). 3) Supervised methods treat LP as binary classification problems. MLLP for LP can be considered as a special instance of supervised learning methods, where we learn the similarity scores using a supervised learning scheme trained on conceptually “infinitely many” training examples. Our work may be most similar to the work (Chen and Zhang 2014), who already explore marginalized denoising for LP, however do not incorporate MLL.

Multi-label learning. Many existing works on MLL explicitly incorporate graph structure into models such as the traditional graph-based semi-supervised models (Zhou et al. 2003) and collective classification methods (Kong, Shi, and Zhang 2014), who already explore marginalized denoising for LP, however do not incorporate MLL.

Comparison of the joint model with the baselines. We define the relative improvement of MLLP over the baselines as \( \frac{\gamma}{\delta} - \frac{a_0}{a_1} \), where \( a_0 \) is the AUC score of MDM for LP or that of GR for MLL, and \( a_1 \) is the corresponding AUC score of MLLP. Fig. 4 shows the relative improvement in percentage under all 25 settings with varying fractions of missing labels (y-axis) and missing links (x-axis). A clear trend emerges showing that the relative improvement of MLLP upon the baselines increases as the fraction of missing labels and/or links increases. When the fraction of missing links is high, e.g. \( \gamma = 0.4 \), MLLP achieves more than 7% boost over MDM on the LiveJournal dataset for LP, and more than 10% improvement over GR on the Yeast and LiveJournal dataset for MLL. The improvement, especially on the Yeast and Human datasets for LP, is less evident as the fraction of missing labels \( \delta \) increases. This is consistent with our previous hypothesis that the labels in these two datasets adds relatively less information as the baseline is able to infer links from the observed links alone.

Running time. The time complexity of MLLP is \( O(n^3) \) due to matrix inversions and multiplications. The algorithm usually converges within 10 to 20 iterations. On a server with two 8-core Intel Xeon E5-2650@2.60GHz CPUs and 128GB RAM, our MATLAB implementation (accelerated with an NVIDIA K40 GPU) required about 170s for a run of MLLP until convergence on the LiveJournal dataset and around 23s on the Human dataset (without GPU 7m and 1m respectively). For perfect reproducibility, we provide the source code and data sets online at http://www.cse.wustl.edu/~chenz/codes/mllp.tar.gz.

Figure 4: Relative improvements of MLLP over the baseline methods MDM for LP (upper row) and GR for MLL (lower row). The plots show the relative increase in LP AUC (%) and MLL AUC (%) as the fractions of missing labels (x-axis) and missing links (y-axis) varies.

\[ \gamma = 0, \text{ where it is comparable to I-BG.} \] It is worth noting that MLLP improves upon the baselines (MDM for LP and GR for MLL) more pronouncedly as either \( \gamma \) or \( \delta \) increases.

Joint link prediction and multi-label learning. Although a collection of methods have been developed to solve the LP and MLL problems independently, little effort has been devoted to solving the two problems jointly. There are works that have alluded to the close connection between the two problems. For example, the graph Laplacian (Zhou and Schölkopf 2004; Fouss et al. 2012) and variants of graph kernels serve as the main building blocks for many similarity-based LP and MLL methods. Bilgic et. al. (2007) proposed to alternately apply LP and MLL algorithms and showed that it can improve the performance of both tasks. However, it is difficult to prevent unreliable predictions being exchanged from one task to the other. This problem is further exacerbated by missing links and labels as shown in our experiment. MLLP, on the other hand, combines LP and MLL seamlessly into a joint optimization and enables both tasks to benefit from each other.
Acknowledgments

ZC and WZ are supported in part by Natural Science Foundation of China (31300999), the municipal government of Wuhan, Hubei, China (2014070504020241 and the Talent Development Program), and an internal research grant of Jianghan University, Wuhan, China, as well as by United States National Institutes of Health (R01GM100364). KQW is supported by NSF grants IIS-1149882, EFRI-1137211. Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the National Science Foundation.

References