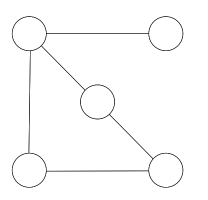
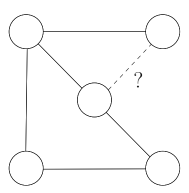
Data Mining 2020 Mining Social Network Data: Link Prediction

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The Link Prediction Problem





Applications

- Biology: protein-protein interaction prediction.
- Recommendation systems, e.g. link recommendation in social networks like Facebook.
- Analysis of criminal/terrorist networks.
- Automatic web hyper-link creation (e.g. discovering missing links in WikiPedia).
- Record linkage/deduplication.
- ...

The Link Prediction Problem

Given a social network G = (V, E) in which an edge $e = (u, v) \in E$ represents some form of interaction between its endpoints at a particular time t(e).

Multiple interactions between the same pair of nodes can be recorded by parallel edges or using a complex time-stamp for an edge.

The Link Prediction Problem

For time $t \leq t'$, let G[t, t'] denote the subgraph of G restricted to the edges with time-stamps between t and t'.

Supervised learning problem: choose a training interval $[t_0, t_0']$ and a test interval $[t_1, t_1']$ where $t_0' < t_1$.

The link prediction problem is to output a list of edges not present in $G[t_0, t_0']$, but are predicted to appear in the network $G[t_1, t_1']$.

Node Neighborhood Based Features

Let $\Gamma(x)$ denote the neighborhood of node x, that is, the set of nodes directly connected to x.

For two nodes x and y, we define:

• Number of shared neighbors:

Common-Neighbors
$$(x, y) = |\Gamma(x) \cap \Gamma(y)|$$

2

Jaccard-Coefficient
$$(x, y) = \frac{|\Gamma(x) \cap \Gamma(y)|}{|\Gamma(x) \cup \Gamma(y)|}$$

A shared neighbor that is itself not heavily connected gets higher weight:

Adamic-Adar
$$(x, y) = \sum_{z \in \Gamma(x) \cap \Gamma(y)} \frac{1}{\log |\Gamma(z)|}$$

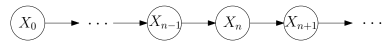
Intermezzo: Markov Chain

Let P be a $k \times k$ matrix with elements P_{ij} . A random process (X_0, X_1, \ldots) with state space $S = \{s_1, \ldots, s_k\}$ is said to be a Markov chain with transition matrix P if for all $i, j \in \{1, \ldots, k\}$ and all $i_0, \ldots, i_{n-1} \in \{1, \ldots, k\}$ we have

$$P(X_{n+1} = s_j \mid X_0 = s_{i_0}, X_1 = s_{i_1}, \dots, X_{n-1} = s_{i_{n-1}}, X_n = s_i)$$

= $P(X_{n+1} = s_j \mid X_n = s_i)$
= P_{ij}

Slogan: the future is independent of the past given the present.



Stationary Distribution

Let $(X_0, X_1, ...)$ be a Markov chain with state space $S = \{s_1, ..., s_k\}$ and transition matrix P. A row vector $\pi = (\pi_1, ..., \pi_k)$ is said to be a stationary distribution for the Markov chain, if it satisfies:

- \bullet $\pi_i \geq 0$ for $i = 1, \ldots, k$ and $\sum_{i=1}^k \pi_i = 1$, and
- $P = \pi$, meaning that

$$\sum_{i=1}^k \pi_i P_{ij} = \pi_j, \text{ for } j = 1, \dots, k$$

The second property implies that if the initial distribution $\mu^{(0)} \equiv P(X_0)$ equals π , then the distribution $\mu^{(1)}$ of the chain at time 1 satisfies:

$$\mu^{(1)} = \mu^{(0)}P = \pi P = \pi,$$

and by iterating we see that $\mu^{(n)} = \pi$ for every n.

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Existence and Convergence

Every irreducible and aperiodic Markov chain has a stationary distribution.

If we run the Markov chain for a sufficiently long time n, then regardless of what the initial distribution $\mu^{(0)}$ was, the distribution $\mu^{(n)}$ at time n will be close to the stationary distribution π . This is often referred to as the Markov chain approaching equilibrium as $n \to \infty$.

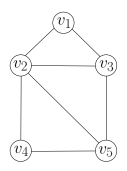
Random Walk on Graph

A random walk on a graph G=(V,E) is a Markov chain with state space $V=\{v_1,v_2,\ldots,v_k\}$.

If the random walker stands at vertex v_i at time n, then it moves at time n+1 to one of the neighbors of v_i chosen at random, with equal probability for each of the neighbors. More formally:

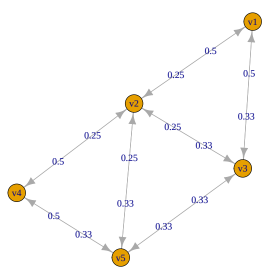
$$P_{ij} = \left\{ egin{array}{ll} rac{1}{|\Gamma(v_i)|} & ext{if } v_j \in \Gamma(v_i) \\ 0 & ext{otherwise} \end{array}
ight.$$

Example Graph and Transition Matrix



$$P = \begin{pmatrix} 0 & \frac{1}{2} & \frac{1}{2} & 0 & 0 \\ \frac{1}{4} & 0 & \frac{1}{4} & \frac{1}{4} & \frac{1}{4} \\ \frac{1}{3} & \frac{1}{3} & 0 & 0 & \frac{1}{3} \\ 0 & \frac{1}{2} & 0 & 0 & \frac{1}{2} \\ 0 & \frac{1}{3} & \frac{1}{3} & \frac{1}{3} & 0 \end{pmatrix}$$

Transition Diagram



Stationary Distribution

The stationary distribution is:

$$\pi = \left(\frac{2}{14}, \frac{4}{14}, \frac{3}{14}, \frac{2}{14}, \frac{3}{14}\right)$$

Check

$$\left(\frac{2}{14}, \frac{4}{14}, \frac{3}{14}, \frac{2}{14}, \frac{3}{14}\right) \begin{pmatrix} 0 & \frac{1}{2} & \frac{1}{2} & 0 & 0\\ \frac{1}{4} & 0 & \frac{1}{4} & \frac{1}{4} & \frac{1}{4}\\ \frac{1}{3} & \frac{1}{3} & 0 & 0 & \frac{1}{3}\\ 0 & \frac{1}{2} & 0 & 0 & \frac{1}{2}\\ 0 & \frac{1}{3} & \frac{1}{3} & \frac{1}{3} & 0 \end{pmatrix} = \left(\frac{2}{14}, \frac{4}{14}, \frac{3}{14}, \frac{2}{14}, \frac{3}{14}\right)$$

For example: in the long run, the random walker will find himself in node v_2 for $\frac{4}{14}\approx 29\%$ of the time.

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Path Based Features

For two nodes x and y:

- Shortest path distance between x and y.
- Katz:

$$\mathsf{Katz}\,(x,y) = \sum_{\ell=1}^{\infty} \beta^{\ell} |\mathsf{paths}_{\mathsf{x},y}^{\langle\ell\rangle}|$$

where $|\mathsf{paths}_{x,y}^{\langle\ell\rangle}|$ is the number of length- ℓ paths from x to y. A very small β yields predictions much like common neighbors, since paths of length three or more contribute very little to the summation.

- Hitting time H(x, y) between nodes x and node y is the expected number of steps in a random walk starting from node x before node y is visited for the first time.
- Commute time: C(x, y) = H(x, y) + H(y, x).

Path Based Features

• Rooted Pagerank(x, y): stationary probability of y in a random walk that returns to x with probability α each step, and moving to a random neighbor with probability $1 - \alpha$.

Vertex Feature Aggregation

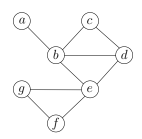
- **1** Preferential Attachment Score between x and y: $|\Gamma(x)| \cdot |\Gamma(y)|$.
- ② Sum of Neighbors of x and y: $|\Gamma(x)| + |\Gamma(y)|$.
- Clustering coefficient:

clustering-coefficient
$$(u) = \frac{2 |\{(v, w) \in E : v, w \in \Gamma(u)\}|}{|\Gamma(u)|(|\Gamma(u)| - 1)}$$

This is the number of neighbor pairs of u that are neighbors of each other, divided by the total number of neighbor pairs of u. For example, if edges represent collaborations between people (nodes), it's the fraction of pairs of a person's collaborators who have also collaborated with one another.

For a pair of nodes, we can use the sum or product of their clustering coefficients as a feature.

Examples



- Shortest path distance between c and e is 2.
- $\Gamma(c) = \{b, d\}$ $\Gamma(e) = \{b, d, f, g\}$
- Common-Neighbors $(c,e) = |\Gamma(c) \cap \Gamma(e)| = |\{b,d\}| = 2$
- Jaccard-Coefficient $(c, e) = \frac{|\Gamma(c) \cap \Gamma(e)|}{|\Gamma(c) \cup \Gamma(e)|} = \frac{|\{b, d\}|}{|\{b, d, f, g\}|} = \frac{2}{4}$
- clustering-coefficient (e) = $\frac{2\times2}{4\times3} = \frac{1}{3}$

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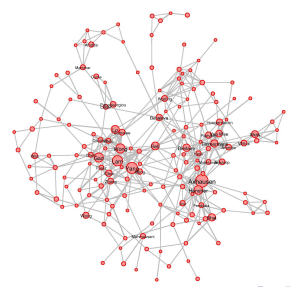
Required Literature

David Liben-Nowell, Jon Kleinberg: *The Link Prediction Problem for Social Networks*, Proceedings of the Twelfth Annual ACM International Conference on Information and Knowledge Management (CIKM'03), November 2003, pp. 556–559.

Mohammad Al Hasan, Vineet Chaoji, Saeed Salem, and Mohammed Zaki: Link Prediction Using Supervised Learning, SDM Workshop on Link Analysis, 2006.

The remaining slides are about the second paper.

Co-authorship Network



Data Sets

Dataset	Number of Papers	Number of Authors
BIOBASE	831,478	156,561
DBLP	540,459	1,564,617

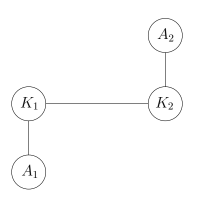
Consider the pairs of nodes not linked in $G[t_0, t'_0]$, and give them class label 1 (positive) if they are linked in $G[t_1, t'_1]$, and class label 0 (negative) otherwise.

- BIOBASE: 5 years of data from 1998 to 2002 (first 4 for training).
- DBLP: 15 years of data, from 1990 to 2004 (first 11 for training).
- Positive/Negative pairs chosen randomly in equal proportion from pairs that qualify.
- Construct feature vector for each pair of authors.

Some Additional Features Used

- Keyword match count (x, y): the number of shared keywords of papers written by x and papers written by y.
- Sum of keyword count: researchers that have a wide range of interests or those who work on interdisciplinary research usually use more keywords. In this sense they have better chance to collaborate with new researchers.
- Shortest distance in author-keyword graph: the author-keyword graph extends the co-authorship graph with nodes that correspond to keywords. Each keyword node is connected to an author node, if that keyword is used by the author in any of his papers. Moreover, two keywords that appear together in any paper are also connected by an edge.

Author-Keyword Graph



- Author 1 wrote a paper with Keyword 1.
- Author 2 wrote a paper with Keyword 2.
- Keyword 1 and Keyword 2 appeared together in some paper.

Experiments

- Seven different classification algorithms, among which classification (decision) trees and naive Bayes.
- Used 5-fold cross-validation to evaluate performance.
- Separate validation set for hyperparameter tuning.
- Measure accuracy, precision, recall, F_1 score.

Results

Classification model	Accuracy	Precision	Recall	F-value	Squared Error
Decision Tree	90.01	91.60	89.10	90.40	0.1306
SVM(Linear Kernel)	87.78	92.80	83.18	86.82	0.1221
SVM(RBF Kernel)	90.56	92.43	88.66	90.51	0.0945
K_Nearest Neighbors	88.17	92.26	83.63	87.73	0.1826
Multilayer Perceptron	89.78	93.00	87.10	90.00	0.1387
RBF Network	83.31	94.90	72.10	81.90	0.2542
Naive Bayes	83.32	95.10	71.90	81.90	0.1665
Bagging	90.87	92.5	90.00	91.23	0.1288

 ${\it Table 2: Performance of different classification algorithms for BIOBASE\ database}$

Classification model	Accuracy	Precision	Recall	F-value	Squared Error
Decision Tree	82.56	87.70	79.5	83.40	0.3569
SVM(Linear Kernel)	83.04	85.88	82.92	84.37	0.1818
SVM(RBF Kernel)	83.18	87.66	80.93	84.16	0.1760
K_Nearest Neighbors	82.42	85.10	82.52	83.79	0.2354
Multilayer Perceptron	82.73	87.70	80.20	83.70	0.3481
RBF Network	78.49	78.90	83.40	81.10	0.4041
Naive Bayes	81.24	87.60	76.90	81.90	0.4073
Bagging	82.13	86.70	80.00	83.22	0.3509

Table 3: Performance of different classification algorithms for DBLP dataset

Feature Ranking

Attribute name	Information gain	Gain Ratio	Chi-Square	SVM feature	Avg. Rank
			Attribute Eval.	evaluator	
Sum of Papers	3	4	3	4	3
Sum of Neighbors	1	3	1	2	2
Sum of KW count	6	6	6	3	5
Sum of Classification	5	5	5	6	5
count					
KW match count	2	1	2	1	1
Sum of log of Sec.	7	7	7	8	7
Neighbor. count					
Shortest distance	4	2	4	5	4
Clustering Index	9	9	9	7	8
Shortest dist. in	8	8	8	9	8
KW-Author graph					

 ${\it Table 4: Rank of different Attributes using different algorithms for BIOBASE dataset}$

Attribute name	Information gain	Gain Ratio	Chi-Square	SVM feature	Avg. Rank
			Attribute Eval.	evaluator	
Sum of Papers	4	4	4	2	4
Sum of Neighbors	3	3	3	4	3
Shortest distance	1	1	1	1	1
Second shortest distance	2	2	2	3	2

 ${\it Table 5: Rank of different Attributes using different algorithms for DBLP \ dataset}$

Feature Distributions

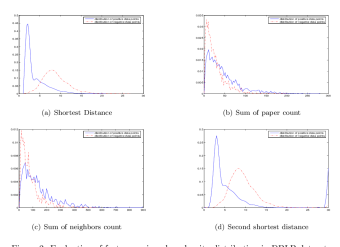
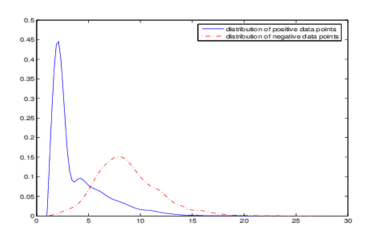


Figure 2: Evaluation of features using class density distribution in DBLP dataset

Feature Distributions



(a) Shortest Distance